Randomized Clinical Trial Assessing the Safety and Immunogenicity of Oral Microencapsulated Enterotoxigenic Escherichia coli Surface Antigen 6 with or without Heat-Labile Enterotoxin with Mutation R192G

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An oral, microencapsulated anti-colonization factor 6 antigen (meCS6) vaccine, with or without heat-labile enterotoxin with mutation R192G (LT_{R192G}) (mucosal adjuvant), against enterotoxigenic Escherichia coli (ETEC) was evaluated for regimen and adjuvant effects on safety and immunogenicity. Sixty subjects were enrolled into a three-dose, 2-week interval or four-dose, 2-day interval regimen. Each regimen was randomized into two equal groups of meCS6 alone (1 mg) or meCS6 with adjuvant (2 μg of LT_{R192G}). The vaccine was well tolerated and no serious adverse events were reported. Serologic response to CS6 was low in all regimens (0 to 27%). CS6-immunoglobulin A (IgA) antibody-secreting cell (ASC) responses ranged from 36 to 86%, with the highest level in the three-dose adjuvanted regimen; however, the magnitude was low. As expected, serologic and ASC LT responses were limited to adjuvanted regimens, with the exception of fecal IgA, which appeared to be nonspecific to LT administration. Further modifications to the delivery strategy and CS6 and adjuvant dose optimization will be needed before conducting further clinical trials with this epidemiologically important class of ETEC.

The significant worldwide burden of diarrheal disease due to enterotoxigenic Escherichia coli (ETEC) in children of developing countries has been well documented (5, 32, 39, 45). In addition, military and civilian travelers are at high risk of acquiring ETEC-associated diarrhea when visiting regions where it is endemic (1, 6, 15, 19, 21), which prompts ongoing vaccine development. Prevalence surveys have documented the significant contribution of CS6-ETEC (ETEC containing surface antigen 6; 10 to 37% of ETEC isolates) relative to the overall global ETEC burden (13, 29, 33, 36, 37, 42, 46, 47). Based on this information, this unique nonfimbrial colonization factor (CF) was selected as a target antigen for vaccine development (17, 18, 24, 25, 27, 32, 43, 48).

ETEC vaccine development has been based on two strategies: blocking adherence and/or toxin activity (32, 39). CFs are necessary for ETEC to adhere to the intestinal mucosal lining. After adherence, heat-labile toxin (LT), heat-stable toxin (ST), or both are expressed, resulting in watery diarrhea. The mucosal adjuvant LT is immunogenic but causes unacceptable gastrointestinal toxicity (3, 28). LT_{R192G} is a mutant form of LT which retains immunogenicity and adjuvanticity with greatly reduced toxicity, as documented by in vitro assays, animal models, and clinical trials (3, 11, 14).

Incorporation of CFs into poly(DL-lactide-co-glycolide) (PLG) microspheres has been attempted in order to improve antigen delivery and uptake at mucosal inductive sites (30). In a previous study, microencapsulated CS6 (meCS6) was safe and well tolerated when delivered orally as either a 1- or 5-mg dose in a three-dose, 2-week interval regimen (23). The highest immune responses were observed in subjects (n = 5) receiving the 1-mg dose in a buffered solution, with 80% antibody-se-
TABLE 1. Baseline characteristics of study participants

| Characteristic | Regimen* | Day 0-14-28 | Day 0-2-4-6 |
|----------------|----------|-------------|-------------|
|                | meCS6+   | meCS6-      | meCS6+      | meCS6-      |
| No. of subjects | 15       | 15          | 15          | 15          |
| Age             | 31 (18-45) | 36 (20-45) | 31 (20-44) | 30 (19-45) |
| Male gender†    | 9 (60)   | 10 (67)     | 11 (73)     | 6 (40)      |
| Race/ethnicity  |          |             |             |             |
|                 |          |             |             |             |
| African-American| 8 (53)   | 8 (53)      | 4 (26)      | 9 (60)      |
| Asian-American  | 1 (7)    | 0 (0)       | 0 (0)       | 1 (7)       |
| Caucasian       | 5 (33)   | 5 (33)      | 7 (47)      | 5 (33)      |
| Hispanic         | 0 (0)    | 0 (0)       | 1 (7)       | 0 (0)       |
| Hispanic, non-Caucasian | 1 (7) | 0 (0) | 2 (13) | 0 (0) |
| Pacific Islander| 0 (0)    | 1 (7)       | 0 (0)       | 0 (0)       |

| Serology          |        |             |            |            |
|-------------------|--------|-------------|------------|------------|
| Anti-CS6 IgA      | 5.6    | 5.2         | 6.6        | 7.1        |
| Anti-LT IgA       | 25.7   | 25.7        | 16.2       | 13.8       |
| Anti-CS6 IgG      | 7.2    | 7.2         | 28.8       | 38.9       |
| Anti-LT IgG       | 69.2   | 104.7       | 49.0       | 63.1       |
| Fecal IgA‡        | 2.6    | 3.3         | 3.4        | 4.6        |
| Anti-CS6 IgA      | 3.2    | 4.5         | 6.7        | 5.9        |
| Anti-LT IgA       | 0.6 (5)| 2.2 (3)     | 0.8 (2)    | 1.0 (1)    |
| Anti-LT IgG       | 1.0 (4)| 18.8 (2)    | 1.0 (3)    | 1.5 (2)    |

| MATERIALS AND METHODS |

Volunteer eligibility and randomization. Sixty healthy male and female subjects, 18 to 45 years of age, were recruited from the Washington, DC, metropolitan area. Exclusion criteria included residential, travel, or occupational factors in which a potential for exposure to ETEC, LT, or cholera toxin existed; absence of any prior allergy to any vaccine. The subjects eligible for the study were randomized using a block size of three into one of four treatment regimens as shown in Table 1.

Vaccines. (i) Vaccine composition. The test articles were meCS6, containing the putative colonization factor CS6, and LT<sub>R192G</sub>, a mucosal adjuvant and modified ETEC enterotoxin. meCS6 consists of purified CS6 antigen encapsulated in PLG microspheres, produced using current good manufacturing practices (cGMP) at the Walter Reed Army Institute of Research (WRAIR), Silver Spring, MD. The four genes required for CS6 expression were cloned into an ETEC strain E8775 plasmid vector pUC19 with ampicillin resistance replaced by kanamycin resistance (48). The plasmid was transformed into E. coli HB101, with the resulting strain, M346, used to prepare the vaccine master cell bank. Bulk purified CS6 was loaded into PLG microspheres by the solvent evaporation technique (7). The final product was lyophilized in single-dose vials containing 81.3 mg total dry weight of microspheres with 0.951 mg CS6 protein. The recombinant E. coli strain JM83(pL326), used as a source of LT<sub>R192G</sub>, was constructed at Tulane University in the laboratory of John Clements (11). LT<sub>R192G</sub> was manufactured under cGMP at the WRAIR. The final product was lyophilized in vials containing 1,000 μg of protein per vial. The test articles were delivered in CeraVacXII (Cera Products, LLC, Jessup, MD), an oral electrolyte solution (osmolality, 270 mmol/liter), containing 1.5 g of sodium bicarbonate, 0.375 g of trisodium citrate, and 5.25 g of rice syrup solids in the administered volume. This product was manufactured under cGMP and USDA/FDA guidelines. CeraVacXII (derivative of CeraVac) was designed and formulated as a buffer to optimize the maintenance of a favorable pH throughout the gastric digestion process, increase gastric emptying, and increase absorption of buffer after completing its course through the stomach (35).

(ii) Vaccine administration. Single-dose vials of meCS6 were reconstituted with 10 ml of CeraVacXII and then mixed with 140 ml of CeraVacXII to a final volume of 150 ml per dose. On each vaccination day, LT<sub>R192G</sub> was reconstituted with sterile water and diluted in phosphate-buffered saline (PBS) to achieve a dose of 2 μg per 0.4 ml solution and added just prior to ingestion (within 30 min). Subjects fasted for 90 min pre- and postvaccination.

(iii) Postvaccination follow-up. Safety monitoring included the following: 30-minute postvaccination observation, 24-hour telephone clinical checks, and diaries (capturing the number and grade of bowel movements). Additional clinical visits occurred through postvaccination day 28. For the three-dose, 2-week dosing interval (Day 0-14-28 regimen), diaries were provided for a 7-day period after each dose. For the four-dose, 2-day dosing interval regimen (Day 0-2-4-6 regimen), diaries were provided during the 6-day vaccination period and for 7 days after the final dose. Safety laboratory assessments (complete blood count and analysis of serum aspartate aminotransferase, alanine aminotransferase, and creatinine) were checked on days 7 and 35 for the Day 0-14-28 regimen and days 9 and 21 for the Day 0-2-4-6 regimen. Adverse events (AEs) were graded as mild (transient discomfort that did not interfere in a significant manner with the subject’s normal function), moderate (produced limited impairment of function and could require therapeutic intervention but produced no sequelae), or severe (resulted in a marked impairment of function and could lead to temporary inability to resume usual life pattern; might also produce sequelae that require prolonged therapeutic intervention).

Laboratory methods. (i) Serology assays. The Day 0-14-28 group had their serum assayed for CS6- and LT-specific IgA and IgG antibodies at baseline, 2 weeks postdose for the first two doses, and 4 weeks postdose for the third dose. The Day 0-2-4-6 group serology assays were done at baseline and 3, 7, 14, and 28 days after the dose series. Anti-CS6 antibodies were measured by enzyme-linked immunosorbent assay (ELISA) (2, 16, 20). Individual microtiter wells (Nunc immunoplates) were coated with 100 μl of a 1.0-μg/ml PBS solution of CS6 antigen, incubated overnight at 37°C, washed with PBS, and blocked with 0.1% bovine serum albumin (BSA) (Sigma). The CS6 antigen was generated at bench scale using the identical identification and purification process as for the cGMP (vaccine) lot, with a purity of >95%, as determined by sodium dodecyl sulfate-polyacrylamide gel electrophoresis and densitometric scanning (10). Threefold serial dilutions of sera were made (starting dilution of 1:5), and plates were incubated at room temperature (RT) for 90 minutes. Rabbit anti-human IgA horseradish peroxidase (HRP) conjugate (Jackson) or rabbit anti-human IgG HRP conjugate (Jackson) were added for 90 minutes of incubation at RT. The wells were developed for 20 minutes with orthophenylenediamine (Sigma), followed by optical density (OD) determination at 450 nm in a Titertek spectrophotometer. The end-point titer was determined as the reciprocal of the interpolated dilution giving an OD of 0.4 OD units above background. Anti-LT antibodies were assayed by GM<sub>1</sub>-ELISA (38, 40). Microtiter wells were coated with 100 μl of a 0.5-μg/ml PBS solution of GM<sub>1</sub> ganglioside (Sigma), incubated at RT overnight, washed with PBS, and blocked, followed by coating with 100 μl of a 0.5-μg/ml solution of LT<sub>R192G</sub> (cGMP vaccine lot) in PBS. The procedure continued in the same manner as for anti-CS6 antibodies. Serum samples that demonstrated a ≥2-fold rise in titer above the baseline were considered to indicate seroconversion, provided that the postimmunization titer was ≥10.

(ii) Fecal IgA assay. The Day 0-14-28 groups had assayed done for CS6- and LT-specific fecal IgA at baseline and weekly thereafter (except for day 49) until day 56. The Day 0-2-4-6 groups had assayed at baseline and at days 4, 6, 9, 14, 21, and 28. Stool samples were collected for up to 8 h after defecation if possible and stored and transported to the study site by the subjects (in a transport container with an ice pack). The samples were frozen at −80°C until the time of assay. Fecal extracts were assayed based on methods previously described (2). For total IgA, Nunc immunoplate wells were coated with 100 μl goat anti-human F(ab′)<sub>2</sub>
IgG (Jackson) in a 1.0 µg/ml PBS solution. The procedure was continued in the same manner as the serum antibody measurements, using threefold dilutions of fecal extracts and 100 µl of goat anti-human IgA HRP (Jackson; 1:3,000 dilution). Anti-CS6 and LT IgA determinations were done in the same fashion, using CS6 (1.0 µg/ml in PBS) and GM3 (0.5 µg/ml in PBS) (Sigma) followed by LT192G or LT192G-Δ (as a coating antigen) and 100 µl of rabbit anti-human IgA HRP dilution 1:2,500 in 0.1% BSA-PBS-0.05% Tween as the secondary antibody. For CS6 and LT, BSA well OD values were subtracted from the test antigen well OD values to adjust for background. A positive response was defined as a ≥2-fold increase in vaccine-specific IgA antibody titer per total IgA between baseline and postvaccine samples. Samples were excluded if the total IgA content was <10 µg/ml, if the difference in total IgA concentration between baseline and postvaccine specimens was >10-fold, or if specific baseline titers were ≥2 standard deviations above the group mean (unless there was a significant difference between baseline and postvaccine titers).

(iii) IgA-ASC assay. Whole blood was collected in EDTA-containing tubes at baseline and 1 week after each dose of vaccine for the Day 0-14-28 groups. For the Day 0-2-4-6 groups, specimens were obtained at baseline and on days 6, 9, 14, 21, and 34. An enzyme-linked immunospot assay was initially conducted in real time using fresh EDTA-blood specimens according to previously described methods (23, 44). Due to technical difficulties in interpreting ASC well counts, ASCs were assayed with cryopreserved cells from the same volunteer specimens. MNCs were isolated by using a Ficoll-Hypaque density gradient and cryopreserved in a commercially available freezing medium (C-6164; Sigma Chemicals) at a controlled rate and stored in liquid nitrogen until assayed (4). Antigen-specific IgA-ASCs were determined using an enzyme-linked immunospot assay. Briefly, after thawing and washing, viability was determined using Guava ViaCount reagent (Guava Technologies: 4000-0041). Viable cells (average viability, 91.1% ± 3.2%; n = 277) were adjusted to 3.3×10^6/ml in medium containing 10% fetal calf serum, 2 mM l-glutamine, and 50 µg/ml gentamicin (complete medium). Multiscreen immunoplates (Millipore SE2M004M99) were coated with antigens. After blocking with 5% fetal calf serum in RPMI, 100 µl of complete medium containing 3.3×10^6 MNCs was added to triplicate. Plates were incubated for 4 to 5 hours at 37°C in a 5% CO2 environment, followed by an additional 2-hour incubation at 37°C after addition of 0.025 µg of alkaline-phosphate-conjugated goat anti-human IgA (KPL). Spots were developed using Nitro Blue Tetrazolium and 5-bromo-4-chloro-3-indolyl phosphate substrate (SigmaFast; Sigma Chemicals). After an additional 15 minutes of incubation, the plates were extensively rinsed with water and air dried. Spots were counted using a CTL spot analyzer (Cellular Technology Ltd.). For each antigen, data were expressed as the number of ASCs per 10^6 MNCs. A positive response was defined as a ≥2-fold increase over the baseline value of antigen-specific ASCs per 10^6 MNCs, when the number of ASCs was ≥0.5 per 10^6 MNCs in the baseline sample and ≥1.0 ASCs per 10^6 MNCs if preimmune ASCs were <0.5 per 10^6 MNCs.

(iv) Statistical analysis. The primary safety outcome was enterotoxicity that was possibly or probably related to the test articles. Enterotoxicity was indicated by postimmunization gastrointestinal AEs, including loose stools, diarrhea (defined as ≥3 loose liquid stools in a 24-hour period), abdominal cramps, nausea, and vomiting. Gastrointestinal AEs with onset within 3 days postvaccination were graded mild. No clinically significant changes from baseline were observed.

RESULTS

Demographics and study conduct. Sixty subjects were enrolled and received at least one dose of vaccine. Volunteer demographics overall were an age range of 18 to 45 years (mean 32), 60% male, and ethnicity as follows: African-American, 48%; Caucasian, 37%; Hispanic, 7%; Asian, 5%; and other, 3%. Table 1 characterizes volunteer demographics by study group (regimen). The four groups were similar in demographic makeup. Fifty-six subjects (93%) completed the entire vaccination series, and 58 (97%) complied with safety monitoring at least 28 days after receipt of their last vaccine dose. Three of the subjects were unable to comply with the study schedule, and one volunteer did not receive vaccine after the first dose due to an AE not related to the test articles.

Safety profile. Overall, the vaccine was well tolerated regardless of dosing regimen or receipt of LT R192G, and no serious AEs occurred. Table 2 shows the frequency of surveyed symptoms possibly or probably related to the test articles. Interim laxative use was the most-common reported symptom (37% of subjects), although only two subjects met the definition for diarrhea. Abdominal cramping (13% of subjects) and nausea (12% of subjects) were the next most commonly reported symptoms; a headache and malaise were each reported by one volunteer. There was no significant difference in the frequency (or number, in the case of number of loose stools) of these events across vaccine groups.

One subject meeting the diarrhea definition reported four grade 4 stools (without associated symptoms) within 24 hours after receiving the fourth dose of meCS6 plus adjuvant in the Day 0-2-4-6 regimen. The diarrhea did not interfere with daily activities and resolved spontaneously. The other subject meeting the diarrhea definition reported four grade 3 stools over a 4-hour period (with associated moderate upper abdominal pain and nausea) 6 days after the first dose of meCS6 without adjuvant (Day 0-14-28 regimen). Resolution of all symptoms occurred within 24 hours, with no symptoms with subsequent doses.

Another subject reported abdominal cramps precluding daily activities in the evening following the first dose of meCS6 with adjuvant (Day 0-2-4-6 regimen). The cramps lasted 90 min, during which time two grade 3 stools were passed, with spontaneous resolution. Reports of moderate grade AEs included two subjects with short-lived abdominal cramps. All other symptoms possibly or probably related to the test articles were graded mild. No clinically significant changes from base-

| No. (%) for regimen: |  |  |  |  |
|----------------------|---|---|---|---|
| AE                   | Day 0-14-28 | Day 0-2-4-6 | Day 0-14-28 | Day 0-2-4-6 |
| meCS6+               | meCS6−       | meCS6+       | meCS6−       | meCS6−       |
| Diarrhea^c           | 0 (0)       | 1 (7)       | 1 (7)       | 0 (0)       |
| Loose stools^d       | 5 (33)      | 5 (33)      | 5 (33)      | 5 (33)      |
| Abdominal cramps     | 1 (7)       | 2 (13)      | 4 (27)      | 1 (7)       |
| Nausea               | 0 (0)       | 4 (27)      | 1 (7)       | 2 (13)      |

^c Limited to actively monitored events determined to be possibly, probably, or definitely related to the vaccine (determined by temporal relationship and no other clearly evident cause). No vaccine-related vomiting or fever was noted. One person in the Day 0-2-4-6 meCS6− group had a headache and malaise that were possibly vaccine related.

^d Three or more loose liquid stools per 24 h.

Two or less loose liquid stools per 24 h. Does not include subjects with diarrhea.
line levels in hematology or chemistry safety laboratory values occurred in any group. Fifty-one (85%) subjects were available for phone follow-ups 6 months after the last vaccination. None reported any serious health problems or hospitalizations since study completion.

**Immune responses.** A total of 57 subjects (95%) received at least two (Day 0-14-28) or three (Day 0-2-4-6) vaccine doses required for inclusion of data in the primary immunologic analyses. Table 3 presents summary response rates and PFRs or maximum numbers of cells (ASCs) for all groups.

**Serology.** Response rates and serology kinetics for anti-CS6 and -LTR192G antigens are presented in Table 3 and Fig. 1 and 2. Baseline titers (Table 1) of anti-CS6 IgG in the Day 0-2-4-6 group were comparable to those in the D. E. Katz study (unpublished data), while those in the Day 0-14-28 group were significantly lower ($P < 0.0001$). Response rates to CS6 were similar (21 to 27%) in three groups, with no responders in the fourth group (unadjuvanted Day 0-14-28). The IgG response rate to CS6 in the Day 0-2-4-6 group without adjuvant trended toward being significant compared with the Day 0-14-28 group without adjuvant (Fisher’s exact test; $P = 0.10$), with titer PFRs significantly higher (Wilcoxon rank-sum test; $P = 0.04$). The ASC response rate was also higher (not significant) in the Day 0-2-4-6 adjuvanted group versus the unadjuvanted group. The median peak number of anti-CS6 ASCs per $10^6$ MNCs was $=3$ across all groups. Forty-two percent of anti-CS6 and 60% of anti-LT peak responses were seen after the third vaccine dose in the Day 0-14-28 adjuvanted regimen. In all other groups, maximum ASC counts were evenly distributed across all vaccine doses.

**DISCUSSION**

Our study demonstrated that meCS6, with or without LT$_{R192G}$ was safe when administered in two different regimens (four-dose, 2-day interval and three-dose, 2-week interval), with an anti-CS6 ASC response rate of 86% when adjuvanted in the three-dose regimen. One episode of diarrhea occurred in 1 of 30 subjects, which was consistent with

### Table 3. Frequency of responses to vaccine antigens by group

| Antigen Response type | Result for regimen: |
|-----------------------|----------------------|
|                       | Day 0-14-28 | Day 0-2-4-6 |
| meCS6+ | meCS6- | meCS6+ | meCS6- |
| CS6 IgA (feces) | 14 (50.4) | 21 (5.1) | 43 (8.2) | 40 (7.2) |
| IgA (serum) | 7 (4.4) | 0 (n/a) | 14 (11.0) | 20 (15.0) |
| IgG (serum) | 21 (3.9) | 0 (n/a) | 21 (4.6) | 27 (3.1) |
| LTR192G IgA-ASC | 86 (1.3) | 36 (1.7) | 57 (2.7) | 47 (2.5) |
| LT IgA (feces) | 50 (10.0) | 57 (8.3) | 71 (7.7) | 53 (6.0) |
| IgG (serum) | 57 (4.1) | 0 (n/a) | 36 (7.4) | 0 (n/a) |
| IgA-ASC | 43 (9.7) | 7 (2.3) | 36 (5.4) | 0 (n/a) |
| IgA-ASC | 71 (5.5) | 0 (n/a) | 57 (3.3) | 7 (2.3) |

*Serology responses are summarized as the percentages of volunteers meeting the responder definition and the median PFRs from the baseline titer among responders (in parentheses). ASC responses are summarized as the percentages of volunteers meeting the responder definition and the median maximum numbers of ASCs among responders (in parentheses). +, with adjuvant; −, without adjuvant.

*Mean Geometric Titer (Log Scale)

*FIG. 1. Serologic responses for Day 0-14-28 regimen. +, with adjuvant; −, without adjuvant.
LT_{R192G}-related enterotoxicity (four-dose adjuvanted group). In contrast, in a Helicobacter pylori killed whole-cell vaccine study (placebo controlled), with and without LT_{R192G} (25 μg), diarrhea occurred in 6 of 31 (19%) subjects, all from adjuvanted vaccine groups (26). Loose stools not meeting the definition of a diarrhea episode, a more-sensitive indicator of entero toxicity, were evenly distributed across the vaccine groups in the current study.

PLG-encapsulated and nonencapsulated CS6 antigens have been evaluated for safety and immunogenicity in recent studies of purified CFs and enterotoxoids. In the study by Güreña-Burguéñ o et al., nonencapsulated CS6 was administered transcutaneously with and without LT (500 μg) at 0, 1, and 3 months with good anti-CS6 serologic and ASC responses in the adjuvanted groups (14). The rationale for incorporating CS6, an antigen intended to target the mucosal immune system, into a microparticle system was informed by encouraging preclinical studies testing oral immunization using PLG vaccines (7, 31). In humans, there has been limited experience with enteric vaccines incorporated into PLG microspheres (41). PLG-encapsulated and nonencapsulated CS6 antigens were administered orally in the study by Katz et al. (23). In five subjects who received 1 mg of unadjuvanted meCS6 with buffer, three were anti-CS6 IgA responders (PFRs of 17 to 77), four were anti-CS6 IgG responders (PFRs of 4 to 153), and four were IgA-ASC responders (peak of 6 to 41 spots per 10^6 MNCs).

In comparing results in the unadjuvanted Day 0-14-28 group (n = 14) in this study with those for the smaller group (n = 5) in the Katz et al. study (no adjuvant used), which received the same regimen, we note considerable differences in qualitative and quantitative responses to CS6. A factor that may have accounted for the difference in immunogenicity in our study is a new lot of meCS6 with some differences in various parameters compared to the lot used by Katz et al. Two characteristics of the two lots for consideration are the proportion of meCS6 with the desired particle size of 4 to 11 μg (94%) in this lot versus the proportion (52%) in the previous lot and the in vitro release of CS6 at 24 h (39%) in this lot versus the release (53%) in the previous lot. The core antigen load of meCS6 was similar in both lots (1.17% in this lot versus 1.01% in the previous lot). Whether any of these differences in vaccine characteristics affected the immunogenicity of the vaccine in humans is not known. The effects of the microencapsulation process on specific epitopes of the CS6 bulk-purified protein in relation to the specific uptake in the human small intestine are unknown. Preclinical immunogenicity measures done on both meCS6 lots demonstrated good immune responses when the vaccine was given subcutaneously to rabbits and intranasally to mice. Mice did not respond to vaccine delivered orally (8). The same lots and clinical formulations used in this study were also tested for safety and immunogenicity in Aotus nancymae monkeys using the same doses and schedules as in the clinical trial. The Day 0-14-28 regimen was more immunogenic (only serologic responses were measured). Six of seven (86%) monkeys given 1 mg of meCS6 in buffer orally without adjuvant had serum IgG responses with mean PFRs of 78.7 ± 7.8 (22). Currently, the sensitivity of in vitro and animal immunogenicity tests for predicting CS6 immune responses in humans is unknown. The vaccination procedures had what are seemingly minor differences. In the Katz et al. study, the vaccine, reconstituted in 100 ml of CeraVacx, was ingested, followed by 50 ml of CeraVacx without vaccine within 5 min. In our study, the vaccine was reconstituted in 150 ml of CeraVacxII (same buffering formula as CeraVacx, with a small amount of flavoring added), with no postdose buffer.

Sample size differences between the two studies (statistical effects) may explain some of the difference in the results in the two studies. The impress serologic and IgA-ASC response rates and the magnitude of responses seen in the small (n = 5) group in the Katz et al. study were unique to that group compared to results for the other groups of three to five subjects, even those who received 5 mg of meCS6 with buffer. Potential differences in the study populations’ prior immunity may explain part of the variability in results. Demographic factors in our study populations were comparable to those of the first meCS6 study by Katz et al. (23): residence in the Washington, DC, metropolitan region; age range of 21 to 44 years (mean, 40); 66% male; ethnicity (African-American, 60%; Caucasian, 21%; Hispanic, 3%; other, 8%); and 80% of subjects completing the entire vaccination series.

There is limited information about the immune response to CS6-ETEC in natural disease. Immune responses to CS6-ETEC infections have been measured in Department of Defense personnel with diarrhea in Incirlik, Turkey. In 28 cases of
ETEC diarrheal illness with CS6 as the only CF and with no other enteric pathogens isolated, mean peak anti-CS6 IgA titers were 20, mean peak anti-CS6 IgG titers were 240, and median peak IgA-ASC counts were 5 spots per 10^6 MNCs (C. titers were 20, mean peak anti-CS6 IgG titers were 240, and ET. diarrheal illness with CS6 as the only CF and with no VOLUME 15, 2008 PHASE 1 STUDY OF A MICROENCAPSULATED ETEC VACCINE 1227

Disease Research Program, NMRC Work Unit Number A0055. The Department of Defense, or the U.S. Government. The immune responses observed in our study were lower in all groups than prospectively determined thresholds we considered necessary for further testing of the current formulation of meCS6. While an 86% anti-CS6 ASC response was observed in the Day 0-14-28 adjuvanted group immune responses are compared to all CS6 subunit vaccine responses, the response rate is at the high end of the spectrum for ASC response (but with a lower magnitude, in general) and the lower end for serologic rates and magnitudes of response (14, 23). This group had higher ASC responses than subjects challenged with ETEC strain B7A, with similar magnitudes in responders and similar serologic responses. The magnitude of response in our study group is similar to that seen in natural CS6-ETEC infections. The ASC response rate and magnitudes in this group are also comparable to those reported against other CFs (CFA/I, CS1/3, CS2, and CS4) present in an inactivated ETEC whole-cell vaccine. Except for anti-CFA/I, serum IgA responses were also similar compared to these other CFs (2).

The immune responses observed in our study were lower in all groups than prospectively determined thresholds we considered necessary for further testing of the current formulation of meCS6. While an 86% anti-CS6 ASC response was observed in the Day 0-14-28 adjuvanted regimen, low peak ASC counts and poor serologic responses accompanied this result. Our study may have been limited by a suboptimal dose of meCS6 or LT102G, a suboptimal ratio of meCS6 to LT102G, or by not having both test articles coformulated in the microspheres. The microencapsulation technology itself may need further refinement before this delivery system can be taken forward into advanced development, such as targeting ligands or surface exposing the antigen (12, 30). Further concerns are the lack of information known about CS6 compared to better-characterized CFs such as CFA/I.

In conclusion, although CS6 is a good vaccine candidate, the product was not as adequately immunogenic in the current formulation as a microsphere vaccine. However, CS6-ETEC strains are prevalent in many areas where ETEC is endemic, and they are often the only pathogen isolated from diarrheal cases. More research is needed to characterize the role of CS6 in the pathogenicity of ETEC infections in order to develop an effective broad-spectrum ETEC vaccine.

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J.A.L. is a military service member. This work was prepared as part of her official duties.

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