Invasive amebiasis, due to the enteric protozoan Entamoeba histolytica, manifests most commonly as amebic colitis or a liver abscess. E. histolytica infection is highly endemic in the most populated areas of the world, such as India, China, Africa, the Middle East, South and Central America, and southeast Asia (35). Recent epidemiological studies in Hue, Vietnam (8), South Africa (1, 36), and Bangladesh (20) revealed that E. histolytica infection and invasive disease are more prevalent than previously appreciated. Up to 70% of adults in an area where infection is highly endemic, such as Durban, South Africa, are challenged by the parasite each year (1); the prevalence of amebic liver abscesses (ALA) is as high as 21 per 100,000 people per year in Vietnam (8). It is estimated that E. histolytica is the third leading parasitic cause of death worldwide (43).

Presently, no vaccine to prevent asymptomatic E. histolytica infection or invasive disease is available. E. histolytica trophozoites attach to colonic mucins, host inflammatory cells, and epithelial cells by a galactose-inhibitable surface lectin (10, 32, 37). The lectin consists of three subunits (16); the heavy subunit contains the galactose-binding activity (33) and is highly antigenic (38). Purified native lectin protein (33) and a cysteine-rich recombinant protein encompassing amino acids 758 to 1134 of the lectin heavy subunit (designated LC3) (40) are effective as a vaccine (in combination with Freund’s adjuvant) in the gerbil model of ALA. Monoclonal antibodies to the lectin heavy subunit inhibit amebic binding to colonic mucins (10), suggesting that an intestinal antilectin immunoglobulin A (IgA) antibody response may be effective in preventing parasite attachment to and invasion of the gut. Studies of asymptomatic E. histolytica intestinal infection in children in Bangladesh (19–21) and in adults in Durban, South Africa (1, 36),

Adherence-Inhibitory Intestinal Immunoglobulin A Antibody Response in Baboons Elicited by Use of a Synthetic Intranasal Lectin-Based Amebiasis Subunit Vaccine

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We designed an amebiasis subunit vaccine that is constructed by using four peptide epitopes of the galactose-inhibitable lectin heavy subunit that were recognized by intestinal secretory immunoglobulin A (IgA) antibodies from immune human subjects. These epitopes are contained in the region encompassing amino acids 758 to 1134 of the lectin heavy subunit, designated LC3. Baboons (Papio anubis) are natural hosts for Entamoeba histolytica; naturally infected baboons raised in captivity possess serum IgA antibodies to the same four LC3 epitopes as humans. Uninfected, seronegative baboons received four intranasal immunizations at 7-day intervals with the synthetic peptide vaccine (400, 800, or 1,600 μg per nostril) with cholera toxin (20 μg) as the adjuvant. As determined by an enzyme-linked immunosorbent assay (ELISA), each dose of the peptide vaccine elicited antipeptide serum IgA and IgG and intestinal IgA antibody responses in all six immunized baboons by day 28, 7 days after the last immunization (P < 0.01 for each dose compared to the cholera toxin control). The peptide vaccine elicited serum IgG and intestinal IgA antibodies that recognized purified recombinant LC3 protein (P < 0.008 and 0.02, respectively) and native lectin protein (P < 0.01). In addition, an indirect immunofluorescence assay with whole trophozoites (P < 0.01) and Western blot analysis confirmed that serum IgG antibodies from vaccinated baboons recognized native lectin protein on the surfaces of axenic E. histolytica trophozoites or from solubilized amebae. All four synthetic peptides were immunogenic; the vaccine elicited dose- and time-dependent responses, as determined by ELISA optical density readings indicating the production of serum and intestinal antibodies (P < 0.02 for antipeptide and antilectin antibodies). As a positive control, intranasal immunization with purified recombinant LC3 protein with cholera toxin as the adjuvant elicited a serum anti-LC3 IgA and IgG antibody response (P = 0.05 and <0.0001, respectively); however, no intestinal anti-LC3 IgA antibody response was observed (P = 0.4). Of interest, serum IgA and IgG antibodies elicited by the recombinant LC3 vaccine did not recognize any of the four putatively protective LC3 peptide epitopes. Both serum and fecal antibodies elicited by the peptide vaccine exhibited neutralizing activity, as determined by their dose-dependent inhibition of the galactose-specific adherence of E. histolytica trophozoites to Chinese hamster ovary cells in vitro (P < 0.001 for each group of antibodies compared to the control). In summary, a lectin-based intranasal polysine-linked synthetic peptide vaccine was effective in eliciting an adherence-inhibitory, intestinal antilectin IgA antibody response in baboons. Future studies with the baboon model will determine vaccine efficacy against asymptomatic E. histolytica intestinal infection.

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demonstrated that intestinal antilectin IgA antibodies mediate the rapid clearance of infection or immunity to new infections. Studies in Durban, South Africa, by our research group (1, 36) revealed that subjects cured of ALA develop high-titer peaks of intestinal antilectin IgA antibodies upon parasite challenge and that these antibody levels are sufficient to prevent or clear asymptomatic *E. dispar* and *E. histolytica* infections. In contrast, control subjects without a history of treatment for invasive amebiasis have lower-titer intestinal antilectin IgA antibody peaks that are delayed, less long-lasting, and insufficient to clear amebic infection (1).

Our strategy for vaccine design was to use synthetic peptides to elicit a mucosal immune response that mimics that of the South African ALA subjects. By using segments of the LC3 protein, expressed as overlapping recombinant proteins, we identified two LC3 fragments that are recognized by the antilectin IgA antibodies present in the sera and feces of immune human hosts (2). Fine mapping using overlapping synthetic peptides led to the discovery of four discrete LC3 peptide epitopes, referred to as peptides 2, 9, 11, and 12. Further studies have demonstrated that LC3 epitope recognition by human IgA antibodies is highly conserved and was exhibited by serum IgA antibodies from subjects cured of colitis in Cairo, Egypt (our unpublished data). Based on the epitope-mapping studies, we constructed a synthetic peptide vaccine containing the four putatively protective LC3 epitopes (peptides 2, 9, 11, and 12).

The systemic immunization of BALB/c mice with purified recombinant LC3 protein elicits murine IgA antibodies with LC3 epitope specificity different from that of human IgA antibodies, with little overlap (2). In addition, most mouse strains exhibit high levels of innate immunity to *E. histolytica* infection (22). Baboons were chosen for this vaccine study as they are naturally infected with *E. histolytica* in the wild (25, 42) and in captivity (45); in addition, baboons have an immune system that best approximates that of humans (39). Cholera toxin (CT) was selected as the adjuvant because of its known effectiveness in inducing a mucosal immune response (30) and because of our prior experience using recombinant LC3 protein to induce the secretion of intestinal adherence-inhibitory, antilectin IgA antibodies in BALB/c mice following oral immunization (7). Intranasal vaccination requires a smaller immunization volume of bovine serum albumin (BSA [Pierce; catalog no. 23210]) at 200 μg/dl was diluted with 500 μl of H2O and kept at 4°C. The DNA sample to be tested (5 μl) was added to 95 μl of the PCR mixture to yield a complete volume of 100 μl. Each DNA sample was tested twice, once using the *E. histolytica* sense primer and once using the *E. dispers* sense primer.

Conventional PCR machine thermocycling conditions were as described previously (2). The specific detection of amplified DNA was achieved by gel electrophoresis. Amplified DNA was separated on a 2% agarose gel containing ethidium bromide.

**ELISA for detection of baboon antilectin, anti-LC3, and antipeptidet antibod-ies.** ELISAs were performed as described previously (36). Recombinant 52-kDa LC3 protein was purified as described previously (40), and native lectin protein was purified by immunoaffinity chromatography from detergent-solubilized trophozoites (33, 40). ELISA 96-well microtiter flat-bottom polystyrene plates were coated with LC3 protein, purified native lectin, or synthetic peptides, and nonreactive sites were blocked with 1% BSA. Serum samples at a 1:100 dilution in phosphate-buffered saline (PBS)-TWEEN (1% BSA) were analyzed for IgG or IgA and incubated for 2 h at room temperature or overnight at 4°C. Each sample was studied in triplicate and tested twice by ELISAs. Alkaline phosphatase-conjugated goat anti-human IgG (Sigma, St. Louis, MO) or IgA (ICN Biomedicals, Costa Mesa, CA) antibodies were diluted (1:5,000 for IgG and 1:2,500 for IgA) in PBS-TWEEN (1% BSA) to give a final volume of 100 μl/well, and the plates were incubated for 2 h at room temperature. Reading of the plates and correction for nonspecific background binding were performed as described previously (38).

ELISAs for the detection of fecal antilectin IgA antibodies were performed using purified native lectin protein as described previously (1, 3). We have previously reported that native lectin is more sensitive than recombinant LC3 protein for the detection of antilectin intestinal IgA antibodies in humans (3, 36). Flat-bottomed microtiter ELISA plates were coated with lectin protein (0.1 μg/well), and nonreactive sites were blocked with 1% BSA (3). Fecal samples were mixed with equal volumes of PBS containing 2.0 mM phenylmethylsulfonyl fluoride, and the mixtures were added to 100 μl/well and incubated for 2 h at room temperature or overnight at 4°C. Alkaline phosphatase-conjugated goat anti-human IgA antibodies (Sigma, St. Louis, MO) diluted (1:5,000) in PBS-TWEEN containing 1% BSA were added for incubation over 2 h at room temperature. Plate reading with correction for nonspecific background binding was performed as described previously (3).
9-fluorenlymethoxy carbonyl chemistry. Peptides were cleaved from the resin, deprotected using reagent R, and then lyophilized. Lyophilized crude peptides were purified by preparative reverse-phase high-performance liquid chromatography using a Beckman 126 system with a Vydac C4 column. Solvent A consisted of 0.1% trifluoroacetic acid in water, and solvent B consisted of 0.1% trifluoroacetic acid in acetonitrile. The purification was carried out with a gradient of 0 to 60% solvent B in 30 min. Purity and quality control of the peptides was done by using an analytical-high performance liquid chromatography HP 1090 system with a C18 column and the same gradient and by using mass spectrometry as on an HP matrix-assisted laser desorption ionization-time of flight system (6). The levels of purity of the four peptides utilized were 97, 100, 90, and 75%, respectively, for peptides 2, 9, 11, and 12.

Peptide polysyme complexes were manufactured by Alpha Diagnostic international and shipped in a lyophilized form. Each polysyme molecule had a set of six peptides, a random mixture of all four peptides, linked to it. The polysyme-linked peptide vaccine preparation was suspended in saline, submitted to a quantitative assay (12), and adjusted with saline according to the desired vaccinon protocol. Each aliquot tube contained one vaccine dose for one animal to avoid repeated thawing and freezing of peptides. The contents of each tube were mixed with the proper amount of adjuvant as defined by the vaccination protocol before use.

Experimental vaccine protocol. Seronegative and PCR-negative baboons were selected by screening, isolated, and then immunized intranasally (in each nostril) four times at 7-day intervals. CT adjuvant (Sigma) at 20 µg per nostril served as a negative control, and purified recombinant LC3 protein (40) at 200 µg per nostril with CT adjuvant was the positive control. An analysis of dose-dependent responses to the polysyme-linked synthetic peptide vaccine at 400, 800, and 1,600 µg with 200 µg of CT per nostril was performed. The definition of a positive response was the recognition of amebic surface lectin by serum IgG antibodies from peptide-vaccinated sera (1/4, 1/8, 1/16, 1/32, 1/64, and 1/128) from peptide-vaccinated and control baboons. Therefore, a total of six baboons per experimental condition were studied. Experiments 1 and 2 included three baboons immunized with adjuvant alone, three baboons immunized with LC3 protein, and three baboons immunized with 400 µg of the peptide vaccine. Experiments 3 and 4 included three baboons immunized with 800 µg and three immunized with 1,600 µg of the peptide vaccine. Therefore, a total of six baboons per experimental condition were studied. Blood and feces were collected from each baboon before vaccination and 7 days after each vaccination (days 7, 14, 21, and 28 of the study). All sera (1/4, 1/8, 1/16, 1/32, 1/64, and 1/128) from peptide-vaccinated and control baboons were prepared in 2% Marvel (dried skim milk) dissolved in PBSM and added at 25 µl/well. Slides were incubated for 2 h in a moist chamber at room temperature. Excess antibodies were rinsed off with PBSM, and the slides were washed. After drying, fluorescein isothiocyanate-labeled goat anti-human IgG antibodies (Sigma) diluted 1:40 in PBSM (containing 33 µl of 1% Evan’s blue/ml) were added, and the slides were incubated for 1 h. After washing, a drop of mounting medium (50% Citifluor and 50% PBS) was added to each well. Fluorescence density (FD) readings were obtained with an immunofluorescence microscope (E clips E 600; Nikon) under blue light; a Photometrics COOL SNAP ES camera was used to read the slides. Each trophozoite was traced separately, and an FD histogram was developed and corrected for background fluorescence. The cutoff point for immunofluorescence positivity was calculated as the mean + 2 standard deviations of the FD reading for control trophozoites submitted to the same procedure.

Effect of antipeptide serum and intestinal antibodies from vaccinated baboons on amebic galactose-inhibitable adherence in vitro. E. histolytica trophozoites, strain HMI:IMSS, were grown in F-12 medium (GIBCO) supplemented with 10% fetal bovine serum (GIBCO), penicillin (100 µg/ml), and streptomycin (100 µg/ml) as described previously (37). Amebae galactose-inhibitable adherence studies were performed by using a rosetting assay as described previously (37). Amebic E. histolytica trophozoites (10⁸/ml) were incubated at 4°C in serial dilutions (1.25, 2.5, 5.0, and 10%) of baboon sera or fecal supernates. Sera and feces were obtained from baboons 7 days after the final vaccination with the 1,600-µg dose of the peptide vaccine. Amebae incubated with galactose (0.05, 0.1, 0.2, and 0.4% dilutions) in RPMI-10% fetal bovine serum were used as a positive control for the adherence-inhibitory effects of galactose. Amebae in RPMI-10% fetal bovine serum only or with sera or fecal supernates from naive, uninfected seronegative baboons served as negative controls. Trophozoites were washed, and 10⁶ were added to CHO cells (2 × 10⁴) in 1 ml of M1998. The mixture was centrifuged at 250 × g and incubated for 2 h at 4°C. After incubation, 0.8 ml of supernate was removed and the cellular pellet was suspended. The total number of amebae that formed or did not form rosettes with CHO cells (three or more adherent CHO cells per ameba) was determined in a hemocytometer chamber. The results were expressed as the percentage of all the amebae counted that formed ameba-CHO cell rosettes.

Statistical methods. We used repeated-measure models to analyze the mean change in antibody response compared to the baseline, as determined by ELISA, induced by vaccination with one of three different doses of synthetic peptides or with purified LC3 recombinant protein. The effects of each vaccination regimen both over the entire follow-up period, as assessed with a type 3 F-test, and at the last follow-up, as assessed with a two-sample t test, were compared to those of a saline control. Degrees of freedom for the statistical tests were determined using the within-between method. An additional repeated-measure analysis using data for all three doses of synthetic peptides was used to perform a type 3 F-test of dose-dependent responses. The analyses described above were repeated for the nine different combinations of antigen (synthetic peptides, recombinant LC3, and native lectin) and antibodies (IgA and IgG in sera and IgA in feces) we investigated.

Our repeated-measure models accounted for correlated errors that may have arisen due to the longitudinal measurements made for each baboon at the baseline time point and at 7, 14, 21, and 28 days after the baseline time point. We used a first-order autoregressive model for the variation in the immune response of each baboon. Since immune responses to experimental vaccines were expected to be more variable than those to the saline control, our analysis estimates separate residual variance parameters for synthetic peptide, LC3, and saline vaccination groups. The models were fit to the data by restricted maximum likelihood using PROC MIXED in SAS version 9.1.

The proportions of trophozoites adhering to CHO cells under different experimental conditions were compared using chi-square tests. For all statistical analyses, P < 0.05 was considered statistically significant.

RESULTS

We screened a total of 131 baboons from the colony and found that 16% were positive for serum antilectin IgG antibodies, 6.5% were positive for serum antilectin IgA antibodies, and 9.9% were positive for intestinal antilectin IgA antibodies. None of the baboons screened by fecal PCR were infected with E. histolytica; however, 22% were infected with E. dispar. We found that antilectin intestinal IgA antibodies from baboons naturally infected in captivity recognized all of the LC3 peptide epitopes included in the vaccine (optical density [OD] readings were 2.76, 2.08, 0.393, and 0.494 for LC3 peptides 2, 9, 11, and 12, respectively). Baboons selected for the study were seronegative and were not infected with E. histolytica or E. dispar.

Antibody responses to the synthetic peptide and recombinant LC3 vaccines were examined by three separate ELISAs using either all four synthetic peptides, purified recombinant LC3 protein, or immunoaffinity chromatography-purified native lectin (obtained from solubilized trophozoites) as the capture antigen. We found that baboons immunized intranasally with the synthetic peptide vaccine demonstrated brisk serum antipeptide IgA, serum antipeptide IgG, and intestinal antipeptide IgA antibody responses in a dose-dependent fashion (P < 0.03 for each type of response compared to those in control baboons) (Fig. 1A, B, and C, respectively). Higher doses of the peptide vaccine (800 and 1,600 µg) elicited a more rapid (P < 0.02) and incrementally greater antibody response than the 400-µg dose (P < 0.003 for each antibody studied).
Of interest, by day 28, 100% of the six baboons studied for each peptide vaccine dose (a total of 18 baboons) developed antipeptide serum IgA and serum IgG antibody responses \((P < 0.01)\) and at day 28 \((P < 0.01)\). The serum antipeptide IgA and IgG antibody responses increased with the vaccine dose \((P < 0.0001)\). The results of vaccination with purified recombinant LC3 protein are also illustrated in Fig. 1. Unexpectedly, the serum IgA and IgG antibodies elicited by the LC3 vaccine did not recognize the synthetic peptide antigen (Fig. 1). Using recombinant LC3 protein as the capture antigen in the ELISA, we found that the peptide vaccine elicited serum IgG antibodies to LC3 protein \((P < 0.008)\) (Fig. 2B); antibody responses were maximal at 28 days and generally dose dependent \((P < 0.001)\) compared to the baseline) (Fig. 2B). Serum IgA antibodies elicited by the peptide vaccine had low levels of reactivity to LC3 protein (Fig. 2A). Intestinal IgA antibodies to LC3 protein were elicited

![Fig. 1. Effects of immunization of baboons with the polylysine-linked synthetic peptide vaccine (400, 800, or 1,600 μg), the recombinant LC3 protein vaccine (200 μg), or a control vaccine consisting of CT adjuvant (20 μg) alone as assessed by an ELISA using the synthetic peptides from the vaccine as the capture antigen. Compared to controls, immunized baboons exhibited serum antipeptide IgA (A) and IgG (B) responses over 4 weeks \((P < 0.03)\) and at day 28 \((P < 0.01)\). The serum antipeptide IgA and IgG antibody responses increased with the vaccine dose \((P < 0.0001)\). (C) The synthetic peptide vaccine antigen at the 800- and 1,600-μg doses \((P < 0.02)\) induced an intestinal IgA antibody response to the peptide as measured over 4 weeks. However, at day 28, positive responses to all doses were observed \((P < 0.02)\) compared to the control), and dose-dependent responses were observed over the 4 weeks \((P = 0.004)\). The LC3 vaccine did not elicit any serum IgA (A) or IgG (B) antibodies to the peptide antigen \((P = 0.80)\) compared to the control. The LC3 vaccine did not induce any intestinal antipeptide IgA antibody response \((P = 0.46)\) (C).](http://iai.asm.org/figure/1.png)
after the fourth immunization with the 400- and 1,600-μg peptide doses (P < 0.02) (Fig. 2C). The recombinant LC3 vaccine induced high ODs indicating high levels of serum IgA and IgG antibodies to itself (P, <0.001 compared to controls) (Fig. 2A and B). However, no measurable intestinal IgA antibody response was elicited by the LC3 vaccine, even at 28 days (Fig. 2C) (P, <0.001 compared to the peptide vaccine and >0.5 compared to saline controls).

Lastly, we studied antibody responses using immunoaffinity chromatography-purified native lectin as the capture antigen in the ELISA. Serum IgA and IgG antibodies elicited by the peptide vaccine reacted with immunoaffinity chromatography-purified lectin, but at low levels as indicated by ODs (Fig. 3). However, the synthetic peptide at two doses (400 and 1,600 μg) elicited intestinal IgA antibodies in a dose-dependent manner (P = 0.03), and after four vaccinations, these antibodies demonstrated high levels of recognition of purified native lectin protein (P, <0.01 compared to those from control baboons at 28 days) (Fig. 3C). Lastly, vaccination with the recombinant LC3 protein elicited serum IgA and IgG antibodies to immu-

FIG. 2. Effects of immunization of baboons with the polylysine-linked synthetic peptide vaccine (400, 800, or 1,600 μg), the recombinant LC3 protein vaccine (200 μg), or the CT adjuvant (20 μg) alone as assessed by an ELISA using purified recombinant LC3 protein as the capture antigen. The peptide vaccine did not elicit a significant serum IgA antibody response (A) to LC3 antigen at any dose, over time or at 4 weeks (P > 0.233), but did elicit a substantial serum anti-LC3 IgG antibody response (B) at all doses (P < 0.001) and in a dose-dependent manner (P < 0.0001). The peptide vaccine at the 400- and 1,600-μg doses elicited an intestinal anti-LC3 IgA antibody response (C) after 4 weeks (P, 0.02 and 0.06 compared to the control, respectively). The LC3 vaccine elicited significant levels of serum anti-LC3 IgA (A) and IgG (B) antibodies over time and at 4 weeks (P, <0.001 compared to the control). The LC3 vaccine did not elicit any intestinal IgA antibody response (C) over 4 weeks or by the end of the study (P = 0.4).
no affinity chromatography-purified lectin protein at high levels as indicated by OD readings \((P < 0.001\) compared to the control\) (Fig. 3A and B). However, no significant intestinal antilectin IgA antibody response was observed at 28 days after four intranasal vaccinations with LC3 protein (Fig. 3C) \((P = 0.80)\).

To determine whether the low level of recognition in the ELISA of immunoaffinity chromatography-purified lectin protein by peptide vaccine-elicited serum IgA and IgG antibodies compared to recombinant LC3 protein-elicited antibodies may have been due to the alteration of the protein epitopes during the detergent solubilization and protein purification process, we performed IFAs using intact axenic \(E.\ histolytica\) trophozoites. Serum IgG antibodies from peptide (1,600-µg-dose)-immunized baboons were reactive to trophozoite surface antigen at serial serum dilutions from 1/8 to 1/128 as evaluated by an IFA \((P = 0.002)\) for each dilution compared to sera from control baboons (Fig. 4). In addition, every baboon immunized with the peptide vaccine demonstrated a positive antilectin

![Graphs A, B, and C showing ELISA results for different doses of the vaccine.](http://iai.asm.org/)
antibody response as determined by the IFA ($P > 0.001$ for each dilution compared to control baboon sera [Δ]). (Means are represented by solid and dotted lines, respectively.) In addition, serial dilutions of immune baboon sera demonstrated a stepwise decrease in IFA-measured fluorescence intensity ($P < 0.01$ for 1/8 to 1/16 and 1/16 to 1/32). Importantly, 100% of study baboons receiving this vaccine dose had positive serum responses as determined by an IFA ($P < 0.001$ compared to control baboons). Control baboon sera did not show a positive IFA result at any dilution compared to results from IFA studies without serum present (data not shown).

FIG. 5. Western blot analysis of responses of human and baboon control and immune sera to whole trophozoite protein and immunoaffinity chromatography-purified lectin. Lanes: 1, molecular mass standard; 2, whole amebic lysate; 3, immunoaffinity chromatography-purified lectin protein. Serum IgG antibodies from subjects cured of ALA recognized the lectin 170-kDa heavy subunit in both whole trophozoites (panel C, lane 2) and immunoaffinity chromatography-purified native lectin (panel C, lane 3) compared to control human sera (A). In contrast, serum IgG antibodies from peptide-immunized baboons recognized the 170-kDa lectin protein in whole trophozoites (panel D, lane 2) but under these experimental conditions were not reactive with immunoaffinity chromatography-purified native lectin (panel D, lane 3). Control baboon sera were entirely nonreactive (B).

adherence. A positive control with galactose and a negative control with test medium combined with sera or fecal supernates from control baboons (receiving adjuvant only) were studied. As expected, galactose exhibited a dose-dependent (0.05 to 0.4%) inhibition of trophozoite adherence to CHO.
cells compared to control media ($P < 0.001$ for each dilution) (Fig. 6). At each dilution, sera from peptide-immunized baboons inhibited trophozoite adherence to CHO cells compared to either control media ($P > 0.0001$) (Fig. 6) or control baboon sera ($P < 0.001$ for each dilution) (Fig. 6A). Increasing concentrations of immune sera, from 1.25 to 10%, demonstrated a stepwise enhancement of inhibition of amebic adherence to CHO cells ($P < 0.03$ for each dilution compared to the previous one) (Fig. 6A). Fecal supernates from baboons receiving adjuvant alone had some inhibitory effect on amebic adherence to CHO cells ($P < 0.01$) (Fig. 6B), but increasing concentrations of the supernates caused no change in the level of inhibition ($P > 0.5$) (Fig. 6B). Fecal supernates from peptide-immunized baboons exhibited greater inhibition of trophozoite adherence to CHO cells than supernate from control baboons ($P < 0.001$ at each dilution) (Fig. 6B). The inhibition of amebic adherence by fecal supernates from peptide-immunized baboons was also concentration dependent ($P < 0.0004$) (Fig. 6B). The results of these studies demonstrate that the synthetic peptide vaccine elicited serum and intestinal antibodies that recognized lectin molecules on the surfaces of viable *E. histolytica* trophozoites and that had substantial adherence-inhibitory activity in vitro, comparable to that of galactose monomers.

### DISCUSSION

In summary, we demonstrated that an intranasal synthetic peptide vaccine which includes the four LC3 epitopes recognized by IgA antibodies from immune human subjects, with CT as the adjuvant, was highly immunogenic in baboons as determined by ELISAs for peptide-specific serum IgA, serum IgG, and intestinal IgA antibodies. The peptide vaccine elicited intestinal IgA antibodies that reacted strongly with immunofinity chromatography-purified native lectin protein as demonstrated by ELISA; peptide vaccine-elicited serum IgA or IgG antibodies exhibited low levels of reactivity to LC3 or immunofinity chromatography-purified lectin. To further address the recognition of lectin molecules by peptide vaccine-elicited antibodies, we performed IFAs and Western blot analysis and studied in vitro amebic adherence to CHO cells. IFAs demonstrated serum IgG antibody recognition of lectin molecules on the surfaces of axenic trophozoites, and immunoblotting demonstrated the recognition of the lectin heavy-subunit protein present in amebic lysates. Therefore, the lower level of reactivity of peptide vaccine-elicited serum antibodies than of recombinant LC3 protein-elicited antibodies to purified lectin protein as evaluated by ELISAs may have been due to the alteration of the peptide epitopes on the native lectin molecule by the detergent solubilization and immunoaffinity chromatography purification process. The recombinant LC3 protein, as a positive control, was effective in eliciting serum IgG and IgA antibodies to itself and to purified native lectin protein. However, the recombinant LC3 vaccine did not elicit a mucosal immune response. In addition, the serum anti-LC3 IgG and IgA antibodies elicited by the LC3 vaccine did not recognize the four peptide epitopes recognized by serum and intestinal IgA antibodies from immune human subjects (2). Therefore, recombinant LC3 protein would not be suitable for use as an intranasal subunit vaccine to elicit a protective mucosal immune response. Importantly, the synthetic peptide vaccine elicited neutralizing serum and fecal antilectin antibodies as evidenced by a dose-dependent inhibitory effect on the galactose-specific in vitro adherence of viable *E. histolytica* trophozoites to CHO cells.

To our knowledge, this is the first experimental study of an intranasal synthetic peptide amebiasis subunit vaccine. Investigators from our laboratory were also the first to report that a purified *E. histolytica* protein, the galactose-inhibitable lectin, is highly efficacious in a gerbil model of ALA (33). Gerbils immunized with native lectin protein exhibited a protective efficacy of 70% against intrahepatic challenge with axenic trophozoites; however, a subset of vaccinated animals developed larger abscesses than controls (33). Concern that vaccination with purified lectin may induce adherence-enhancing antibodies (34) or be immunosuppressive were raised. Therefore, the need for a more defined amebiasis subunit vaccine became evident. We subsequently reported that purified recombinant LC3 protein under the same experimental conditions was equally effective in the gerbil ALA model (40) without any paradoxical worsening of disease in those immunized (40). The gerbil ALA model has been utilized for the study of numerous amebiasis subunit vaccines, including native and recombinant forms of the galactose-inhibitable lectin (34, 40), the EhADH112 protein (29), and recombinant *Yersinia enterocolitica* expressing hybrid type III proteins containing the cysteine-rich section of the lectin heavy subunit (26). A DNA vaccine expressing the serine-rich protein SREHP (45) and a 24-mer derived from the lectin heavy subunit chemically coupled to keyhole limpet hemocyanin (27) are also protective in experimental gerbil or hamster liver abscess models. However, immunity to experimental ALA is dependent on cellular rather than mucosal immune mechanisms (24), bypasses the natural gastrointestinal route of infection, and thus is not analogous to the study of asymptomatic intestinal infection in humans.

In a susceptible-mouse (C3H) model, native galactose-inhibitable lectin and, to a lesser extent, a 64-kDa fragment of the lectin heavy subunit with Freund’s adjuvant provide protection against experimental cecal inoculation (22). Studies with resistant mouse strains indicated that neutrophil and innate immune mechanisms are protective (5). Passive transfer with antibodies to the amebic proteophosphoglycan, an abundant surface molecule, prevented colitis in human intestinal xenografts placed in SCID mice (46). As mice are not natural hosts of *E. histolytica* and have vaccine immunity mediated by
secretory IgA-independent mechanisms and epitope recognition differing from that observed following human infection (2, 40), we chose to use a primate model.

Baboons get infected with *E. histolytica* and *E. dispar* in the wild and in captivity (23, 25, 42) and are susceptible to amebic colitis (25, 42). They have the same immunoglobulin subclasses as humans (39). Upon the screening of a large, asymptomatic baboon colony, we found that antilectin IgA antibodies from

![Graph A](image1.png)

**FIG. 6.** Inhibition of amebic galactose-specific adherence to CHO cells by sera (A) or fecal supernates (B) from peptide (1,600-μg-dose)-immunized or control baboons. Axenic *E. histolytica* trophozoites were incubated at 4°C with or without increasing concentrations of galactose, sera, or fecal supernates. As expected, increasing concentrations of galactose demonstrated a dose-dependent inhibition of ameba-CHO cell rosette formation (*P* <0.01 for each concentration compared to control media). Sera from peptide-immunized baboons (*P* <0.01 at each dilution) and sera from baboons receiving adjuvant only (*P* <0.001 at each comparable dilution) demonstrated significant inhibition of amebic galactosespecific adherence compared to control media. The adherence-inhibitory effects of immune baboon sera were no different from those of galactose. Fecal supernates from peptide-immunized baboons inhibited amebic adherence to CHO cells compared to fecal supernates from baboons receiving CT adjuvant only (*P* <0.01 at each dilution). The adherence-inhibitory effects of fecal supernates from immunized baboons were also concentration dependent (*P* <0.01 for each dilution compared to the proceeding one) and comparable to the adherence-inhibitory effects of galactose.
naturally infected baboons recognized the same four LC3 peptide epitopes as antibodies from seropositive humans, further confirming the suitability of baboons as an experimental vaccine host. The nasal route of vaccination offers many advantages, including easy access, lack of the need for sterile syringes, and the successful induction of mucosal and systemic immune responses, and requires lower doses of antigen than the oral route (13, 31). Antigen-specific B and T lymphocytes migrate from nose-associated lymphoid tissues to regional lymph nodes and distant effector sites (31, 45). Intranasal vaccination is more effective than the intragastric route in generating a rapid, retained, and strong mucosal immune response (44); mucosal immune responses in the gastrointestinal tract do occur after intranasal vaccination (11).

Adjuvants known to be effective in animal models of intranasal vaccination include CT, the subunit of CT (CTB), Escherichia coli heat-labile enterotoxin, CTA1-DD (a protein produced by the fusion of the A1 subunit of CT [CTA1] with a staphylococcal protein A-derived protein that targets B cells), interleukin-10 [IL-10], and IL-12 (4, 11, 30). Microbiologically derived antigens that have been successful in experimental intranasal vaccines include E. coli outer membrane particles, the human papillomavirus, Mycobacterium bovis, tetanus toxoid, the VP6 protein of rotavirus, influenza virus, schistosoma antigen, and human immunodeficiency virus, among others (11, 18, 30). CT is a very effective mucosal adjuvant for vaccines administered via the intranasal route (18, 31) due to the induction of CD4+ T helper cells secreting IL-4 and IL-5; these TH2 cells support the development of systemic IgG1 and IgG2b subclasses and mucosal secretory IgA responses (28). However, when delivered to mice, CT and CTB accumulate in the olfactory nerves and the olfactory bulb (17). This accumulation is due to the binding of CTB to gangliosides and the direct neural connections of the olfactory nerves to the olfactory bulb, raising the possibility of nervous tissue damage (9). CTA1-DD holds promise as a genetically engineered B-cell-specific adjuvant that should avoid such toxicity, as it does not traffic to the olfactory bulb (15). Therefore, intranasal vaccination appears promising as a safe method for the induction of protective mucosal immunity with the minimum use of vaccine antigen.

In summary, based on investigations of antilectic mucosal immunity in prospective field studies (1, 36), followed by IgA epitope mapping (2), we designed a polylysine-linked synthetic peptide amebiasis subunit vaccine. When delivered intranasally to baboons with CT as the adjuvant, this peptide vaccine was successful in eliciting high levels of neutralizing intestinal antilectic IgA antibodies. Studies to develop a reproducible baboon model of asymptomatic E. histolytica intestinal infection so as to determine vaccine efficacy are in progress. In addition, the use of other biomaterials and adjuvants to enhance the immunogenicity and safety of this subunit peptide vaccine will be necessary before conducting studies with humans.

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