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Investigation of Role of Nitric Oxide in Protection from
Bordetella pertussis Respiratory Challenge
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The mechanism whereby whole-cell pertussis vaccines (WCV) confer protection against Bordetella pertussis is still not fully understood. We have previously reported that macrophage activation produced by vaccination with WCV is associated with induction of NO synthesis by macrophages in response to in vitro stimulation with B. pertussis antigens. To determine whether NO production is an effector of protection or simply a marker of activation, the susceptibility of inducible nitric oxide synthase (type II, iNOS) knockout mice to infection with B. pertussis was examined. We showed that iNOS knockout mice were more susceptible to B. pertussis respiratory challenge than wild-type mice. iNOS-deficient mice also developed a less effective protective response than wild-type mice after the same immunization with WCV. This suggests that NO plays an important role in effecting protection against B. pertussis challenge.

Bordetella pertussis is a human pathogen which possesses tropism for the respiratory system, causing an acute and sometimes persistent disease. Although pertussis vaccines have been in use for mass vaccination in most countries for many years and have led to a major decrease in the incidence of pertussis, the mechanism by which they induce protection against pertussis in children is still unclear. Recent evidence indicates that B. pertussis is a facultatively intracellular organism and that clearance involves activated macrophages (4, 15, 20, 21). The mechanism whereby macrophage activation results in the killing of facultatively intracellular pathogens is still incompletely determined. However, it has become increasingly apparent in recent years that NO and reactive nitrogen intermediates (nitrite and peroxynitrite) are potentially important mediators of the immune system (1). Production of NO by activated murine macrophages has been implicated as an antimicrobial effector mechanism against several pathogens (2, 5, 9). We have reported previously that macrophage activation produced by vaccination with a whole-cell pertussis vaccine (WCV) is associated with induction of NO synthesis by macrophages in response to in vitro stimulation with B. pertussis antigens (20). The presence of small quantities of active pertussis toxin seems to be important for this process (21). The relationship between NO induced in macrophages in response to in vitro culture with bacterial antigens and protection in vivo in the mouse intracerebral challenge model indicates that macrophage activation is involved in protective immunity (20). However, it is not clear from these studies whether NO is an effector of protection or simply a coincidental marker of activation.

To clarify further the role of NO in protection against B. pertussis challenge, the induction of NO synthesis by macrophages and protection in vivo against aerosol challenge induced by a conventional WCV and the new-generation acellular pertussis vaccine (ACV) was investigated in inducible nitric oxide synthase (iNOS)-deficient mice.

MATERIALS AND METHODS

Vaccines. A WCV (National Institute for Biological Standards and Control [NIBSC] reagent 88/522, 3rd British Reference Preparation; potency, 50 IU/ampoule) (14) and a commercially available three-component ACV containing 25 g of pertussis toxoid (PT) chemically detoxified with formaldehyde and glutaraldehyde, 25 g of filamentous hemagglutinin (FHA), and 8 g of pertactin (PRN) per single human dose (SHD), in combination with diphtheria and tetanus toxoids (DTPa), was used for the immunization. All other reagents were of analytical grade.

Animals. iNOS-deficient mice and their wild-type littermates were generated as described previously (17). The murine iNOS gene was disrupted by homologous recombination in 129sv embryonic stem (ES) cells. The recombinant allele was passed through the germ line following mating of ES cell chimeras with 129sv (Harlan UK Ltd., Oxford, United Kingdom). The homozygous, heterozygous, and wild type littermates of the 129sv strains were used at the ages of approximately 3 to 4 weeks.

Immunogenicity study. Groups of five mice were immunized (intraperitoneally [i.p.]) with ACV at 0.25 SHD per dose and with WCV at 0.125 IU per dose (which is equivalent to approximately 0.03 SHD), and both vaccines were diluted in phosphate-buffered saline (PBS). Mice in the control group received PBS. Mice were terminally bled at 4 weeks postimmunization, and sera from individual animals were assayed for total immunoglobulin G (IgG) antibodies to the B. pertussis antigens PT, FHA, and PRN by a standard enzyme-linked immunosorbent assay (ELISA). The geometric mean ELISA units (EU) of the antibody to each antigen were calculated against the First World Health Organization (WHO) International Reference Anti-B. pertussis Serum (Mouse) (19). All the serum samples were always analyzed in parallel with the reference antiserum on the same plate. Relative concentrations of IgG1 and IgG2a specific for the B. pertussis antigens PT, FHA, and PRN were measured by using sheep anti-mouse IgG subclass-biotin and horseradish peroxidase-avidin conjugates (PharMingen) (11). Specific responses for each subclass were presented as the ratio of the optical density at 492 nm (OD492nm) of the test sample to the OD492nm of the reference serum used in each plate.

Bacterial antigens. Heat-killed B. pertussis 18.323 cells (HKC) were prepared by incubation of bacterial cells (5 × 10⁸/ml) in PBS at 80°C for 30 min (20). Purified detoxified PT, FHA, and PRN were kindly provided by GlaxoSmithKline, Rixensart, Belgium.

Macrophages. Mice were immunized with WCV or ACV at the indicated doses. Control mice received PBS. Macrophage cultures were prepared according to the method described by Torre et al. (16). In brief, mice were terminally bled on the indicated day postimmunization. The peritoneal cavity was then
were regarded as statistically signi
cance of differences between two groups.

Griess reagent (16), and the viability was checked by trypan blue exclusion before and after incubation.

phages were cultured in a total volume of 1 ml/well with 2°C under 5% CO2 in 90% humidity for 24 h. Cell cultures were incubated at 37°C for 18.323 for 5 min by use of a custom-made aerosol apparatus (22). The lungs and tracheas were removed from each group at the indicated time point and were homogenized in 1 ml of 1% casein solution by means of a mini-bead beater using 2.5- to 3.5-mm-diameter glass balls. Viable counts were performed on the homogenate by diluting across microtiter plates and a mini-bead beater using 2.5- to 3.5-mm-diameter glass balls. Viable counts were performed by four replicates. Each data point represents the mean CFU per lung. Error bars, standard deviations.

In vivo aerosol challenge. Bacterial suspensions (B. pertussis 18.323), stored at −70°C in 5% glycerol, were spread on charcoal agar plates (charcoal agar base plus 10% defibrinated horse blood) and incubated at 37°C for 2 days. Two further subcultures were performed with incubation for 16 to 18 h at the same conditions. Bacterial cells were harvested and resuspended in 0.9% saline containing 1% casein and adjusted to an OD625 of 0.2 by using a spectropho-
tometer (MSE-Fisons, Loughborough, United Kingdom). The suspension was kept on ice until it was used for aerosol challenge. Aerosol challenge was performed on groups of five previously immunized mice that were exposed to B. pertussis 18.323 for 5 min by use of a custom-made aerosol apparatus (22). The lungs and tracheas were removed from each group at the indicated time point after challenge and were homogenized in 1 ml of 1% casein solution by means of a mini-bead beater using 2.5- to 3.5-mm-diameter glass balls. Viable counts were then performed on the homogenate by diluting across microtiter plates and plating onto charcoal agar plates. The mean viable count per lung from the two groups of mice, approximately 2.5 to 3.0 log units by day 7 to 10. However, bacterial counts subsequently declined more rapidly in wild-type mice than in iNOS knockout mice. Thus, by day 18, bacterial counts in iNOS-deficient mice were at least 0.5 log unit higher than those in wild-type mice, and this difference (P < 0.05) was maintained until the end of the experiment.

To assess the ability of murine peritoneal macrophages to generate NO in response to in vitro stimulation with HKC, mice were immunized with WCV or ACV. The peak time for production of NO in the macrophage cultures was approximately 10 to 15 days postimmunization (data not shown). Therefore, in this study all macrophages were isolated from mice on day 15 post immunization.

Peak NO concentrations in macrophage cultures from the wild type group were approximately threefold higher than those from the mutant group. Furthermore, macrophages from mice lacking iNOS produced lower NO concentrations in their cultures after in vitro restimulation with HKC than cells recovered from wild-type mice (Fig. 2a). This corresponded closely to the resistance to aerosol challenge observed in vivo in the two types of mice (Fig. 2b), where the CFU count per lung was approximately 2 log units lower in wild-type than in mutant mice.

Macrophages from mice immunized with ACV were also assayed for NO production with and without the addition of HKC. Unlike those from mice immunized with WCV, macrophages from mice immunized with ACV did not produce NO concentrations higher than those for the control group before the stimulation in vitro (baseline) (Fig. 3a). However, these macrophages were able to produce NO in response to B. pertussis HKC stimulation, although the levels were lower than those for the WCV-immunized group. NO concentrations in cultures from the wild-type group were more than threefold higher than those in cultures from the homozygous group. This also closely paralleled the pattern of in vivo protection against aerosol challenge, where CFU counts in the lungs of the mutant group were approximately 1.5 log units higher (P < 0.05) than those in the lungs of the wild-type group (Fig. 3b).

To investigate the relationship of the NO produced by acti-
vated macrophages to host defense against B. pertussis challenge, different immunization doses were used. Figure 4a shows that the synthesis of NO by macrophages in response to HKC restimulation was immunization dose dependent, and again, macrophages from the wild-type group had higher NO production than those from the mutant group. It was noted in the protection study that when mice were immunized with WCV at 1.0 IU/dose, similar reductions (of approximately 2.5 to 3.0 log units) in lung CFU counts occurred in both wild-type and mutant groups in comparison with the control group (Fig. 4b). This was in spite of the fact that cells from wild-type mice produced higher NO levels in their cultures than those from the mutant group. Reduction of the immunization dose from 1.0 to 0.25 IU resulted in different protection profiles for these two groups of mice, which corresponded with the NO concentrations achieved in their macrophage cultures. That is, the mutant mice showed a lower level of protection than the wild-type mice.

To investigate further the cellular and humoral immune responses induced by immunization of these two types of mice with WCV and ACV, pertussis-specific antibody production,
macrophage activation, and in vivo protection were assessed 4 weeks after immunization. Macrophages from both wild-type and mutant mice immunized with WCV produced approximately 5.5-fold-higher NO concentrations in their cultures after stimulation with HKC than macrophages from the ACV-immunized group (P < 0.05) (Fig. 5a). The NO concentration in macrophage cultures from the mutant group immunized with WCV was approximately half that for the wild-type group.

There was no difference (P > 0.05) in NO induction between the mutant group immunized with ACV and the control group.

Titers of antibody to PT, FHA, and PRN (69 kDa) were much higher in the ACV group than in the WCV group (Table 1). The ratio of IgG1 to IgG2a showed that mice immunized with WCV gave a response shifted towards Th1, whereas those immunized with ACV gave a response biased towards Th2. No difference in antibody production was found between wild-type
and mutant mice immunized with WCV. However, among mice immunized with ACV, lower levels of antibodies to FHA and PRN were observed in the mutant group than in the wild-type group. Aerosol challenge of mice at 4 weeks after immunization showed that mice immunized with WCV had developed better protection \( (P < 0.05) \) than mice immunized with ACV despite higher antibody responses developing in the latter group (Fig. 5b). Comparison of wild-type mice and mutant mice showed that the former group were better protected from the challenge than the latter group \( (P < 0.05) \) when immunized with WCV or ACV.

**DISCUSSION**

Infection by *B. pertussis*, usually manifesting as pertussis (whooping cough), is still an important cause of morbidity and mortality among children in many parts of the world (18). It is also being recognized increasingly as a significant agent of respiratory disease in adults (Editorial, Lancet 339:526–527, 1992). The results of recent clinical trials have indicated that both established WCVs and the new generation of ACVs can stimulate protection in children (6, 7). We have previously reported that macrophage activation produced by vaccination...
with WCV is associated with induction of NO synthesis by macrophages in response to in vitro stimulation with *B. pertussis* antigens (20, 21). In the present study, INOS knockout mice were used to further examine the role of NO in protection from *B. pertussis* respiratory challenge.

It has been reported previously that INOS-deficient mice produced more gamma interferon (IFN-γ) and less interleukin 12 (IL-12) than the wild-type mice following antigenic stimulation (10, 12, 13). In the present study, the results showed that elimination of *B. pertussis* from the lungs of infected INOS knockout mice was slower than elimination from the lungs of wild-type mice after respiratory challenge. The INOS-deficient mice also showed a lower level of protection than the wild-type mice following the same immunization with WCV or ACV in spite of producing a greater IFN-γ response to the bacterial antigens in vitro (data not shown). This provides a further indication that NO is an important effector molecule in protection against *B. pertussis* challenge.

Macrophages from wild-type and mutant mice immunized with the WCV all produced NO in response to in vitro stimulation with bacterial cells. This suggests that this type of vaccine is a very powerful inducer of NO synthase even in INOS-deficient mice. That the latter still produced some NO in spite of the gene disruption may be attributable to the operation of compensatory mechanisms through other pathways. Constitutive NOS may also have contributed a background level of NO.

It should be noted that the INOS gene-targeting construct was produced by terminal extension and integration into the INOS gene. This disrupts the gene and should prevent expression but may not completely eliminate it (3, 8). In the present study, these mice produced substantially less NO than wild-type mice. However, NO may not be the only effector of protection. It was notable that when mice were immunized with higher doses of the vaccine, there was no difference in protection between wild-type and mutant mice, even though the wild-type mice produced more NO in their macrophage cultures. This may have been because the high vaccine doses stimulated an adequate NO response in the knockout mice and the greater amount produced by the wild-type mice added nothing further to protection, or it may suggest that at high vaccine doses another mechanism comes into play which is not dependent on the bactericidal action of NO.

Macrophages isolated from mice immunized with WCV produced larger amounts of NO than those from the control group without additional stimulation. Our previous studies showed that this NO production was increased by adding HKC but not by IFN-γ and that NO induced by HKC was less effectively blocked by concentrations of anti-IFN-γ which completely blocked NO production in control cell cultures (20). Taken together, these results suggested that these macrophages that had already been activated in vivo. It is noteworthy that macrophages from mice immunized with ACV did not produce NO in vitro in the absence of stimulant. However, NO production was significantly increased by the addition of HKC, and this was clearly associated with protection in vivo. These results suggested that there might be a difference in the degree of macrophage activation produced in vivo by immunization with these two different types of vaccines. Furthermore, mice immunized with ACV produced lower IFN-γ levels (data not shown) in culture than those immunized with WCV after stimulation in vitro. This may indicate that mice immunized with WCV developed a stronger Th1-type response than those that received ACV.

Although pertussis vaccination is used throughout the world and has made a major contribution to decreasing morbidity and mortality from pertussis, its precise mode of action is still unclear. There is, however, increasing recognition of the importance of cell-mediated immunity in protection against *B. pertussis*. The present study using INOS knockout mice has provided direct evidence that the reactive nitrogen intermediates play an important role in the immune response induced by both WCV and ACV and that this is associated with protective immunity in vivo. This adds further weight to the hypothesis that activation of the killing mechanisms of macrophages helps to eliminate intracellular *B. pertussis* and hence to clear infection.

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