Intranasal Immunization with SAG1 and Nontoxic Mutant Heat-Labile Enterotoxins Protects Mice against Toxoplasma gondii

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Effective protection against intestinal pathogens requires both mucosal and systemic immune responses. Intranasal administration of antigens induces these responses but generally fails to trigger a strong protective immunity. Mucosal adjuvants can significantly enhance the immunogenicities of intranasally administered antigens. Cholera toxin (CT) and heat-labile enterotoxin (LT) are strong mucosal adjuvants with a variety of antigens. Moreover, the toxicities of CT and LT do not permit their use in humans. Two nontoxic mutant LTs, LTR72 and LTK63, were tested with Toxoplasma gondii SAG1 protein in intranasal vaccination of CBA/J mice. Vaccination with SAG1 plus LTR72 or LTK63 induced strong systemic (immunoglobulin G [IgG]) and mucosal (IgA) humoral responses. Splenocytes and mesenteric lymph node cells from mice immunized with LTR72 plus SAG1, but not those from mice immunized with LTK63 plus SAG1, responded to restimulation with a T. gondii lysate antigen in vitro. Gamma interferon and interleukin 2 (IL-2) production by splenocytes and IL-2 production by mesenteric lymph node cells were observed in vitro after antigen restimulation, underlying a Th1-like response. High-level protection as assessed by the decreased load of cerebral cysts after a challenge with the 76K strain of T. gondii was obtained in the group immunized with LTR72 plus SAG1 and LTK63 plus SAG1. They were as well protected as the mice immunized with the antigen plus native toxins. This is the first report showing protection against a parasite by using combinations of nontoxic mutant LTs and SAG1 antigen. These nontoxic mutant LTs are now attractive candidates for the development of mucosally delivered vaccines.

The intracellular protozoan parasite Toxoplasma gondii infects all mammalian cells and is responsible for toxoplasmosis. Toxoplasmosis is generally harmless in an immunocompetent host, but it can cause severe damage to the fetus during pregnancy and is often lethal for immunodeficient subjects unless they are treated (22). T. gondii infection is also an important problem in animal breeding because it causes the deaths of many fetuses in cattle and sheep, for example (6). The natural site of infection for T. gondii is the mucosal surface of the intestine. Protective immunity obtained after a natural infection with T. gondii points to the importance of developing a vaccine that stimulates mucosal defenses. One major mechanism of protection against toxoplasmosis is considered to be systemic cell-mediated immunity with gamma interferon (IFN-γ) induction (3, 4, 33, 34, 46). SAG1 is a vaccine candidate of interest for a protective immune response in toxoplasmosis. Partial protective immunity has been produced by immunization with the SAG1 antigen, the major T. gondii surface antigen, which accounts for 3 to 5% of the total parasite protein (5, 13, 14, 35). The ability to increase systemic and mucosal responses against T. gondii may be of great importance for the development of an efficient vaccine. Furthermore, the intranasal route requires less antigen than the oral route because there is much less proteolytic activity in the nasal cavity. This route effectively promotes the production of both systemic and mucosal immune responses to an antigen (49, 61–63). Many if not most antigens trigger only weak or poor mucosal immune responses when given alone. The use of a mucosal adjuvant such as cholera toxin (CT) from Vibrio cholerae or heat-labile enterotoxin (LT) from toxigenic strains of Escherichia coli is necessary to enhance an immune response (14, 20, 21, 29, 39). The amino acid sequences of CT and LT are 80% identical (12). They have two functional domains, the A subunit, with ADP ribosyltransferase activity, and the pentameric B subunit, which is responsible for the toxin binding to the GM1 ganglioside receptor at the cell surface (28). The A subunit is toxic for eucaryotic cells, as it activates the Gs protein that binds to GTP. The Gs protein regulates the intracellular production of cyclic AMP. An increase in cyclic AMP stimulates the secretion of electrolytes and the osmotic movement of water in the gut lumen, which is responsible for profuse watery diarrhea (60).

CT and LT cannot be included in vaccine formulations for use in humans because of their toxicity. Therefore, the construction of nontoxic mutant of E. coli is important for the development of mucosal vaccines. Several mutant LTs have been constructed by site-directed mutagenesis (48). Two of these have been tested in our model of vaccination against T. gondii. The mutant toxin LTR72 (substitution of Arg for Ala at position 72) has reduced enzymatic and toxic activities (25). The other mutant, LTK63 (substitution of Ala for Ser at position 63 of the catalytic site) has neither enzymatic nor toxic activities (15, 25). LTR72 and LTK63 act as appropriate mucosal adjuvants following oral and intranasal immunization with various antigens and trigger local and systemic immune
responses (15, 47). Intranasal immunization with influenza virus hemagglutinin in combination with LTR72 induces serum immunoglobulin G (IgG) and mucosal IgA antibodies and neutralization of the virus with the development of a systemic Th1 response (2). These adjuvants have also been used with bacterial antigen and protect against infection with *Helicobacter pylori* after intragastric vaccination (41), with *Streptococcus pneumoniae* after nasal vaccination (30), and with *Bordetella pertussis* as strongly as bacterial antigen with LT (52). These results indicate that ADP ribosyltransferase activity is not necessary for adjuvant activity.

In previous studies, we have shown that intranasal vaccination with SAG1 plus CT protects mice against *T. gondii* (14). Since CT is too toxic to be used in humans, we have now investigated the capacities of mutant LTR72 and LTK63 to enhance the immunogenicity of intranasally administered SAG1.

**MATERIALS AND METHODS**

**Animals.** Pathogen-free female inbred CBA/J mice were used at 6 to 8 weeks of age (Janvier, Le Genest St. Isle, France).

**Immunization.** Mice were divided into groups of five. Two of the RH strain of *T. gondii* were harvested from the peritoneal fluid of Swiss OF1 mice that had been intraperitoneally infected 3 to 4 days earlier. They were used to prepare *T. gondii* lysate antigen (TLA). Cysts of the 76K strain of *T. gondii* were obtained from the brains of CBA/J mice infected 1 month previously.

**Adjuvants and antigen.** Wild-type CT and LT were purchased from Sigma, and the nontoxic mutant LTs (LTR72 and LTK63) were kindly provided by Chiron (Siena, Italy). They were used as adjuvants in combination with SAG1 protein purified from TLA by immunoaffinity (14). The mutant LTs LTR72 and LTK63 were obtained as previously described (25).

**Immunization.** The mucosal immunogenetics and adjuvant activities of mutant toxins were tested by immunizing groups of 10 mice intranasally two times at 28-day intervals with SAG1 (10 μg), LTR72 (1 μg), LTK63 (1 μg), LT (1 μg), or CT (1 μg) alone (defined as control groups) or with the combination SAG1 plus LTR72, SAG1 plus LTK63, SAG1 plus LT, or SAG1 plus CT (10 μg of protein and 1 μg of toxin). Each dose of immunogen was diluted to a final volume of 16 μl in phosphate-buffered saline (10 mM phosphate, 140 mM NaCl [PBS]) and was instilled into the nostrils of anesthetized mice with a micropipettor (6 μl/nasal). The experimental design included a group of untreated mice. The day before and 10 days after the boost, blood was collected by retro-orbital puncture. All samples were kept frozen (−20°C) until assayed for antibody activity.

**Antigen-driven cell-proliferative responses.** Three mice per group were sacrificed at day 42. Splenectomized lymph nodes, and nasal-associated lymphoid tissue (NALT) were harvested under sterile conditions and processed through a nylon mesh. Single-cell suspensions were obtained by filtration through nylon mesh to remove tissue debris. Hypotonic shock (0.155 M NH₄Cl [pH 7.4]) was used to remove splenic erythrocytes. The cells were then suspended in RPMI 1640 medium (GIBCO) supplemented with 5% fetal calf serum, HEPES (25 mM; Sigma), l-glutamine (1 mM; BioWhittaker), sodium pyruvate (1 mM; Sigma), and penicillin-streptomycin (1 mM; Sigma) and seeded at 5 × 10⁶ cells per well in triplicate in flat-bottomed 96-well microtiter plates (Costar) in 200 μl of culture medium alone or with various concentrations of TLA or 10 μg of concanavalin A per ml as a positive control of proliferation. The plates were incubated in 5% CO₂ at 37°C for 4 days, and 1 μCi of [³H]thymidine (NEN, Paris, France) was added for the final 18 h of incubation at 37°C. The plates were washed in 5% CO₂ at 37°C. The plates were washed with 1 mg of p-nitrophenylphosphate per ml in diethanolamine buffer (pH 9.8), and the optical density (OD) of each sample was read at 405 nm (1420 multilabel counter Victor; Wallac).

**SAG1-specific antibody responses.** Serum IgG antibody to SAG1 was measured by ELISA. Flat-bottomed 96-well plates (Nunc) were coated with TLA (10 μg/ml) in sodium carbonate buffer overnight. The plates were washed and blocked with PBS–4% bovine serum albumin (BSA). Serial dilutions of sera, in PBS, were added, and the plates were incubated for 1 h at 37°C. The plates were then washed in PBS–0.05% Tween 20 and incubated with alkaline phosphate–conjugated goat anti-mouse IgG (γ-chain specific) diluted 1:1,000 in PBS–4% BSA for 2 h at 37°C. The plates were washed with 1 mg of p-nitrophenylphosphate per ml in diethanolamine buffer (pH 9.8), and the optical density (OD) of each sample was read at 405 nm (1420 multilabel counter Victor; Wallac). The antigen-specific antibody titer was given as the reciprocal of the highest dilution whose absorbance was 2.5-fold greater than the absorbance of the sera of control mice at the same dilution. Results are expressed as the means of log₂ titers ± standard deviations (SD).

The Ig subclasses of the antibodies were determined with the alkaline phosphatase conjugates IgG1, IgG2a, and IgG2b, and IgG3 (1:500; Cappel) and developed as described above. Reactions were stopped when the OD at 405 nm for total IgG was 2 U and compared between groups.

**Assay of ASC by ELISPOT assay.** Nasal lymphocytes were obtained by dissecting out the NALT, identified as two small longitudinal strips of tissue (57). Cell suspensions were obtained as described above. Cells were suspended in culture medium containing 1% supernatant from concanavalin A-stimulated rat spleen cells. B cells secreting Ig specific for the SAG1 protein of *T. gondii* were detected by enzyme-linked immunosorbent (ELISPOT) assay (11). Briefly, a 96-well plate (Costar) was coated overnight at 4°C with TLA at 10 μg/ml in sodium carbonate buffer. The wells were then blocked with 1% BSA for 1 h, and the diluted lymphocytes were added to each well, centrifuged, and incubated for 4 h at 37°C in 5% CO₂. The plate was washed three times in H₂O and three times in PBS–0.05% Tween 20 and incubated overnight at 4°C with alkaline phosphatase–conjugated goat anti-mouse IgG (Sigma), with rinses in TLA and TNP after each step. An alkaline phosphatase conjugate IgG MAb was added to detect IgG.

**Measurement of ASC by ELISPOT assay.** Nasal lymphocytes were obtained by dissecting out the NALT, identified as two small longitudinal strips of tissue (57). Cell suspensions were obtained as described above. Cells were suspended in culture medium containing 1% supernatant from concanavalin A-stimulated rat spleen cells. B cells secreting Ig specific for the SAG1 protein of *T. gondii* were detected by enzyme-linked immunosorbent (ELISPOT) assay (11). Briefly, a 96-well plate (Costar) was coated overnight at 4°C with TLA at 10 μg/ml in sodium carbonate buffer. The wells were then blocked with 1% BSA for 1 h, and the diluted lymphocytes were added to each well, centrifuged, and incubated for 4 h at 37°C in 5% CO₂. The plate was washed three times in H₂O and three times in PBS–0.05% Tween 20 and incubated overnight at 4°C with alkaline phosphatase–conjugated goat anti-mouse IgG (Sigma), with rinses in TLA and TNP after each step. An alkaline phosphatase conjugate IgG MAb was added to detect IgG.

**Challenge infection.** Two weeks after the last immunization, mice were infected orally with 70 cysts of the 76K strain. The mice were killed 1 month after the challenge, and their brains were removed. Each brain was homogenized in 5 ml of PBS containing 1 mM phenylmethylsulfonyl fluoride (Sigma), β-mercaptoethanol (50 μM), and penicillin-streptomycin (1 mM; Sigma) and seeded at 5 × 10⁶ cells per well in triplicate in flat-bottomed 96-well microtiter plates (Costar) in 200 μl of culture medium alone or with various concentrations of TLA or 10 μg of concanavalin A per ml as a positive control of proliferation. The plates were incubated in 5% CO₂ at 37°C for 4 days, and 1 μCi of [³H]thymidine (NEN, Paris, France) was added for the final 18 h of culture. The cells were then harvested on glass fiber filters using an automatic cell harvester (Tomtec; Wallac), and the amounts of [³H]thymidine incorporated into the DNA of proliferating cells were determined in a liquid scintillation β-counter (Microbeta Trilux; Wallac). Proliferation was expressed as the stimulation index (SI) (counts per minute for stimulated cells/counts per minute for stimulated cells).

**Detection of cytokines in cell supernatants.** Cytokines released from spleen and mesenteric lymph node cells stimulated in vitro with 15 μg of TLA per ml were measured in the culture supernatants collected at 48 h for interleukin 2 (IL-2) and IL-4, 72 h for IL-10 and IL-5, and 96 h for IFN-γ detection. Levels of cytokines were determined with a commercial enzyme-linked immunosorbent assay (ELISA) kit (Opteia set: R&D Systems). In brief, 96-well (flat-bottomed) plates (Nunc) were coated with anti-mouse cytokine in PBS overnight at room temperature. Free sites were then blocked with block buffer, and supernatants (1/2 diluted) were added and incubated for 2 h at room temperature. A standard curve was also created with the recombiant cytokine. Cytokine was detected using biotinylated anti-mouse cytokine. Horseradish peroxidase–streptavidin was added and incubated for 20 min. Bound antibodies were visualized with a tetramethylbenzidine substrate (Sigma). The enzymatic reaction was stopped with 2 N sulfuric acid, and absorbances were read at 450 nm (1420 multilabel counter Victor; Wallac).

**RESULTS**

**Proliferative activity after intranasal immunization using LTR72 as the adjuvant.** Mice were primed with two intranasal immunizations with SAG1 protein mixed with the LTR72,
Spleens and mesenteric lymph nodes were removed from three mice of all immunized groups 15 days after the last immunization and prepared as single-cell suspensions. They were then stimulated with various concentrations of TLA in vitro. Cells from untreated mice or mice immunized with SAG1 alone or with toxin alone were also prepared similarly and cultured in parallel. Splenocytes from all groups of mice immunized intranasally with LTR72, LT, or CT combined with SAG1 responded by increased dose-dependent antigen proliferation in vitro. In contrast, cells from mice immunized with LTK63 plus SAG1 and from control mice did not respond to antigen restimulation (Fig. 1A). The SI for LTR72 plus SAG1 (SI = 6.3) was significantly higher than that for CT plus SAG1 (SI = 4) but lower than that for LT plus SAG1 (SI = 13.5) (Fig. 1) (P < 0.001). Splenocytes cultured in the absence of specific antigen showed no enhanced cell proliferation.

The cellular response in mesenteric lymph nodes was also analyzed. Mesenteric lymph node cells from animals immunized with LTR72 plus SAG1, LT plus SAG1, and CT plus SAG1 (Fig. 1B) were also stimulated dose dependently. The SI for mice immunized with LT plus SAG1 (SI = 11.2) was higher than the SI for mice immunized with LTR72 plus SAG1 (SI = 6.8) or with CT plus SAG1 (SI = 6) (P < 0.01 and P < 0.001, respectively). The mesenteric lymph node cells from mice immunized with LTK63 plus SAG1 did not respond to antigen restimulation. These results demonstrate that intranasal administration of a mixture of SAG1 and LTR72 triggered systemic and mucosal cellular immunity. LTK63, which is devoid of any toxic and enzymatic activity, does not trigger a detect-
plus SAG1 or to that with CT plus SAG1 and splenocytes and mesenteric lymph node cells, unstimulated further explore the immune response. Culture supernatants of patterns obtained after cellular stimulation were studied to three booster immunizations. Itation in our model, and it may be necessary to perform two to activity might account for its poor stimulation of cell prolifer-
able cellular response. The absence of ADP ribosyltransferase activity might account for its poor stimulation of cell prolifer-
able in our model, and it may be necessary to perform two to three booster immunizations.

**Analysis of culture supernatants for cytokines.** The cytokine patterns obtained after cellular stimulation were studied to further explore the immune response. Culture supernatants of splenocytes and mesenteric lymph node cells, unstimulated and stimulated with 15 \( \mu \text{g} \) of TLA per ml, were assayed for cytokines. Only groups immunized with SAG1 plus the toxins, except LTK63, produced significant amounts of both IFN-\( \gamma \) and IL-2 compared to levels produced by the control groups (Table 1). Mesenteric lymph node cells from mice immunized with LTR72, LT, or CT with SAG1 (Table 1) produced only IL-2. IL-4, IL-5, and IL-10 were undetectable in any of the supernatants from spleen and mesenteric lymph node lymphocytes stimulated with TLA in vitro (data not shown).

**Serum antibody responses.** Mice given two 1-\( \mu \text{g} \) doses of toxin with 10 \( \mu \text{g} \) of SAG1 protein produced a specific IgG antibody response to SAG1 when they were immunized with SAG1 plus LTR72, LTK63, LT, or CT (Fig. 2A) after the first immunization. The second immunization enhanced the antibody response. No antibody was detected in the control group. The titers of anti-SAG1 IgG were high in all groups of mice immunized with one of the toxins plus SAG1. However, the antibody titer in the group given LTR72 plus SAG1 was significantly lower (\( P < 0.05 \) compared to the titer with LT plus SAG1). The IgG isotype pattern of the SAG1-specific antibodies elicited after two immunizations was analyzed. Both IgG2a and IgG2b isotypes were detected in all groups immunized with toxin and SAG1 (Fig. 2B). No IgG1 or IgG3 was detected in significant quantities in any group.

**IgA in nasal, lung, and intestinal washes.** The local humor-
al IgA response induced by intranasal immunization with the nontoxic mutants LTR72 and LTK63 was analysed using nasal, lung, and intestinal washes taken after the second immunization. IgA responses were visualized by Western blotting on a nitrocellulose membrane of TLA. Bands corresponding to SAG1 protein (30 kDa) were observed in samples from the nasal, lung, and intestinal washes (Fig. 3) of mice immunized with SAG1 plus wild-type CT and LT showed that they also trigger an IgA response. Control mice and mice immunized with SAG1 alone or toxin

![Image](http://iai.asm.org/)
alone never showed any nasal, lung, or intestinal antibody responses.

**IgG and IgA ASC in nasal mucosa.** The numbers of IgG and IgA ASC specific for SAG1 were determined in nasal mucosal tissues after the second immunization. Intranasal immunization always induced IgA-specific ASC in all mice immunized with SAG1 plus any of the toxins. The numbers of ASC in the nasal mucosa of mice immunized with LTR72 plus SAG1 and CT plus SAG1 were similar and lower than in mice immunized with LT plus SAG1 ($P<0.01$ compared to numbers with LTR72 plus SAG1; $P<0.05$ compared to numbers with CT plus SAG1) (Table 2). Mice immunized with LTK63 plus SAG1 had only a few IgA ASC. Mice immunized with SAG1 plus native toxins were the only mice with IgG ASC in the nasal mucosa. There were significantly more ASC in mice immunized with LT plus SAG1 than in mice immunized with CT plus SAG1 ($P<0.01$).

**Protection.** The protection provided by vaccination with these nontoxic mutant LTs, LTR72 and LTK63, to act as mucosal adjuvants. They were mixed with SAG1 protein and used for intranasal immunization of CBA/J mice. They were tested by assessing the protective immune response to a challenge with *T. gondii*. The protection against *T. gondii* was measured by the brain cyst load. Mice immunized with LTR72 plus SAG1 or LTK63 plus SAG1 had significantly fewer cysts than controls (Fig. 4). This decrease was as great as with the wild-type toxin and the homolog CT, as previously reported (14). The ADP-ribosylating holotoxins CT and LT are powerful adjuvants. Initially, the A subunit was considered to be crucial for adjuvant activity (40), but the construction of nontoxic mutants devoid of any enzymatic or toxic activity showed that these mutants still had their adjuvant and immunological properties (15, 17, 18, 25).

**DISCUSSION**

We have investigated the capacities of two nontoxic mutant LTs, LTR72 and LTK63, to act as mucosal adjuvants. They were mixed with SAG1 protein and used for intranasal immunization of CBA/J mice. They were tested by assessing the protective immune response to a challenge with *T. gondii*. The protection against *T. gondii* was measured by the brain cyst load. Mice immunized with LTR72 plus SAG1 or LTK63 plus SAG1 had significantly fewer cysts than controls (Fig. 4). This decrease was as great as with the wild-type toxin and the homolog CT, as previously reported (14). The ADP-ribosylating holotoxins CT and LT are powerful adjuvants. Initially, the A subunit was considered to be crucial for adjuvant activity (40), but the construction of nontoxic mutants devoid of any enzymatic or toxic activity showed that these mutants still had their adjuvant and immunological properties (15, 17, 18, 25).

**TABLE 2. ASC specific for SAG1 in the nasal mucosa**

<table>
<thead>
<tr>
<th>Group</th>
<th>No. of ASC/10^6 cells ± SD secreting:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IgA</td>
</tr>
<tr>
<td>Untreated</td>
<td>0</td>
</tr>
<tr>
<td>SAG1</td>
<td>0</td>
</tr>
<tr>
<td>LTR72</td>
<td>0</td>
</tr>
<tr>
<td>LTR72 + SAG1</td>
<td>$142.6 \pm 71.6$</td>
</tr>
<tr>
<td>LTK63</td>
<td>0</td>
</tr>
<tr>
<td>LTK63 + SAG1</td>
<td>$13 \pm 3.6$</td>
</tr>
<tr>
<td>LT</td>
<td>0</td>
</tr>
<tr>
<td>LT + SAG1</td>
<td>$270 \pm 73$</td>
</tr>
<tr>
<td>CT</td>
<td>0</td>
</tr>
<tr>
<td>CT + SAG1</td>
<td>$158 \pm 23$</td>
</tr>
</tbody>
</table>

*Fifteen days after the last immunization, the nasal mucosa was removed and ELISPOT analysis was performed for detecting IgA and IgG ASC. Results from one of two similar experiments are shown.*
ably through the production of IFN-γ. CD8 T lymphocytes help mediate resistance to toxoplasmosis (7–9, 38). Their protective capacity depends on epithelial lymphocytes, which form the first functional barrier to the site of natural infection by the parasite. The cells that are important in the gut during infection by T. gondii are intraepithelial lymphocytes, which form the first functional barrier to toxoplasmosis (7–9, 38). Their protective capacity depends on the production of IFN-γ (7). CD4 and CD8 T lymphocytes are also implicated. Experiments with T cells from the spleen (46) and with mice depleted of T cells (23) indicate that CD4 and CD8 T lymphocytes help mediate resistance to T. gondii, probably through the production of IFN-γ.

Several reports indicate that LTR72 elicits a Th2-like response (16, 52). In contrast, the cellular response in our experiments had more of a Th1 pattern in terms of cytokines produced. We detected, by ELISA, IFN-γ and IL-2 in culture supernatants of restimulated cells from mesenteric lymph nodes and splenocytes taken from mice immunized with LTR72 plus SAG1. A response with a Th1 cytokine pattern was also obtained with LT or CT plus SAG1. This observation can be supported by the fact that IFN-γ is an important cytokine in the immune response, conferring resistance to the development of toxoplasmic encephalitis (54–56), and that IFN-γ-mediated cellular immunity is required for the survival of mice in acute and chronic stages of infection with T. gondii (24, 26). Last, T. gondii-specific Th1 cells activate, via IFN-γ production, infected macrophages to kill intracellular parasites (51). B cells are also crucial in the resistance of the host cell to T. gondii. An experiment with mice lacking B cells showed decreased resistance to infection, indicating that antibody production by B cells prevents the persistent replication of tachyzoites in the brain and lung (32). It has also been reported that antibody can inhibit intracellular proliferation (44) and that antibody-coated tachyzoites are killed by macrophages in vitro (1). In vitro, monoclonal antibodies to SAG1 can inhibit murine enterocyte infection (45). The protection we obtained was correlated with high titers of anti-SAG1 IgG in serum. The specific IgGs produced were IgG2a and IgG2b subclasses (Fig. 2B), as previously observed with CT (14). The IgG2a subclass was specific for a Th1 response. IgG2b production is known to be selectively induced by transforming growth factor β (TGF-β) (53), and this TGF-β can also direct the switching of B cells to the IgA isotype (19).

Intranasal immunization is more effective than intragastric immunization, as it generates an earlier and stronger mucosal immune response (27, 64). Intranasal immunization also delivers the antigen directly to the site of uptake (37). The stim-
ulation of cells from the spleen and mesenteric lymph nodes after nasal immunization points to the existence of a common mucosal immune system. This implies that cells stimulated in the NALT by the antigen presented by antigen-presenting cells can leave this site for mucosal effector sites (36, 42, 50). Nasal immunization triggers pulmonary immunity. Antigen administered by the nasal route can reach the tracheal area, or dendritic cells loaded with the antigen may have migrated from the NALT to the pulmonary lymph node, where they can initiate an immune response (58). Intranasal immunization triggers both mucosal and systemic T- and B-cell responses (64) and can be used to target pathogens that invade far from the immunization site, such as the gut (10, 14).

The response had a Th1 cytokine pattern and IgG subclass profile protective against a challenge with the parasite. The most important finding is a protective immunity obtained without a detectable cellular response by using the association of SAG1 and LTK63. LTR72 in association with SAG1 triggers strong cellular and humoral responses to the protein SAG1. These mutant LTs could thus be attractive mucosal adjuvants to obtain immune responses at systemic and mucosal sites following vaccination against T. gondii. The response had a Th1 cytokine pattern and IgG subclass profile protective against T. gondii infection. Nevertheless, LTK63, which is devoid of toxicity and which produced only a humoral response under our experimental conditions, is as good an adjuvant as LTR72 to promote protection, but more has to be known of the exact immune mechanisms involved in protection. The importance of a humoral or IFN-γ cytokine must be studied in our future work with deficient mice.

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