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DEVELOPMENT OF NEW INMUNOGENS AND A CONTROLLED RELEASE DELIVERY SYSTEM FOR URAL INMUNIZATION AGAINST STAPHYLOCOCCAL ENTEROTORIN B

Project 6160-XXV

Annual Report

- Jay K. Staas, John H. Eldridge, Richard H. Gilley, Thomas R. Tice

December 6, 1991

Supported by

US ARMY MEDICAL RESEARCH AND DEVELOPMENT COMMAND Fort Detrick, Frederick, ND 21702-5012

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DEVELOPMENT OF NEW IMMUNOGENS AND CONTROLLED RELEASE DELIVERY SYSTEM FOR IMMUNIZATION AGAINST STAPHYLOCOCCAL EXTEROTOXIN B

I. EXECUTIVE SUMMARY

During this year of the contract, we obtained a great deal of knowledge about the immunization of rhesus monkeys with SEB toxoid. A preliminary study was performed to confirm the existing data rogarding the effectiveness of intramuscular (IM) immunization with alum precipitated SEB toxoid and oral immunization with high doses of toxoid in solution, to raise circulating anti-SEB toxin antibodies. Additionally, several studies involving the immunization of rhesus monkeys with SEB toxoid microspheres have also been performed.

The preliminary monkey experiment yielded the following results: Immunization with alum-precipitated SEB toxoid did result in the appearance of circulating IgG antibodies, which were detected in low levels in the BAW fluids. Oral immunization with SEB toxoid solution induced only the transient appearance of low levels of IgM and IgA anti-SEB toxin antibodies in circulation.

Results from the first two studies (EX-Rh-101 and EX-Rh-102) in which rhesus monkeys were immunized with microencapsulated SEB toxoid are included in this report. These data suggest that an IM primary immunization followed by an intratracheal (IT) or oral secondary immunization provides the highest antibody response and best level of protection. It has been reported to us that the monkeys receiving an IM primary and IT secondary immunization in both studies survived aerosol challenge. In addition, the monkey receiving an oral primary and IT secondary immunization from Study EX-Rh-101 and the monkey receiving both an IT primary and IT secondary immunization in Study EX-Rh-102 also survived the aerosol challenge. Official documentation of these results has not been received by Southern Research or the University of Alabama in Birmingham.

These results are very encouraging as not only have high antibody titers been achieved in the rhesus monkeys, but as indicated above, several of the monkeys have been shown to be protected from aerosol challenge.

II. STATEMENT OF THE PROBLEM UNDER STUDY

The staphylococcal enterotoxins are extracellular proteins produced by <u>Staphylococcal aureus</u> that have been demonstrated to be a major cause of food poisoning. Serological methods have differentiated five classes of enterotoxins, A, B, Cl, C2, D, and E (1-6) (Note: References can be found in Section VII, see Page 22). Despite extensive study of the structure and properties of these enterotoxins, their precise mode of action is

unclear (7). While detection of the toxins and prophylaxis of toxemia is of obvious importance to the food industry and medical community, there is also significant concern from a military perspective about the potential use of staphylococcal enterotoxins as biological-warfare agents.

Over the last 20 years, studies at the United States Army Medical Research Institute of Infectious Diseases (USAMRIID) on the pithophysiological effects of staphylococcal enterotoxin B (SEB) have demonstrated its lethal toxicity at low doses (25 μ g/kg body weight) in rhesus monkeys (8), with presumably similar effects in man. Studies by Liu et al. (9, 10, 11) have shown that SEB administered intravenously to rhesus monkeys in high doses (1 mg/kg body weight) results in death within 20 hours due to pulmonary dysfunction and edams, with secondary effects on cardiovascular, hepatic, and renal function. Under a biological-warfare scenario, military personnel could be exposed to large quantities of SEB, probably in an aerosol form, which might result in lothal toxemis more rapidly than similar intravenous doses, given the sensitivity of the lungs to the toxin. Immunization with toxoids of SEB that result in high levels of neutralizing antibodies might be an effective and inexpensive defense against possible exposure to SEB by military personnel.

The objective of this research program is to develop an SEB toxoid microsphere formulation with biocompatible, biodegradable polymers. After oral administration of these microspheres, the microsphere formulation will be designed to target and control the release of SEB toxoid in the Peyer's patches, so as to elicit a protective secretory immune response.

III. BACKGROUND AND REVIEW OF THE LITERATURE

A. <u>Secretory IgA</u>

The mucosal surfaces of man and other mammals are in direct continuity with the external environment and represent the major bodily size of antigenic exposure and recognition. Mucosal secretions covering these tissues represent a major host defense mechanism, which is often times underestimated in importance. It was recognized only 20 years ago that large differences exist in the proportions of the various immunoglobulin classes present in external secretions as opposed to serum. The discovery by Tomasi and colleagues (12, 13) and Hanson (14) that IgA is the major isotype in human and other mammalian external secretions provided the impulse for numerous studies on the mucosal immune system in both health and disease. The immunoglobulins secreted at these sites are primarily produced by local plasma cells which heavily infiltrate the lamina propria regions of mucosal sites such as the gut, masal passage and salivary glands.

Secretory IgA (SIGA) antibodies are structurally and functionally distinct from the immunoglobulins which make up the recirculating pool in serum. SIGA exists primarily as a dimer in association with a molecule of secretory component (70,000 daltons) and a molecule of J Chain (15,000 daltons). The covalently bound secretory component is hypothesized to wind around the Fc portions of the two IgA molecules and, "hypothesized secondary

interactions, stabilize the IgA molecules against proteolytic cleavage. This structure may provide a distinct advantage in the efficacy of sIgA in the gut and oral cavities, which commonly contain bacteria that produce proteolytic excenzymes. IgA is neither opsonic nor does it fix complement (both characteristics of IgH and most classes of IgG antibodies), but it is highly effective in viral and toxin neutralization (15, 16) and in inhibiting the adherence of bacteris to epithelial coll surfaces (17). The unique characteristics of sIgA are doubtlessly a reflection of the need for this immunoglobulin to function outside the body where it prevents antigen adherence and penetration.

3. Antibodies in the Respiratory Tract

Antibodies present within the respiratory tract originate from two different sources. IgA predominates in the mucus which bathes the nesopharynx and bronchial tree (18, 19). More than 90% of this IgA is in the 115-dimeric form and has attached secretory component and \overline{J} chain, while only a small amount is in the 75-monometric form (20). This distribution is in contrast to the serum IgA of humans, which is virtually all monomeric. The molecular weight of sIgA 390,000-395,000 daltons is well above the approximately 200,000-dalton cutoff isposed by the capillary-alveolar membrane, and the igA/albumin ratio in the bronchial-alveolar wash (BAW) fluids is substantially higher than in serum, indicating that the bulk of bronchial IgA is locally produced (21). In this regard, IgA plasma cells have been shown to line the lamina propris of the sirvay wall and are particularly concentrated about the bronchial glands (18). Dimeric IgA from these plasms cells is bound to secretory component expressed; on the basal surface of bronchial epithelial cells and reaches the external secretions via active transport through this epithelium (22). In contrast to the nasopharynx and bronchial tree, the bronchioli and alveoli predomimantly contain IgG (23). The ability of IgG (158,000 daltons) to freely pass the capillary-alveolar membrane, plus the fact that the IgG/albumin ratio in BAW fluids is the same as that in serum, indicates the bulk of pulmonary IgG is passively derived from the intravascular pool (21, 23).

C. <u>Gut-Associated Lymphoraticular Tissue and the Common</u> <u>Mucosal Immune System</u>

The finding by Graig and Gebra (24) that Peyer's patches which are distinct lymphoreticular follicles along the gastrointestinal tract (GI tract) possess IgA precursor B cells which can repopulate the lamina propria regions of the gastrointestinal and upper respiratory tracts and differentiate into mature IgA synthesizing plasma cells, suggested that the induction of IgA responses is not necessarily a local phenomenon. The Peyer's patches contain a large subpopulation of B lymphocytes that are committed to IgA synthesis, all categories of regulatory T lymphocytes, and functional accessory cells, i.e., macrophages and dendritic cells (25, 26). In addition, Peyer's patches are covered by a unique epithelium which contains microfold cells (M cells) with highly developed pinocytotic channels that allow sampling of antigens from the gut lumen, and transport to cells in the underlying dome region, with subsequent stimulation in the B cell (follicles) and T cell (parafollicular) zones.

Heremans and Bazin et al. (27, 28) measured the development of IgA responses in mice orally immunized with soluble (28) or particulate (27) antigen. A sequential appearance of antigen-specific IgA lymphoblasts occurred, first in mesenteric lymph nodes, later in the spleen, and finally in the lamins propris of the GI tract. Subsequent studies have shown that oral administration of antigen leads to production of sIgA antibodies in the gut and also in secretions distant to the gut, e.g., in bronchial washings, colostrum, milk, saliva, and tears (29-32). Coupled with the findings of Craig and Cebra (24). these results suggested that Peyer's patches are enriched sources of precursor IgA B cells, which, subsequent to antigen sensitization, follow a circular migrational pathway and account for the expression of IgA at distant mucosal surfaces. This circular traffic pattern provides a common mucosal immune system by continually shuttling sensitized B cells to mucosal sites for responses to gutencountered environmental antigens and potential pathogens (33). This circulatory pathway helps explain the presence of naturally occurring IgA antibodies to microorganisms at sites where local antigenic stimulation would not be expected to occur. This is best exemplified by the presence of sIgA antibodies in human colostrum to gut Escherichia coli (32) and the oral pathogen Streptococcus mutans (34).

Of particular importance to this project is the ability of oral immunization to induce protective antibodies in the respiratory tract. In this regard, studies have demonstrated that the ingestion of various antigens by humans (35, 36), primates (37), rabbits (38), rats (39), and mice (36) results in the appearance of antigen-specific sIgA antibodies in bronchial and/or masal washings. Experiments by Waldman and colleagues (36) have shown that when mice are immunized with live influenza virus via the nasal. oral, or rectal routes, protection against aerosol challenge is conferred. The protection correlated with the level of antibodies secreted into the respiratory tract, but no correlation with serum antibody levels was found. In contrast, paranteral immunization was found to be a good method to induce humoral antibodies against influenza, but a poor method to induce pulmonary entibodies and was non-protective. Extension of these studies with human volunteers confirmed that oral administration of influenza vaccine was effective at inducing secretory anti-influenza antibodies in nasal secretions (40).

Several investigations have shown that secretory antibody levels in humans (41, 42) and mice (43, 44) correlate with protection against pulmonary viral infection to a significantly greater extent that do circulating antibody titers. This offers a valid explanation for the observation made over 50 years ago by Bull and McKee (45) that intranasal immunization of rabbits with killed pneumococci resulted in resistance to pulmonary challenge with live homologous pneumococci in the absence of circulating antibodies. Therefore, it appears that oral immunization to stimulate the pulmonary immune system may have many advantages, including effectiveness, safety, decreased side effects, and the potential for an almost unlimited number and frequency of boostings.

IV. RATIONALE USED IN THE CURRENT STUDY

The use of microencepsulation to protect sensitive bioactive agents from degradation and to control their release over extended periods of time in vivo has become quite prevalent. The technique involves the coating of a bioactive agent (solid or liquid) with a protective wall material. The wall materials are usually polymeric in nature. The microsphere product is a free-flowing powder of spherical particles. The agent to be encapsulated can be coated with a single wall of the polymeric material, or it can be homogeneously dispersed within a polymeric matrix. The amount of agent inside the microspheres can be very small or can range to as high as 95t of the microsphere composition. The diameter of microspheres can be less that 1 μ m or as large as 3 mm.

The use of microspheres to deliver vaccine antigens to the Peyer's patches offers several advantages. First, a microsphere formulation can be designed to protect the antigens from degradation during passage through the gastrointestinal tract and then facilitate uptake into the Peyer's patches. After uptake, the microspheres can release the vaccine antigens at a controlled rate over a period of hours to months.

One microsphere system of particular interest involves the use of poly(lactide-co-glycolide)s (PLGs) (46-47). PLGs are biocompatible, biodegradable polyesters and are from the same class of material used in resorbable sutures. They biodegrade in vivo into lactic acid and glycolic acid, eventually carbon dioxide and water. The mechanism of degradation is by hydrolysis of the ester linkages. The rate of degradation for these copolymers is primarily determined by the ratio of lactide to glycolide in the copolymer (58). For instance, DL-FLG with a 50:50 mole ratio of DL-lactide-to-glycolide will completely biodegrade in vivo when administered subcutaneously (SC) or intramuscularly (IM) within about 6 weeks, while poly(DL-lactide)(DL-FL) completely biodegrades in about 10 to 12 months.

One of the major advantages of DL-FLG microsphere systems is the flexibility allowed in formulating the specific duration of release. It is well known that specific antigens require different lengths of exposure to elicit a strong primary response. The time required before re-exposure to elicit the most potent secondary response also varias with different antigens. By using different combinations of DL-PLGs. delivery systems that demonstrate the proper release rates or release program can be prepared for virtually any antigen. More specifically, a formulation can be designed so that part of the antigen is released soon after the microspheres are taken up by the PP to elicit the primary response. Then, no additional antigen will be released from the microspheres until sufficient time has elapsed to obtain an efficacious secondary response. At this time (as a result of degradation of the polymer), additional antigen will be released, potentiating the secondary response. If desired, multiple releases of antigen at different times could be incorporated into the final microsphere formulation. The times at which the antigen would be released from the formulation would depend on the DL-PLGs used in the formulation.

V. EXPERIMENTAL METHODS

. <u>Mice</u>

BALB/c (original breeders obtained from the Jackson Laboratories, Bar Harbour, ME) were bred and maintained in our facilities at the University of Alabama at Birmingham. All mice used in these studies were 8 to 12 weeks of age at the initiation of the procedures and were of mixed sexes.

B. Rhesus Monkey

Hacaqua mulatta monkeys were obtained from the Texas Frimate Center. All monkeys were 9 to 12 months of age upon receipt.

C. <u>Immunologic Respents</u>

Solid-phase absorbed and affinity-purified goat IgC antibodies specific for murine IgM, IgC, and IgA were obtained from (Southern Biotechnology Associates, Birmingham, AL). Their specificity was confirmed in radioimmunometric assays (RIA) using purified monoclonal antibodies and myeloms proteins as substrates. Hybridoma cell lines producing monoclonal antibodies specific for the murine antigens Thy 1.2 [30-H12, rat IgG_{2b} (59)], Ly-1 [53-7.313, rat IgG_{2a} (59)], Lyt-2 [53-6.72, rat IgG_{2a} (59)], L3T4 [GK 1.5, rat IgG_{2b} (60)]. B 220 [RA 3-3A1/6.1, rat IgM (61)], IgM [331.12, rat IgG_{2b} (62)] and MAC-1 [M1/70.15.11.5, rat IgG_{2b} (63)] were obtained from the American Type Culture Collection. (Rockville, MD). All lines were propagated in vitro and the antibodies in the culture supernatants were purified by sequential precipitation at 50% saturation in ammonium sulphate, anion-exchange chromatography (DE-52, Whatman, Kent, England), and sizing on AcA 34 (LKB, Bromma, Sweden) for IgG and AcA 22 for IgM.

D. Radioiodination and Immunoradiometric Assays

Solid-phase absorbed and affinity-purified goat IgG antibodies specific for mouse total immunoglobulins, IgM, IgG and IgA were radioiodinated with carrier-free Na ¹²⁵I (Amersham, Arlington Heights, IL) using the chloramine-T method modified to reduce oxidative damage to proteins (64). Radioimmunometric assays (RIA) were performed in Immulon assay strips (Dynatech, Chantilly, VA) coated with SEB toxoid at 5 μ g/mL in pH 8.4, borate-buffered saline (BES) overnight at 4 °C. Control strips were left uncoaced, but all strips were blocked for 2 h at room temperature with 1% (w/v) bovine serum albumin (Sigma, St. Louis, MO) in BBS, which was also used as the diluent for all samples and ¹²³I-labeled reagents. Various 2-fold dilutions of test sera were added to washed triplicate

replicate wells and incubated for 6 h at 25 °C. After washing, 100,000 cpm of 125 I-labeled isotype-specific anti-immunoglobulin reagent was added to each well and incubated overnight at 4 °C. Following removal of the unbound 125 I-antibodies by vashing, the wells were counted in a Gamma 5500 spectrometer (Beckman Instruments, Irvine, CA). The results have been presented as the reciprocal of the serum dilution producing a signal greater than 3 times that of the group-matched prebleed at the same dilution (end-point titration).

E. <u>Characterization of Microspheres</u>

The core loading for each batch of microspheres was determined by first extracting the protein (SEB toxoid) from a known amount of microspheres. The extracted protein was then quantified using a colorimetric protein assay. The antigen-containing microspheres were also characterized with respect to in vitro release kinetics using the procedures outlined in our First Annual Report, Pages 8 to 10.

F. Procedure for Sampling Rhesus Monkey Lung Secretions

As the major goal of the rhesus studies is to investigate the induction of SEB toxin-neutralizing antibodies in the lungs, the first group of primates was used to develop the procedures for collecting samples of pulmonary secretions. In order for these samples to be useful in the evaluation of the various immunization schemes to be tasted under this contract effort, they must contain a level of total immunoglobulin which is sufficient to allow detection of the antigen-specific component. Further, antibodies of the IgG and IgA isotypes must be well represented in order to draw firm conclusions about the relative contribution of locally synthesized versus blood circulation derived responses.

Initial attempts to obtain secretions using a pediatric bronchoscope were unsuccessful. The internal tx_heal diameter of the young rhesus monkeys was found to be unexpectedly small by comparison to a human infant of comparable weight. The result was that the 4-mm outer diameter of the bronchoscope completely occluded the traches upon insertion. Although samples could be obtained from the proximal airway by lavage, they were of insufficient volume and antibody content to be of use. In addition, the monkeys rapidly became cyanotic, necessitating that the work be performed over such a short period of time as to prevent careful and reproducible sample collection.

To overcome the above limitations the sampling procedure was changed to one which employs the insertion of a tracheal tube, through which a suction catheter may be passed. This approach has proven to be highly successful, and has been adopted for these primate studies. In brief, fasted monkeys were anesthetized by the intramuscular injection of ketamine hydrochlorids. After obtaining a blood sample, and gastric contents were removed by suction through an 8-French x 14-inch pediatric feeding tube inserted into the stomach. Intratracheal intubation was accomplished with a 3-French x 12-cm pediatric tracheal tube which was inserted through the

glottis with the aid of a 6-French intubating stylet. After taping the trachesl tube in place, the animal was briefly ventilated with humidified 100% oxygen to prevent hypoxemia during the subsequent steps. One mL of phosphate buffered saline (PBS) was placed into the hub of the tracheal tube and allowed to pass into the lungs by normal respiratory action. An 8-French x 14-inch suction catheter, attached to a vacuum line through a 15-mL trap, was immediately passed through the tracheal tube to recover the wash fluid. The suction catheter was then withdrawn and the mucus adhering to the bore of the tube was washed into the trap with 2 mL of additional PBS. The samples were clarified by centrifugation. Sodium aride, phenylmethyl-sulfonyl fluoride, and fetal calf serum were added as preservative, protease inhibitor, and alternate substrate for protease activity, respectively. All samples were stored at -70 °C until assayed.

VI. RESULTS

A. Preparation of SEB Toxoid Microspheres

During this reporting period, we prepared the SEB toxoid microspheres for use in the rhesus monkey studies. The initial tatch of microspheres (Composite F787-074-00), that we prepared were too small (97.3% < 5.3 μ m in diameter). The in vitro release profile also indicated too rapid a release of the SEB toxoid. These data are shown in Figure 1 (Appendix B). A second composite batch of microspheres was then prepared, F787-110-00. This batch consisted of microspheres that were more appropriately sized and had more favorable in vitro release characteristics. These data are illustrated in Figure 2.

One final batch of SEB toxoid microspheres was prepared. This batch (Composite F787-124-00), was prepared to ensure that a sufficient supply of microspheres would be on hand for the monkey studies. The in vitro release profile and size distribution for this batch of microspheres are shown in Figure 3.

All of the SEB toxoid microspheres prepared during this reporting period are described in Table 1. Size distributions for these microspheres are summarized in Table 2.

B. Immunization Results of Monkey Study EX-Rh-100

The preliminary rhesus monkey study, EX-Rh-100, was designed to provide a bridge to the existing data regarding the ability of IM immunization with alum-precipitated SEB toxoid and oral immunization with high doses of toxoid in solution to raise circulating anti-SEB toxin antibodies. This experiment also provides information about the levels of anti-toxin in lung secretions after immunization by these two methods. The toxoid forms, doses and the timing of the doses were selected to approximate the studies of Bergdoll (37).

For Study EX-Eh-100, three groups of two monkeys each were immunized as follows:

- Group 1: Monkeys 89 Bl09 and 89 B093 Normal controls
- Group 2: Monkeys 89 B106 and 89 B115 100 µg of SEB toxoid on alum, administered by IM injection on Days 0, 49, and 105.
- Group 3: Monkeys 89 B043 and 89 B103 10 mg of SEB toxoid in 0.7% bicarbonate, administered by oral gavage on Days 0, 49, and 105.

Plasma samples were collected on Day 0 and at 7-day intervals through Day 98. BAW samples were collected on Day 0, 28, 49, 77, and 98.

1. Plasma anti-SEB toxin responses

End-point titration in ELISAs employing solid-phase adsorbed SEB toxin confirmed the absence of toxin-reactive antibodies of any isotype in the probleeds of all the monkeys employed in this study. The control monkeys, 89 Bl09 (Figure 4) and 89 B093 (Figure 5), remained serologically negative in all isotypes to SEB toxin at the lowest tested dilution of plasma (1:50) throughout the period of this study.

The two monkeys immunized with SES toxoid precipitated on an alum slurry, 89 B106 (Figure 6) and 89 B115 (Figure 7), responded with the production of SEB toxin-specific circulating antibodies in each of the three isotypes. IgH anti-SEB toxin titers reached maximal levels 14 days after the primary immunization in both monkeys, and were detectable throughout the primary responses. Following boosting on Day 49, Monkey 89 B115 demonscrated a secondary rise in IgH anti-toxin, while Monkey 59 B106 did not. However, late in the course of the secondary response, IgH anti-toxin titers in both monkeys became undetectable. The bulk of the circulating anti-SEB toxin responses in the monkeys receiving alum precipitated toxoid were of the IgC class. Maximal IgG anti-SEB toxin titers were achieved in both monkeys by Day 28. Monkey 89 BlO6 mounted an earlier and more vigorous IgG response which reached a peak titer of 102,400 on Day 21, while Monkey 89 B115 exhibited a peak titer of 25,600 which was not reached until Day 28. However, both monkeys exhibited a secondary response which achieved a maximal titer of 409,600 on the 14th day following boosting. Circulating IgA anti-SEB toxin antibodiss were only observed in low titers after immunization with the alua-precipitated toxoid. One monkey, 89 Bl06, produced a low and transient IgA response which coincided with the peak of the secondary response. The other monkey, 89 \$115, produced a low IgA response across the later portion of the primary response, but no response was detectable after boosting. Thus, the two monkeys injected with alumprecipitated toxoid responded somewhat differently to the primary immunization. Monkey 89 B106 responded more rapidly and produced 4- to 8-fold higher levels of IgN and IgG anti-toxin than Monkey 89 Bll5. However, following secondary immunization, both monkeys achieved the same circulating titer of IgG anti-toxin.

In contrast to the high titers of circulating anti-toxin antibodies achieved by the systemic injection of $100-\mu g$ doses of toxoid on alums, the two monkays receiving 10-mg doses of toxoid orally exhibited only sporadically detectable circulating anti-toxin antibodies. As shown in Figure 8, Nonkey 89 B043 produced low, but clearly detectable, IgA anti-toxin antibodies on Day 28 after the primary administration and 7 days after the secondary administration. Monkey 89 B103 also exhibited a low circulating IgA response (Figure 9) 7 days after the secondary administration and an additional transient IgH response on Day 35 and 42 following the secondary administration. Overall, the administration of even high doses of SEB toxoid in solution appears to have been virtually ineffective at the induction of circulating anti-SEB toxin antibodies.

2. Bronchial-alveolar wash anti-SEB toxin responses

Assays of the BAW fluids obtained from the monkeys in EX-Rh-100 revealed that SEB-specific antibodies could only be detected in the samples of the lung fluids obtained from the two monkeys which had received IM immunization with the alum-precipitated toxoid (Table 3). These antibodies were restricted to the IgG isotype and were not detected until Day 49. They were present at the highest level (titers of 80) on Day 77, and fell to titers of 10 on Day 98. Following a tertiary IM immunization on Day 105, these monkeys resisted aerosol challenge with SEB toxin.

C. Immunization Results of Monkey Study EX-Rh-101

Experiment EX-Rh-101 was designed to investigate the efficacy of immunizing with SEB toxoid microspheres. More specifically, 10 monkeys were employed in testing SEB toxoid microspheres prepared with a 53:47 DL-FLG excipient. These microspheres, composite Batch F787-110-00, contained 0.41 wt % SEB toxoid and were 1 to 10 μ m in diameter. One of the 10 monkeys was used as a nonimmunized control, and the remaining 9 monkeys were administered SEB toxoid microspheres in the nine possible combinations of primary and secondary immunizations using the IM, oral and intratracheal (IT) routes of administration. A summary of the immunization regimens is given in Table 4.

The rhesus monkeys were bled for plasma samples on Day 0, and every 7 days thereafter through Day 98. BAW samples were obtained on Days 0, 28, 49, 77, and 98. All samples were tested for SEB toxin specific antibodies of the IgM, IgG, and IgA isotypes by end-point titration.

1. Plasma anti-SEB toxin responses following primary immunizations

The nonimmumized control monkey (89 5088) did not exhibit detectable p issue anti-SEB toxin antibodies of any isotype at any time point tested (Figure 10).

a. Primary IN immunization (Monkeys 89 B096. 89 B161. and 89 B210)

Each of the monkeys that were administered a primary IM immunization with 100 μ g of microencapsulated SEB toxoid mounted a brisk circulating anti-toxin response which was observed in all isotypes, but which was predominantly IgG. These immune responses are graphically illustrated in Appendix B as indicated:

Honkey	Figure	
89 B096	11	
89 B161	12	
89 B210	13	

The responses by Monkeys 89 B161 and 89 B210 were similar in that their maximal primary IgG anti-toxin titers of 102,400 were attained on Day 28 and remained at this level through Day 49 when booster immunizations were administered. In contrast, Monkey 89 B096 mounted a plasma IgG response which arose more slowly, but which staadily increased through Day 49 to a titer of 409,600. Thus, each of the monkeys immunized with 100 μ g of encapsulated SEB toxoid attained primary IgG anti-toxin levels which equalled or exceeded the peak primary IgG anti-toxin titer reached by the monkeys making the highest response to immunization with 100 μ g of SEB toxoid precipitated on alum. In addition, the monkeys immunized with the microencapsulated toxoid maintained their anti-toxin levels better, and by Day 49 their plasma IgG anti-toxin titers were 2 to 16 times higher than the monkeys immunized with the alum-precipitated toxoid.

b. Primary IT immnization (Monkeys 89 5079. 89 5001. and 89 5116)

Primary IT immunization with 100 μ g of microencapsulated SEB toxoid induced a circulating anti-SEB toxin response in each of the three monkeys. These responses are given in Appendix B as follows:

Monkey	Figure	
89 B079	14	
89 BOO1	15	
89 B116	16	

In two of the three monkeys (89 B079 and 89 B001), the response was exclusively of the IgM isotype. The third monkey, 89 B116, produced an early IgM response which was followed by an IgA response. The IgM responses were quite individual in nature: Monkey 89 B079 produced an early, high response, Monkey 89 B001 produced a late response of intermediate level, and Monkey 89 B166 produced an early but low response. None of the IT-immunized monkeys produced a measurable plasma IgG anti-toxin response at any of the time points examined.

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Note: The complete absence of a circulating IgG response, and the presence of only a low IgA response in one of the monkeys, is in contrast to the extremely vigorous responses observed in mice following IT immunization with SEB toxoid microspheres (reference Third Annual Report, page 12). This prompted us to review the menner in which the primary IT immunization of these three monkeys had been performed. After review, we realized that the microsphere suspension had been instilled through a catheter passed deep into one lung. This likely delivered the microspheres to only one segment of one lobe. In contrast, the IT immunizations employed in the mouse studies involved the instillation of the suspension into the traches. This technique resulted in a more uniform deposition in both the upper and lower respiratory tract. All subsequent IT immunizations of the rhesus monkeys (beginning with the EX-Rh-101 booster immunizations) were performed by passing the catheter just beyond the end of the intratracheal tube. Significantly improved responses have resulted, as shown in the EX-Rh-101 monkeys receiving secondary immunizations via the IT route (data discussed below).

c. Primary oral immunization (Monkeys 89 8060, 89 8064, 89 8034)

Following oral immunization with the microencapsulated SEB toxoid, one of the monkeys (89 B060) did not exhibit a demonstrable plasma anti-toxin response. However, the other two monkeys (89 B064 and 89 B034) did produce circulating antibodies. The responses are illustrated in Appendix B as indicated:

Monkey	Figure
69 B060	17
89 8064	15
89 8034	19

Monkey 89 B034 produced an IgM response which was present at a titer of 800 on Days 28 through 49. In contrast, Monkey 89 B064 responded with an early IgM response which peaked on Day 14 and fell to an undetectable level on Day 42. In addition, this monkey produced an IgG anti-toxin response beginning on Day 28 and peaking on Day 4 at a titer of 1,600. Although 2 of the 3 monkeys immunized with the microencapsulated SEB toxoid produced clear responses in the plasma, none exhibited detectable IgA antitoxin activity.

2. Plasma anti-SEB toxin responses following secondary innumizations

a. Secondary impunizations of IN-primed monkeys

As a group, the monkeys administered primary IM immunizations with microencapsulated toxoid, and boosted with the SEB toxoid microspheres either systemically or by the two mucosal routes, exhibited good levels of circulating anti-toxin antibodies. The orally-boosted monkey (89 B096, Figure 11) maintained an IgG antibody level which was of the same order of magnitude as monkeys administered primary and secondary immunizations with toxoid precipitated on alum. The IT-boosted monkey (89 B161, Figure 12)

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and the IN-boostad monkay (49 M10, Figure 13) produced antibadise at substantially higher lavels that these induced by two immutations with the alum-productated teneid. In addition, the monkays boosted with the microsphere formulation by either monaal route sublited directating IgA anti-toxic responses which were of a long duration. A point workly of mention in the consideration of the lovels of circulating IgA anti-toxic induced through the mussal boosting of IN-prime monkays. Is the level of plasma IgC anti-toxic antibodies that are present in the plasma accepter. The exceptionally high IgC antibody levels (siters of all, 500 to 1, 515, 400) must successfully compate for the solid-phase antigms in the sectors. meaking the true IgA anti-toxic level. Thus, the plasma IgA responses and the IgH responses, is these monkays must be considered a minimum sectors.

(1) Oral boarding of Bearings Robert 11 1014

No invesse is the level of circulating applements antibadies was observed following and administration of i ap of microssisponiated SEB consid to one of the IN-primed membrys (40 5056, Figure 11). The Bay 49 IgO anti-taxin titar of 409,600 foll to 31,70 by Bay 70 and then remained at that level through Day 91. It is difficule to descende from these dates if the plateau in circulating IgO antibadies cheeved on Bay 90 through 51 represents maintenance through offersive orbit heasting of mouse bis of the antibady decay after the primery IN immulsation. However, this membry, restring the IN-offer is a detectable plasme IgA anti-conte is one is throughout the secondary response.

(2) IT hearting of Illentined Mechan-II-III

It beesting of previously IN-immuted Nonicey 89 6161 with 100 mg of microencepsulated SKE tonuid atimulated a proppy and vigorous lasterse in the elreulating IgU anti-tonic antibudy level. As shown in Pigure 19, a titer of 1,638,400 was achieved on bay 63. This high titer uss maintained through the time which the secondary responses were followed, and had only decreased to a titer of 619,200 on bay 98. As in the case of the bookey administered the IN-oral lammination schedule, this monkey schibited eirculating IgA anti-toxin antibudies through bay 98 when the anxiyois was terminated.

The unseptionally good beening obtained by the if rouse in shis member, and in the other two members which were benared by the if rouse (19 A001, Figure 15 and 49 8064, Figure 18), indicate the the charge in the method of 1T instillation, discussed previously, has significantly improved the effectiveness of this immutation route.

(1) IN beauting of IN original Montay At Ailo

IN boosting of previously IN-impunited Monkey 69 5810 stimulated approximately a 16-fold rise in the plasme igo anti-texts isvel from a titer of 103,400 on Day 49 when the boost was administered, to a titer of 1,638,400 on Day 43 (Pigura 13). A plateau titer of \$19.300 was plateau on Days 70 through 98. In contrast to the IN-immunized monkeys which restived a montrast to the IN-immunized monkeys which restived a mostal booster immunizations, this monkey did not achibit a datactably plasme IgA anti-toxin response after boosting

b. <u>Sacondary immunizations of IT-primed monkeys</u>

For the reasons discussed previously, the monkeys receiving a primary IT immunization with the microencapsulated SEB toxoid were poorly primed. Havever, both the IT and IM routes proved to be effective boosters in these monkeys. The levels of IgG anti-toxin antibodies produced by these secondary immunizations were greater than those obtained in 2 of the 3 monkeys receiving a primary IN immunizations.

(1) Oral boosting of IT-primed Monkey 89 B072

Oval buosting of this IT-primed monkey did not result in the induction of a secondary circulating anti-toxin response that was detectable in any isotype, at any time point tested (Figure 14). However, there is evidence that the oral boosting did stimulate the appearance of specific anti-toxin antibodies at low level in the BAW fluid samples of this monkey.

(2) IT boosting of IT-primed Monkey 89 B001

IT hussting of Monkey 89 BOOL, which had received a primary IT immunisetion, induced the appearance and a rapid rise in the level of plasma IgG setioniss (Figure 15). On experimental Day 70, these antibodies reached a titer of 204,800, and this level was maintained through Day 98.

(3) IN housting of IT-primed Monkey 89 B116

Following an IM boost with SEB toxold microspheres, IT-primed Monkey #9 Blis anhibited a rapid rise in both IgM and IgG anti-toxin antibodies in the SiFoulation (Figure 16). With the exception of the IgM component, the Following by this morkey to the IT-IN regimen was quite similar to that seen in the general administered the IT-IT schedule. The IgG anti-toxin titer was 204, 500 on Day 98 of the study.

8. Essendary incunizations of orally-primed monkeys

The montesys administered a primary oral immunization with the microenseperates a Sta compile were clearly responsive to secondary immunization via systeple (IM) and muchanal (IT) routes. The 204,800 titer of circulating is anti-texts achieved by both IT (Monkey 89 B064) and IM (Monkey 89 B034) weessing elearly shows that systemic tolerance has not been induced through ersi immuliastion with the alcroancapsulated toxoid. That both these assistants achieved sirculating titers greater than those induced in 2 of the 3 asakeys receiving primary IM immunization with the microencapsulated SEB tangid suggests that a degree of systemic priming has taken place. Somewhat sugging is the total lack of a response in Monkey 89 B060 to both primary and ascendary oral immunization with the SEB toxoid microspheres, when two anhar monkeys made clear responses to a single oral immunization. Fauen segether with the fact that Monkey 89 B079 was clearly nonresponsive we were howevering, these data suggest a threshold effect in which only some of the muchays are excelving an effective amount of vaccine. To what estent genetic factors may play a role is not known.

(1) Ural boosting of orally-primed Monkey 89 8060

Monkey 89 B060 failed to produce a detectable plasma response following either the primary or secondary oral immunization with microencapsulated SEB toxoid (Figure 17).

(2) IT boosting of orally-primed Monkey 89 B064

IT boosting of Monkey 89 B064, which had made the greatest plasma response to a primary oral immunization, resulted in an immediate rise in the levels of IgG anti-toxin antibodies. This response reached a titer of 204,800 on Day 56 (Figure 18). The titer remained steady at this level through Day 98. This indicates that the modification to the IT immunization procedure provided an effective immunization with the microencapsulated SEB toxoid.

(3) IN boosting of orally-primed Monkey 89 B034

This monkey, which had also responded to the primary oral immunization, produced a clear response to secondary IM immunization (Figure 19). Unlike the orally-primed monkey that was boosted by the IT route, this monkey's titer of IgG anti-toxin rose progressively through Day 77 to a titer of 204,800. On Experimental Days 77 through 98, the plasma IgG antitoxin level remained steady.

3. BAW anti-SEB toxin responses from EX-Rh-101

At no time did any of the samples of BAW fluids from the control, nonimmunized monkey (89.8088) contain detectable anti-SEB toxin antibodies. These results are presented in Table 5, which contains all of the BAW fluids analysis for EX-Rh-101.

The BAW fluids from the monkeys which received a primary IM immunizations with microencapsulated SEB toxoid (89 B096, 89 B161 and 89 B210) were without anti-toxin activity except for the Day 49 sample from Monkey 89 B096. This sample contained specific anti-SEB toxin IgG antibodies at a titer of 80. In contrast, 2 of the 3 monkeys administered a primary immunization by the IT route (89 B079, 89 B001 and 89 B116) exhibited IgH anti-toxin antibodies in their BAW fluids samples at a titer of 40. (Note: These are the monkeys which had received the primary IT immunization as a bolus deposition of the SEB toxoid microspheres deep into one lung lobe. The method of IT immunization was altered for later immunizations. The monkeys must be considered poorly primed by this immunization). The monkeys administered a primary oral immunization (89 B060, 89 B064 and 89 B034) did not mount a primary immune response which was detectable in their BAW fluids, in any isotype, at the time points tested.

Booster immunization of the IM-primed monkeys, regardless of the route of administration, resulted in enhanced levels of IgG anti-toxin antibodies in the BAW fluids. However, only the monkey boosted via the JT route showed a detectable IgA anti-toxin response.

Booster immunization of the IT-primed monkeys by the IT route (39 B001) and the IM route (39 B116) resulted in the appearance of IgG anti-toxin antibodies in the BAW fluids, while the BAW fluids from the orally-boosted monkey continued to contain IgM.

The orally-primed monkeys which were boosted by the oral (89 B060) and IM (89 B034) routes failed to produce detectable responses in their BAW fluids. However, IT boosting of orally-primed Monkey 89 B064, resulted in both IgG and IgA anti-toxin antibodies that were still detectable on Day 98. The antibody titers were 80 and 20 for the IgG and IgA respectively.

D. Immunization Results of Monkey Study EX-Rh-102

The immunization schedule for the monkeys in this experiment is the same as that for the previous study. This schedule is summarized in Table 6.

1. Plasma anti-SEB toxin responses following primary immunizations

The nonimmunized, control monkey (90 8001) did not exhibit detectable plasma anti-SEB toxin antibodies of any isotype, at any time point tested. This is illustrated in Figure 20.

a. Primary IM immunization (Nonkeys 90 B002, 90 B003 and 90 B010)

Each of the monkeys which were administered a primary immunization with 100 μ g of microencapsulated SEB toxoid mounted a brisk circulating anci-toxin antibody response. This response was primarily in the IgG isotype, but also in the IgM isotype. These immune responses are graphically depicted in the indicated figures:

Monkey	Figure
90 BOO2	21
90 BOO3	22
90 BO10	23

The responses by all the monkeys receiving a primary IM immunization with the SEB toxoid microspheres were similar in that their maximal primary IgG anti-toxin titers of 51,200 to 102,400 were attained on Day 21 and remained at this level through Day 49 when booster immunizations were administered. Thus, as in Study EX-Rh-101, each of these monkeys attained primary IgG anti-toxin levels which equal'ed or exceeded the highest peak primary IgG anti-toxin titer achieved by a monkey immunized with an equal dose of SEB toxoid precipitated on alum. In addition, the monkeys immunized with the microencapsulated toxoid exhibited no decline in their Day 21 antibody titers.

b. Primary IT immunization (Monkeys 90 B023, 90 B009 and 90 B021)

It should be noted that the altered procedure for IT immunization was used for these primary immunizations. Microspheres containing 100 μ g of SEB toxoid, administered via the IT route, resulted in circulating anti-SEB toxin responses in all three monkeys. The plasma IgG titers (102,400 to 409,600) equalled or exceeded the responses induced by IM immunization with alum-precipitated or microencapsulated SEB toxoid. Figures showing the immune responses for these monkeys are as follows:

Monkey	Figure
90 8023	24
90 8009	25
90 8021	26

In addition to the strong IgG anti-toxin response induced via IT immunization with the microencapsulated SEB toxoid, this approach also induced circulating IgM responses which tended to peak early and fall to undetectable levels by Day 35. IgA anti-toxin antibodies were also detectable through Day 49 when the booster immunizations were administered.

c. Primary oral immunization (Monkeys 90 B014. 90 B005 and 90 B018)

Following oral immunization with SEB toxoid microspheres, all three of the monkeys produced circulating anti-SEB toxin antibodies. There were some differences in the responses, which are illustrated in the indicated figures.

Monkey	Figure
90 B014	. 27
90 8008	25
90 B018	29

Monkeys 90 B014 and 90 B018 produced IgM anti-toxin antibodies on Day 14 followed by the appearance of IgG antibodies on Day 21. However, Monkey 90 B014 exhibited these circulating IgG antibodies through Day 49, while the IgG response in Monkey 90 B018 declined to undetectable levels on Day 42. In contrast, Monkey 90 B008 produced circulating IgM and IgA antitoxin antibodies that were only detected on Day 35. Although none of these monkeys mounted high titers of circulating anti-toxin antibodies, when the results from this experiments are combined with those from Study EX-Rh-101, 5 of 6 monkeys receiving the microencapsulated SEB toxoid orally, have produced circulating antibody responses. This is in contrast to Study EX-Rh-100, in which neither of the two monkeys administered nonmicroencapsulated SEB toxoid exhibited any detectable response.

2. Plasma anti-SEB toxin responses following secondary immunizations

. Secondary immunizations of IM-primed monkeys

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As a group, the monkeys administered primary IM immunizations with microencapsulated SEB toxoid, and boosted either systemically or by the two mucosal routes, exhibited good levels of circulating anti-toxin antibodies. The orally-boosted monkey, 90 B002 (Figure 21), maintained an IgG antibody level which was of the same magnitude as monkeys administered primary and secondary immunizations with SEB toxoid precipitated on alum. The ITboosted monkey, 90 B003 (Figure 22) and the IM-boosted monkey, 90 E010 (Figure 23) produced antibodies at substantially higher levels than those induced by two immunizations with the alum precipitated toxoid.

In addition, the monkey boosted with the SEB toxoid microspheres by the IT route, exhibited circulating IgA anti-toxin responses which were of a long duration. A point worthy of mention in the consideration of the levels of circulating IgA anti-toxin induced through the mucosal boosting of IM-primed monkeys, is the level of plasma IgG anti-toxin which is present in the plasma samples. The exceptionally high IgG antibody levels (titers of 819,200 to 1,638,400) must successfully compete for the solidphase antigen in the assays, masking the true IgA anti-toxin level. Thus, the plasma IgA and IgH responses in these monkeys must be considered minimum estimates.

(1) Oral boosting of IM-primed Monkey 90 B002

Following the oral administration of 1.0 mg of microencapsulated SEB toxoid to this IM-primed monkey, a booster response was evident in the IgM isotype on Days 56 through 70 (Figure 21). Although the circulating IgG anti-toxin titer fell from its Day 63 high, it remained on a plateau from Day 77 through Day 98. This suggests that the oral booster may have helped maintain the circulating antibody level.

(2) IT boosting of IN-primed Monkey 90 8003

An IT boost consisting of 100 μ g of microencapsulated SEB toxoid resulted in a prompt and vigorous response in this monkey. As shown in Figure 22, the circulating IgG anti-toxin antibody level increased to a titer of 1,638,400 on Day 63. This high titer of IgG anti-toxin activity was maintained through the time which the secondary response was followed. The response had only decreased to a titer of 819,200 on Day 98. In addition, this monkey exhibited circulating IgA anti-toxin antibodies ten days after IT boosting. These IgA antibodies rose steadily to a titer of 1,600 on Day 77 and remained at this level though Day 98.

(3) IN boosting of IM-primed Monkey 90 B010

IM boosting of previously IM-immunized Monkey 90 B010 stimulated approximately a 16-fold rise in the plasma IgG anti-toxin level. As illustrated in Figure 23, the IgG anti-toxin antibody titers increased from 102,400 on Day 49 to 1,638,400 on Day 63. A plateau titer of 409,600 was present on Days 77 through 98. In contrast to the IM-immunized monkeys which received mucosal booster immunizations, this monkey did not exhibit a detectable IgA anti-toxin response following the booster immunization.

b. Secondary immunizations of I"-primed monkeys

The monkeys which received IT primary immunizations with microencepsulated SEB toxoid were very well primed. This demonstrated that the change in the IT installation procedure has been effective. Both the IT and IM routes proved to be very effective boosters in these monkeys. The longterm high levels of IgG antibodies in the monkeys receiving an oral booster suggests that an effective amount of vaccine was delivered by this route.

(1) Oral boosting of IT-primed Monkey 90 B023

Oral boosting of this IT-primed monkey did not result in the induction of a clearly demonstrable booster response (Figure 24). However, good maintenance of the circulating IgG anti-toxin levels are consistent with the delivery of an effective amount of vaccine.

(2) IT boosting of IT-primed Monkey 90 B009

Following the IT boost, a rapid rise in the levels of plasma IgG and IgA anti-toxin antibodies was detectable (Figure 25). The IgG antibodies reached a titer of 819,200 on Day 63 and remained at this level through the duration of the experiment. The IgA antibodies reached a peak titer of 6,400 at a point 10 days after the IT boost, and were present at elevated titers throughout the experiment.

(3) IN boosting of IT-primed Monkey 90 B021

After an IN boost with SEB toxoid microspheres, this IT-primed monkey exhibited a rapid rise in IgG anti-toxin antibodies in the circulation (Figure 26). The secondary response by this monkey was very similar to that seen in the monkey receiving two IM immunizations. The IgG anti-toxin titer rose to 1,638,400 on Day 63 and then remained at a plateau level of 204,800 through Day 98. However, this secondary response differs significantly from that exhibited by the IT-IT immunized monkey in that it completely lacks a detectable IgA response.

c. <u>Secondary izmunizations of orally-primed monkeys</u>

Each of the three monkeys administered a primary oral immunization responded with circulating anti-toxin antibodies. In addition, all were clearly responsive to secondary immunization via the systemic and mucosal routes. The titers of circulating IgG anti-toxin achieved by both IT (90 B008) and IM (90 B018) boosted monkeys indicated that systemic tolerance had not been induced through oral immunization with the microencapsulated SEB toxoid.

(1) Oral boosting of orally-primed Monkey 90 B014

In the first 21 days following oral boosting of the orally-primed monkey, a four-fold rise in the circulating IgG anti-toxin antibodies to a titer of 800 was observed (Figure 27). Once attained, this titer remained constant through Day 98. However, this immunization method did not result in the appearance of detectable IgA anti-toxin activity at any tested time point.

(2) IT boosting of orally-primed Monkey 90 B008

IT boosting of this monkey resulted in a rise in the levels of IgG anti-toxin antibodies which was evident 14 days after boosting. The increase continued through Experimental Day 77, when the titer reached 102,400. Thereafter the titer decreased to 25,600 on Day 98 (Figure 28).

(3) IN boosting of orally-primed Monkey 90 B018

This monkey exhibited a rapid response to a secondary IM immunization which was characterized by both IgM and IgG components (Figure 29). The IgM response was of short duration and had fallen to an undetectable level 28 days after boosting. In contrast, the IgG anti-toxin response steadily increased through Day 77 to a titer of 25,600 and remained at that level through Day 98.

3. BAW anti-SEB toxin responses from EX-Rh-102

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As in Study EX-Rh-101, the nonimannized control monkey in this experiment did not exhibit detectable anti-toxin antibodies of any isotype in any of the samples tested. Table 7 contains this data as well as BAW data from all the monkeys in Study EX-Rh-102.

Home of the monkeys administered a primary IN (90 B002, 90 B003 and 90 B010) or primary oral (90 B014, 90 B008 and 90 B018) immunization with the microencapsulated SEB toxoid produced an anti-toxin response that was detectable in the BAW fluids. In contrast, all the monkeys receiving a primary immunization via the IT route, responded with relatively high levels of IgA and IgG anti-toxin antibodies in their BAW fluids.

Bocster immunization of the IM-primed monkeys by the oral route (90 B002) and IM route (90 B010) was ineffective at stimulating an antitoxin response in the BAW fluids. However, boosting by the IT route (90 B003) induced appearance of IgG and IgA anti-toxin antibodies at titers of 12,800 and 320, respectively.

Each of the monkeys primed by the IT route continued to have measurable anti-toxin antibodies after boosting. The antibody levels declined, despite the boost in the orally-boosted monkey (90 B023). In the ITboosted monkey (90 B009), the IgA antibodies in BAW fluids increased and the IgG antibodies decrease after boosting. The converse was evident for the IM-boosted monkey (90 B021).

None of the orally-primed monkeys (90 B014, 90 B008 and 90 B018) mounted a detectable response in the BAW fluids regardless of the route of booster immunization.

VII. DISCUSSIONS AND CONCLUSIONS

During this year of the contract, we obtained a great deal of knowledge about the immunization of rhesus monkeys with SEB toxoid. A preliminary study was performed to confirm the existing data regarding the effectiveness of systemic (IN) immunization with alum precipitated SEB toxoid and oral immunization with high doses of toxoid in solution, to raise circulating anti-SEB toxin antibodies. Additionally, several studies involving the immunization of rhesus monkeys with SEB toxoid microspheres have also been performed.

The preliminary monkey experiment yielded the following results: Immunization with alum-precipitated SEB toxoid did result in the appearance of circulating IgG antibodies, which were detected in low levels in the BAW fluids. Oral immunization with SEB toxoid solution induced only the transient appearance of low levels of IgM and IgA anti-SEB toxin antibodies in circulation.

Results from the first two studies (EX-Rh-101 and EX-Rh-102) in which rhesus monkeys were immunized with microencapsulated SEB toxoid are included in this report. These data suggest that an IM primary immunization followed by an IT or oral secondary immunization provides the highest antibody response and best level of protection. It has been reported to us that the monkeys receiving an IM primary and IT secondary immunization in both studies survived aerosol challenge. In addition, the monkey receiving an oral primary and IT secondary immunization from Study EX-Rh-101 and the monkey receiving both an IT primary and IT secondary immunization in Study EX-Rh-102 also survived the aerosol challenge. Official documentation of these results has not been received by Southern Research or the University of Alabama in Birmingham.

These results are very encouraging as not only have high antibody titers been achieved in the rhesus monkeys, but as indicated above, several of the monkeys have been shown to be protected from aerosol challenge.

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IX. AGENOVLEDOMENTS

44 Georgi D. Hudson, Assessch Chemical Technician, prepared and chossossistical die SBR testeld minrespheres. Mr. Orlan B. Finch, Research decisions, is responsible for maintaining the mouse-breeding colony, saidel hemiling, tissue procurement, and respont preparation. Co. Chasicope K. Jones, Assessch Assistant II, is responsible for objects Scheling, Rissospilet antioning, immunchistochemical procedures. End Exceptional envisory production and purification.

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APPENDIX A

TABLES

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TABLE 1. SEB MICROSPHERES PREPARED DURING THIS REPORTING PERIOD

	Batch	Yield.	Core los	ding. Eb	efficiency. Z of
Batch"	size, g	h4	Theoretical	Actual	theoretical
787-046-00	1.01	68.8	1.0	06.0	30.0
787-052-00	1.01	62.0	1.0	0.35	35.0
787-055-00	2.02	80.6	1.0	0.42	42.0
787-071-00	2.02	77.3	1.0		
787-073-00	2.02	79.3	1.0	8 9 1	
787-074-00	:		1.0	0.48	48.0
787-082-00	2.02	81.2	1.0		
187-089-00	2.02	75.9	1.0	8	:
187-091-00	2.02	1.11	1.0	t e	•
00-200-00	2.02	79.9	1.0	•	
787-095-00	2.02	78.7	1.0	6	:
787-110-00 ⁴	•	•	1.0	0.41	41.0
/87-117-00	2.02	:	1.0	•	•
787-119-00	2.02	:	1.0	;	
787-121-00	2.02	:	1.0	:	
187-123-00	2.02	1	1.0	•	. 1
187-124-00*	•	78.0	1.0	0.35	35.0

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"All microspheres were prepared with a 53:47 DL-PLG excipient. Lot BPI-043-62-1.

b*..* = Not determined.

"Batch F787-074-00 is a composite of Batchas F787-071-00 and F787-073-00.

^dBatch F787-110-00 is a composite of Batches F787-082-00, -089-00, -091-00, -093-C0, and -095-00. "Batch F787-124-00 is a composite of Batches F787-117-00, -119-00, -121-00, and -123-00.

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TABLE 2. SIZE-DISTRIBUTION DATA FOR SEB MICROSPHERES PREPARED DURING THIS REPORTING PERIOD

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		y volume. 3 of microsphe	res
Batch	Less than 4.6 µm in diameter	From 4.6 to 9.6 µm in diameter	Greater than 9.6 µm in diameter
F787-046-00		4 3	0.0
F787-052-00	63.2	33.8	3.0
F787-055-00	87.2	12.8	0.0
F787-074-00	91.7	\$.3	0.0
\$787-082-00	۲۲ ۲	40.0	6.5
F787-110-00	63.7	35.7	0.6
F787-124-00	73.2	25.5	1.3
F787-074-00 F787-082-00 F787-110-00 F787-124-00	91.7 53.5 63.7 73.2	\$.3 40.0 35.7 25.5	

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TABLE 3. DAW ANTI-SED TOXIN RESPONSE TO SEB TOXOID, EX-RH-100

Monkey	Imunization		DAV 28		Brenchi	VIS-IS VAU	Colar V	ash anti	Dav 77	oxin ti	ter	Dav 98	
number	schedule	N	IEC	IEA	1cH	166	IEA	1eM	Je C	IEA	M	lec	IEA
89 B109	Control	<10	<10	<10	<10	<10	<10	<10	<10	<10	<10	<10	<10
89 B093	Control	<10	<10	<10	<10	01⊳	<10	01 >	410	<10	40	01≥	40
69 B 106	1M- 1H*	<10	<10	<10	<10	40	40	<10	09	<10	<10	10	<10
89 8115	HI-HI	<10	<10	<10	<10	20	<10	<10	0	<10	<10	10	<10
89 8043	0-0	<10	<10	<10	<10	40	<10	<10	<10	<10	01≥	40	<10
89 B103	0-0	<10	<10	<10	<10	40	<10	<10	<10	<10	<10	410	<10
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A-3

"Anti-toxin titer determined by end-point titration.

^bNot-**immunize**d.

^c Immunized on Days 0 and 49 by IM injection of 100 μg of SEB toxold on aluman.

^d Immunized on Days 0 and 49 by oral gavage of 10 mg of SEB toxoid in a 0.7% bicarbonate solution.

PROPRIETARY INFORMATIO

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	Primary in Day	v O	Secondary is Day	Annization
Monkey number	Route	Dose	Route	Dose
89 B088	None	None	None	None
89 3096	IM	100 µg	Oral	1 mg
89 B161	IM	100 µg	IT	100 ##
89 B210	IM	100 µg	IM	100 µg
89 8079	IT	100 µg	Oral	1 mg
89 BOO1	IT	100 µg	IT	100 µg
89 B116	IT	100 µg	IM	100 µg
89 B060	Oral	1 mg	Oral	1 mg
89 BO64	Oral	1 ng	IT	100 µg
89 8034	Oral	1 ag	IM	100 µg

TABLE 4. SUMMARY OF INMUNIZATIONS FOR STUDY EX-RH-101

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TABLE 5. BAW ANTI-SEB TOXIN RESPONSE TO SEB TOXOID MICROSPHERES, EX-RH-101

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Mor	ıkey	imuniza tion		Day 28		Bronch	Dav 45	Ceoler V	ash ant	Dav 7	toxin El	ter	Dav 96	
an i	ber	. schedule	<u>Та</u>	IEC	IEA	Na1	140	IcA	Leff	Ê	IEA	Mal	US I	1sA
89	8993	Control ^b	<10	40	<10	<10	<10	<10	<10	410	<10	<10	<10	<10
89	B096	0-WI	<10	<10	<10	<10	80	<10	20	1260	<10	<10	<10 <10	₹10
89	8161	- TI-MI	<10	<10	<10	40	410	40	<10	320	9	<10	320	64
89	B 210	NI - NI	<10	<10	<10	<10	<10	<10	<10		<10	<10	04	50
69	B079	11-0	40	40	<10	- 4	<10	₹10	40	<10	40	40	40	<10
89	B001	11-11	<10	<10	<10	40	<10	<10	40	20	<10	9	<10	66
83	8116 8	11 - IN	<10	<10	<10	<10	<10	<10	<10	20	<10	40	20	<10
89.	B060	0-0	<10	<10	<10	<10	€10	<10	<10	₹10	40	<10	<10	<10
89	B 064	11-0	01>	<10	<10	<10	₹10	<10	97	320	9	<10	8	20
8 9	B 034	WI-O	<10	<10	01 20	410	<10	≺10	<10	<10	<10	<10	<10	<10

PROPRIETARY INFORMATION (COMPLETE PAGE)

^b Not - **immunized**.

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"Anti-toxin titer determined by end-point titration.

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	Primary in Day	munization 7 0	Secondary in	munization
Monkey number	Route	Dose	Route	Dose
90 8001	None	None	None	None
90 8002	IM	100 µg	Oral	1 mg
90 8003	IH	100 µg	IT	100 µg
90 8010	IN	100 µg	IM	100 µg
90 8023	IT	100 µg	Oral	1 mg
90 8009	IT	100 µg	IT	100 µg
9 0 8021	IT	100 µg	IM	100 µg
90 8014	Oral	l ag	Oral	1 mg
90 8008	Oral	1 mg	IT	100 µg
90 B018	Oral	1 mg	IM	100 µg

TABLE 6. SURGARY OF INDUNIZATIONS FOR EXPERIMENT EX-RH-102

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TABLE 7. BAW ANTI-SEB TOXIN RESPONSE TO SEB TOXOID MICROSPHERES, EX-RH-102

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Honkey	Immulzation		Day 2			Day 4			Day 7.	~		Day 9	8
number	schedule	H	IEG	IEA	Hal	1 EG	IEA	Ien	IEG	IEA	LeH	Irc	E
1008 06	Control*	<10	<10	<10	<10	610	<10	<10	40	<10	<10	40	4
90 B 002	0-WI	<10	<10	<10	<10	015	<10	<10	<10	40	<10	~10	ž
00 B003	IN-IT	<10	40	<10	<10	61	₹0	<10	320	160	<10	1260	32(
0 5010	NI-NI	<10	<10	<10	<10	410	<10	40	<10	<10	<10	01> .	Ă
90 B023	0-11	10	1280	160	<10	320	50	<10	50	40	<10	20	Ā
00 B009	11-11	<10	640	94	<10	1280	04	<10	973	320	<10	320	16(
0 8021	11-11	<10	160	8	61 0	160	0	<10	320	30	<10	320	Ă
00- B014	0-0	40	₹10	<10	410	10	<10	<10	<10	<10	<10	410	₹.
8008 O	0-11	<10	<10	10	<10	10	<10	01≥	<10	<10	<10	<10	\$
90 B018	0- IK	<10	<10	<10	<10	<10	<10	<10	410	<10	<10	<10	¥,

PROPRIETARY INFORMATION (COMPLETE PAGE)

^bNot-**immized**.

APPENDIX B.

FIGURES

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Figure 1. Cumulative in vitro release of SEB toxoid (above) and microsphere size distribution (below): Batch F787-074-00.

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Figure 2. Cumulative in vitro release of SEB toxoid (above) and microsphere size distribution (below): Batch F787-110-00.

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Figure 3. Cumulative in vitro release of SEB toxoid (above) and microsphere size distribution (below): Batch F787-124-00.

8-3





B-4



Figure 5. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B093: Non-immunized control.

B-5

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Figure 6. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 8106: Immunized by IM injection on Days 0 and 49 with 100 micro rams of SEB toxoid precipitated on Num.

8--6



Figure 7. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B115: Immunized by IM injection on Days 0 and 49 with 100 micrograms of SEB toxoid precipitated on alum.

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Figure 8. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 BO43: Immunized by oral gavage, on Days 0 and 49 with 10 mg of SEB toxoid in 5.0 mL of 0.7% bicarbonate.

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Figure 11. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B096: Primary immunization (Day 0)—100 micrograms microencapsulated SEB toxoid by IM injection. Secondary immunization (Day 49)—1.0 mg of microencapsulated SEB toxoid by oral gavage.

B-11



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Figure 12. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B161: Primary immunization (Day 0)---100 micrograms microencapsulated SEB toxoid by IM injection. Secondary immunization (Day 49)---100 micrograms of microencapsulated SEB toxoid by IT instillation.

B- 12



Figure 13. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B210: Primary immunization (Day 0)—100 micrograms microencapsulated SEB toxoid by IM injection. Secondary immunization (Day 49)—100 micrograms of microencapsulated SEB toxoid by IM injection.

8-13



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Figure 14. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B079: Primary immunization (Day 0)—100 micrograms microencapsulated SEB toxoid by IT instillation. Secondary immunization (Day 49)—1.0 mg of microencapsulated SEB toxoid by oral gavage.

B-14



Figure 15. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 BOO1: Primary immunization (Day O)—100 micrograms microencapsulated SEB toxoid by IM injection. Secondary immunization (Day 49)—100 micrograms of microencapsulated SEB toxoid by IT instillation.

9-15



Figure 16. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B116: Primary immunization (Day 0)—100 micrograms microencapsulated SEB toxoid by IT instillation. Secondary immunization (Day 49)—100 micrograms of microencapsulated SEB toxoid by IM injection.

B- 16



Figure 17. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B060: Primary immunization (Day 0)—1.0 mg microencapsulated SEB toxoid by oral gavage. Secondary immunization (Day 49)—1.0 mg of microencapsulated SEB toxoid by oral gavage.

B-17



Figure 18. Plasma IgM, IgG, and IgA anti-SSB toxin there obtained from rhesus Monkey 89 B064: Primary immunization (D++ 0)-+ 1.0 mg microencapsulated SEB toxoid by oral gavage. Secondary immunization (Day 49)---100 micrograms of microencapsulated SEB toxoid by IT instillation.

B- 18

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Figure 19. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 89 B034: Primary immunization (Day 0)—1.0 mg microencapsulated SEB toxoid by oral gavage. Secondary immunization (Day 49)—100 micrograms of microencapsulated SEB toxoid by IM injection.

B- 19





B-20



PLASMA ANTI-TOXIN TITER



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Figure 22. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 90 B003: Primary immunization (Day 0)—100 micrograms microencapsulated SEB toxoid by IM injection. Secondary immunization (Day 49)—100 micrograms of microencapsulated SEB toxoid by IT instillation

B--22




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Figure 24. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 90 B023: Primary immunization (Day 0)—100 micrograms microencapsulated SEB toxoid by IT instillation. Secondary immunization (Day 49)—1.0 mg of microencapsulated SEB toxoid by oral gavage.

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B-25

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Figure 26. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 90 BO21: Primary immunization (Day 0)—100 micrograms microencapsulated SEB toxoid by IT instillation. Secundary immunization (Day 49)—100 micrograms of microencapsulated SEB toxoid by IM injection.

3-26

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Figura 27. Plasma igM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 90 B014: Primary immunization (Day 0)—1.0 mg microencapsulated SEB toxoid by oral gavage. Secondary immunization (Day 49)—1.0 mg of microencapsulated SEB toxoid by oral gavage.

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Figure 29. Plasma IgM, IgG, and IgA anti-SEB toxin titers obtained from rhesus Monkey 90 8018: Primary immunization (Day 0)-1.0 mg microencepsulated SEB toxoid by oral gavage. Secondary immunization (Day 49)-100 micrograms of microencepsulated SEB toxoid by IM injection.

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