From mouse to man: safety, immunogenicity and efficacy of a candidate leishmaniasis vaccine LEISH-F3+GLA-SE

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Key antigens of Leishmania species identified in the context of host responses in Leishmania-exposed individuals from disease-endemic areas were prioritized for the development of a subunit vaccine against visceral leishmaniasis (VL), the most deadly form of leishmaniasis. Two Leishmania proteins—nucleoside hydrolase and a sterol 24-c-methyltransferase, each of which are protective in animal models of VL when properly adjuvanted—were produced as a single recombinant fusion protein NS (LEISH-F3) for ease of antigen production and broad coverage of a heterogeneous major histocompatibility complex population. When formulated with glucopyranosyl lipid A-stable oil-in-water nanoemulsion (GLA-SE), a Toll-like receptor 4 T helper 1 (T helper 1) promoting nanoemulsion adjuvant, the LEISH-F3 polyprotein induced potent protection against both L. donovani and L. infantum in mice, measured as significant reductions in liver parasite burdens. A robust immune response to each component of the vaccine with polyfunctional CD4 T cell responses characterized by production of antigen-specific interferon-γ, tumor necrosis factor and interleukin-2 (IL-2), and low levels of IL-5 and IL-10 was induced in immunized mice. We also demonstrate that CD4 T cells, but not CD8 T cells, are sufficient for protection against L. donovani infection in immunized mice. Based on the sum of preclinical data, we prepared GMP materials and performed a phase 1 clinical study with LEISH-F3+GLA-SE in healthy, uninfected adults in the United States. The vaccine candidate was shown to be safe and induced a strong antigen-specific immune response, as evidenced by cytokine and immunoglobulin subclass data. These data provide a strong rationale for additional trials in Leishmania-endemic countries in populations vulnerable to VL.

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Leishmaniasis, a sandfly-transmitted series of poverty-related diseases caused by infection with Leishmania species parasites, threatens about 350 million people in 98 countries around the world. It has a wide range of clinical presentations that are dependent on the infecting strain and may cause cutaneous, mucosal or visceral leishmaniasis (CL, MCL or VL). VL, also known as kala-azar, is the most severe form of leishmaniasis. Two Leishmania proteins—nucleoside hydrolase and a sterol 24-c-methyltransferase, each of which are protective in animal models of VL when properly adjuvanted—were produced as a single recombinant fusion protein NS (LEISH-F3) for ease of antigen production and broad coverage of a heterogeneous major histocompatibility complex population. When formulated with glucopyranosyl lipid A-stable oil-in-water nanoemulsion (GLA-SE), a Toll-like receptor 4 T helper 1 (T helper 1) promoting nanoemulsion adjuvant, the LEISH-F3 polyprotein induced potent protection against both L. donovani and L. infantum in mice, measured as significant reductions in liver parasite burdens. A robust immune response to each component of the vaccine with polyfunctional CD4 T cell responses characterized by production of antigen-specific interferon-γ, tumor necrosis factor and interleukin-2 (IL-2), and low levels of IL-5 and IL-10 was induced in immunized mice. We also demonstrate that CD4 T cells, but not CD8 T cells, are sufficient for protection against L. donovani infection in immunized mice. Based on the sum of preclinical data, we prepared GMP materials and performed a phase 1 clinical study with LEISH-F3+GLA-SE in healthy, uninfected adults in the United States. The vaccine candidate was shown to be safe and induced a strong antigen-specific immune response, as evidenced by cytokine and immunoglobulin subclass data. These data provide a strong rationale for additional trials in Leishmania-endemic countries in populations vulnerable to VL.
The use of drugs plus whole parasites or crude antigens adsorbed with alum plus bacillus Calmette-Guérin or appropriate adjuvants has demonstrated that vaccination against leishmaniasis is possible.\(^2\) However, difficulty in standardizing formulation and dosing of live vaccines or crude preparations makes these approaches impractical and inconsistent. A selection of defined antigens including LEISH-111f developed by our group has been studied as vaccine antigen candidates for leishmaniasis in animal models and clinical studies. Using a combination of reverse vaccinology and bioinformatics approaches, we previously identified 43 Leishmania proteins recognized by antibodies in the serum of Sudanese VL patients as potential vaccine antigen candidates.\(^15\) Our selection criteria for vaccine development included sequence conservation across Leishmania species, but is absent in mammals.\(^19\) When appropriately formulated with a Toll-like receptor 4 (TLR-4) adjuvant, SMT provides protection against both L. infantum and L. major experimental infections.\(^15,19,20\)

In general, recombinant proteins are poorly immunogenic and require the addition of an adjuvant to elicit adaptive immune responses. To address this issue, we developed a synthetic TLR-4 ligand, glucopyranosyl lipid A (GLA).\(^21,22\) When formulated in a stable oil-in-water nanoemulsion (SE), the GLA-SE adjuvant system induces antigen-specific T\(_H1\) immune responses that are associated with efficacy in several animal models of infection, including tuberculosis, malaria and influenza.\(^23-35\) Completed and ongoing clinical trials have shown our GLA-based formulations to be both safe and immunogenic in vaccine strategies for infectious diseases, allergy and cancer.\(^36,37\)

Here we report recognition of NH and SMT in the sera of Leishmania-infected individuals from disease-endemic areas of Bangladesh and the development of a recombinant fusion antigen containing these two proteins, LEISH-F3. We characterized the immune responses induced by immunization of LEISH-F3 formulated with GLA-SE (LEISH-F3+GLA-SE), and demonstrate in mouse models of VL that prophylactic immunization confers protection against both causative strains of VL, L. donovani and L. infantum. We also report results for a first-in-human study that evaluated LEISH-F3+GLA-SE in healthy adults from a non-endemic region. As predicted by our preclinical studies, immunization with LEISH-F3+GLA-SE in a human clinical trial proved to be safe and highly immunogenic; in immunized subjects, we observe a substantial anti-LEISH-F3 serum antibody response and production of key protective cytokines by LEISH-F3-recalled peripheral blood mononuclear cells (PBMCs), which may be potent contributors to vaccine efficacy.

RESULTS

Immune recognition of NH and SMT by individuals from a VL-endemic area in Bangladesh

To validate the use of the NH and SMT proteins as potential vaccine candidates, we first evaluated whether they were recognized by antibodies in sera from Leishmania-exposed individuals from a disease-endemic area in Bangladesh (Figure 1). Endemic area residents...
were defined as either having active VL or as asymptomatics, and compared with healthy individuals from a non-endemic area (USA). To determine if sera from *Leishmania*-exposed individuals recognize NH or SMT, we performed enzyme-linked immunosorbent assays (ELISAs) for total immunoglobulin G (IgG) (Figure 1a) and compared recognition of our antigens with recognition of *L. donovani* soluble lysate antigen (SLA). Both VL and asymptomatic individuals had significantly higher anti-NH, anti-SMT and anti-*L. donovani* SLA serum IgG compared with non-endemic individuals (Figure 1a). Asymptomatic individuals had lower anti-*L. donovani* SLA compared those with VL, but had higher anti-NH and anti-SMT serum IgG compared with VL (Figure 1a).

The antigenicity of NH and SMT was then evaluated by ex vivo stimulation of PBMCs collected from asymptomatic individuals. Cells were used to characterize CD4+ T cells producing IFNγ, TNF and IL-2, following stimulation with *L. donovani* SLA, NH and SMT. PBMCs from non-endemic US subjects were used as negative controls. As expected, CD4+ T-cell IFNγ, TNF and IL-2 cytokine responses were significantly higher in the asymptomatic individuals from a VL hyperendemic region when recalled with the antigens as compared with the non-endemic healthy controls (Figure 1b).

**Generation and characterization of the LEISH-F3 fusion protein construct**

Having demonstrated IgG and cell-mediated responses against NH and SMT in *Leishmania*-infected individuals, as well as immunogenicity and protection against *L. donovani*, a straightforward approach was to link in tandem the open reading frames of both genes such that

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**Figure 2** LEISH-F3 construct and characterization. (a) Schematic of LEISH-F3 (NS) fusion protein. (b-e) SDS-PAGE and immunoblot of LEISH-F3 (NS). (b) LEISH-F3 was run in reducing and nonreducing conditions on a 4–20% Tris-glycine gel. (c) Immunoblot of LEISH-F3 with rabbit polyclonal antibody to LEISH-F3; 100 ng each of LEISH-F3, NH and SMT; 27 μg of *L. donovani* whole-cell lysate (WCL). (d) Immunoblot of 2 μg of LEISH-F3 and HMS174 E. coli lysate with antibody to HMS174 E. coli. (e, f) Immunoblot of LEISH-F3 and its component antigens, NH and SMT, with mouse monoclonal antibody (mAb) against (e) NH or (f) SMT; 100 ng each of LEISH-F3, NH and SMT were loaded.
the final construct would result in the generation of a single recombinant antigen, NS/LEISH-F3, comprising both antigens for evaluation as a VL vaccine candidate (Figure 2a). The L. donovani sequence for NH and the L. infantum sequence for SMT were used for the LEISH-F3 construct design. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis analysis of purified, scaled-up fermentation lots of LEISH-F3 revealed minor protein aggregation and a single major band at the expected molecular mass of 75 kDa. A similar band profile was observed for samples in reducing and nonreducing conditions, demonstrating the relative absence of protein aggregates (Figure 2b). The identity of the major band, relative absence of aggregates and large protein fragments were confirmed by immunoblotting with a mouse polyclonal serum raised against

Table 1 Homology of LEISH-F3 antigens, NH and SMT, among *Leishmania* species

| Leishmania species | NH | SMT |
|--------------------|----|-----|
| L. donovani        | 100| 99  |
| L. infantum        | 99 | 100 |
| L. major           | 96 | 97  |
| L. tropica         | 98 | NA  |
| L. mexicana        | 93 | 94  |
| L. braziliensis     | 84 | 86  |
| L. guyanensis      | 84 | 86  |
| L. amazonensis     | NA | 94  |
| L. aethiopica      | 30 | NA  |

Abbreviations: NA, not available; NH, nucleoside hydrolase; SMT, sterol 24-c-methyltransferase.

by immunoblotting with *Escherichia coli* LEISH-F3 was recognized by monoclonal sera raised against a single

to determine the immunogenic potential of NH, SMT and LEISH-F3 as vaccines, we formulated the single components and the fusion antigen with the synthetic nanoemulsion GLA-SE for immunization. Mice received three immunizations at 3-week intervals and antigen-specific antibody and T-cell responses were measured 4 weeks after the final immunization and compared with saline or LEISH-F3-alone controls (Supplementary Figure S1). Compared with control immunizations, LEISH-F3+GLA-SE generated higher anti-LEISH-F3 IgG1 and IgG2a serum antibodies; LEISH-F3+GLA-SE generated significantly higher titers of IgG2a than IgG1 (Figure 3a). A positive IgG2a: IgG1 ratio is indicative of an IFN-γ-dependent isotype switch in IgG subclasses typical of a Th1-biased immune response.

To demonstrate formally that antigen-specific Th1 responses had been induced in immunized mice, splenocytes were recalled with LEISH-F3 protein ex vivo. We observed a Th1-skewed memory response with IFN-γ, but not IL-5, producing cells readily detected in LEISH-F3+GLA-SE-immunized mice (Figure 3b and Supplementary Figure S1). The inverse was observed upon recall of cells from mice immunized with LEISH-F3 alone, indicating the importance of the GLA-SE adjuvant to subvert a Th1 response in favor of Th1 (Figure 3b). This was also reflected in the amount of total cytokines produced by LEISH-F3-recalled splenocytes isolated from LEISH-F3+GLA-SE-immunized mice, which had enhanced production of key Th1 cytokines IFN-γ, TNF-α and IL-2 and low levels of the Th12/Treg cytokines IL-5 and IL-10 (Figure 3c).

We further characterized the immune response generated following our immunization regimen with LEISH-F3+GLA-SE by analyzing the LEISH-F3-specific T-cell memory response using multiparameter flow cytometry and intracellular cytokine staining. Although LEISH-F3-specific CD8 T cells responses were not observed, LEISH-F3+GLA-SE immunization generated LEISH-F3-specific CD4 T cells as indicated by the higher percentage of these cells expressing CD154 (a ligand expressed by recently activated CD4 T cells) following ex vivo recall with LEISH-F3 (Figure 3d). LEISH-F3-specific CD4 T cells from LEISH-F3+GLA-SE mice proved to be Th1-skewed and polyfunctional (Figures 3e and f). Mice that received the LEISH-F3+GLA-SE vaccine showed a significant increase in the proportion of LEISH-F3-specific CD4 T cells producing all three of the key Th1 cytokines (IFN-γ, IL-2 and TNF; 0.23% of CD4 T cells) or some combination of two (0.40% of CD4 T cells) compared with the untreated control; a total of 0.63% of CD4 T cells from LEISH-F3+GLA-SE mice were polyfunctional in contrast to 0.11% of the LEISH-F3-alone immunization. Thus, immunization of mice with LEISH-F3+GLA-SE induces polyfunctional antigen-specific Th1 responses, which are classically associated with protection against experimental *Leishmania* infection.

We also conducted CD4 T-cell epitope mapping of LEISH-F3- in the LEISH-F3+GLA-SE-immunized mice. A panel of 164 peptides, each 15-amino-acid long and overlapping by 11 residues, were organized into a two-dimensional matrix of 26 peptides pools, so that individual peptides were present in two independent pools. When a positive response was observed from a peptide in both peptide pools, it was further evaluated in follow-up splenocyte recalls to determine the actual individual peptide that induced positive responses. We found that BALB/c mice had several CD4 epitopes for LEISH-F3 in both the NH and SMT portions of the fusion antigen (Figure 3g).

**Immunization with LEISH-F3+GLA-SE induces protection against experimental VL infection with both *L. donovani* and *L. infantum***

To evaluate the protective potential of immunization with NH, SMT or LEISH-F3, each formulated with GLA-SE, BALB/c and C57BL/6 mice were immunized and experimentally infected with *L. donovani* or *L. infantum* 4 weeks after the final immunization. Control mice received saline injections, as our prior experience demonstrated that injection with GLA-SE alone in the same regimen does not change infection relative to treatment with saline (data not shown). Parasite burdens in livers were quantified 4 weeks after infection. A significant reduction of *L. donovani* parasites was observed in both BALB/c and C57BL/6 mice immunized with NH, SMT or LEISH-F3 formulated with GLA-SE compared with mice given the control preparations.
Owing to the high cross-Leishmania species homology of NH and SMT (Table 1), we evaluated if immunization with LEISH-F3+GLA-SE could also reduce parasite burdens in mice infected with the other causative strain of VL, *L. infantum*. We found that the LEISH-F3+GLA-SE-immunized mice had significantly reduced *L. infantum* parasite burdens (Figure 4c).

The importance of memory CD4 T cells generated by LEISH-F3+GLA-SE immunization in protection against *L. donovani* infection was examined by depleting either CD4 or CD8 T cells in mice subjected to the standard immunization regimen before infection with *L. donovani*. In LEISH-F3+GLA-SE-immunized mice, depletion of CD4 T cells, but not CD8 T cells, resulted in the loss of protection and parasite liver burdens comparable to the unprotected saline-immunized mice (Figure 4d).

Having demonstrated protection against the causative strains of VL, *L. donovani* and *L. infantum*, with prophylactic immunization in preclinical mouse models, the LEISH-F3+GLA-SE vaccine was advanced to clinical trials to evaluate the safety, tolerability and immunogenicity of the vaccine. Thirty-six volunteers were randomized, enrolled and received at least one study injection; 12 subjects received the vaccine 20μg LEISH-F3+2μg GLA-SE, 12 subjects received the vaccine 20μg LEISH-F3+5μg GLA-SE and 12 received 20μg LEISH-F3 protein alone (see Supplementary Table 1). Each volunteer received three intramuscular injections, administered at 28-day intervals. Eight of 12 subjects in the 20μg LEISH-F3+2μg GLA-SE group, 10 of 12 in the 20μg LEISH-F3+5μg GLA-SE group and 9 of...
12 subjects in the 20 μg LEISH-F3-alone group completed the final day 421 visit. The remainders withdrew consent or were lost to follow-up before completing the study (Figure 5).

Safety evaluation. The vaccine was safe and well tolerated in healthy adults, with no serious adverse events (AEs), no AE of special interest and no AE of grade 3 or grade 4 toxicity observed in any of the study groups. The most frequent local reactogenicity AE was tenderness/pain, which occurred in 95.8% of vaccine recipients and 41.7% of antigen-only recipients (Table 2a). The most frequent systemic reactogenicity AE was fatigue, which occurred in 50.0% of vaccine recipients and 33.3% of antigen-only recipients (Table 2b). The most common AE were injection site tenderness/pain, fatigue and decreased hemoglobin (Table 2c). Injection site tenderness/pain was the only AE found to have a statistically significant difference in frequency between the groups that received the vaccine and antigen alone (P<0.001). There were no clinically significant changes in clinical chemistry or hematology values. These data indicate that the vaccine was safe and well tolerated at either dose level.

Vaccine-induced immune responses. The antigen-specific immune responses of volunteers who received all three study injections were evaluated. In terms of both the humoral and cellular responses to LEISH-F3 (Figure 6 and Supplementary Figure S2), the vaccine was immunogenic at both 2 and 5 μg of GLA-SE doses. Although subjects vaccinated with the LEISH-F3 protein alone did not develop antigen-specific antibody responses, subjects vaccinated with LEISH-F3+GLA-SE had significant levels of antigen-specific IgG antibodies in their serum. These elevations were evident at day 35 of observation and were maintained up to the last study time point of day 168 (Figures 6a and b). An assessment of IgG subclasses revealed a preferential increase in IgG1 and IgG3 subclasses in the GLA-SE-containing groups (Figure 6a). Antigen-specific IgE was not detected at any time nor in any group (Figure 6a). As the induction of IgG1 is IFN-γ-dependent and IgG4 is dependent on IL-4, a preferential induction of isotypes supported by TH1-like cytokines, is indicative of a T H1 response driven by the addition of GLA-SE.41,42

Whole blood assay (WBA) of subjects injected with protein alone did not demonstrate any antigen-specific cytokine responses at any time point. However, at day 35, the secretion of IFN-γ, TNF and IL-2 in response to antigen was observed by WBA of subjects injected with LEISH-F3+GLA-SE. Secretion of these cytokines was further elevated at days 63 and 84, indicating that additional immunizations boosted the cellular response (Figure 6c). Responses were still detected at day 168, although they were lower than those measured at day 84 for IFN-γ and TNF, indicating a slight waning of the response over time. IL-2 cytokine responses alternatively remained consistently high through the last time point tested at day 168. Unlike observations in immunized mice, secretion of IL-5 and IL-10 was observed by WBA of subjects injected with LEISH-F3+GLA-SE (Figure 6c).
However, IL-10 responses showed a marked decrease after the peak response following day 63. Additionally, in the GLA-SE-containing groups, a significant difference was observed in the group that received 5 μg GLA-SE in terms of increased TNF production at days 63 and 84 and day 168 for IL-2 when compared with the 2 μg GLA-SE group. Furthermore, we evaluated the cytokine responses of PBMC from LEISH-F3-vaccinated subjects when recalled with the LEISH-F3 components (Supplementary Figure S2). As expected, subjects immunized with LEISH-F3+GLA-SE had significant levels of antigen-specific IgG antibodies to both the NH and SMT components compared with the antigen-alone group. No differences were observed in the antibody responses to the components between the 2 and 5 μg adjuvant dose groups of GLA-SE. The cytokine responses to the components reflected the pattern seen with the fusion protein LEISH-F3. An increased TH1 and TH2 cytokine response profile to the LEISH-F3 components were seen for the adjuvanted groups compared with antigen alone at day 63, and this was further characterized at day 168 by a decrease in IL-10 and IL-5 and a persistence of IL-2 (against NH and SMT) and TNF (against NH) responses (Supplementary Figure S2). Taken together, these data indicate that LEISH-F3+GLA-SE vaccination is safe in humans and induces antigen-specific cellular responses that could protect against Leishmania infection.

**DISCUSSION**

Strategic framework to eliminate VL in South-East Asia by 2015 has reported a decline in incidence of ~ 35% owing to implementation of single dose AmBisome and indoor residual spraying of insecticides. Coordinated efforts to improve active case searches as well as early detection and diagnostics services have also reported declining incidences in East Africa (http://www.unitingtocombatNTDs.org). Nevertheless, resistance to affordable first-line treatment options such as pentavalent antimonials has developed, and vector control efforts on disease incidence and transmission have yielded varied results. Control of leishmaniasis could be achieved by developing new vaccines or less toxic drugs. Regardless of the strategies adopted for elimination, we believe that, based on the intransigent nature of VL, it is important to develop tools to ensure that elimination is complete. To date, no disease has been eliminated without an effective vaccine.

We designed the subunit vaccine candidate, LEISH-F3+GLA-SE, to be a cost-effective measure against VL, and performed preclinical analyses of its immunogenicity, safety, toxicity and efficacy against challenge with *L. donovani* and *L. infantum*. LEISH-F3 consists of the fusion of two antigens each of which were recognized by humans and were previously shown to confer partial protection in mouse models. LEISH-F3+GLA-SE immunization induced antigen-specific antibody and T-cell immune responses in mice and humans. Induction of predominantly TH1-type CD4 T-cell responses was associated with reduced parasite burdens in the livers of vaccinated mice. Furthermore, we demonstrated that immunization with LEISH-F3+GLA-SE in mouse models was protective against both of the causative species of VL, *L. donovani* and *L. infantum*.

A considerable number of Leishmania antigens including TSA, LmST11, A2, LelF, Leish-111f, KMP-11, KSAc, CPB, HbR, H2A/H2B, p36/LACK and ORFF have been tested as subunit protein or nucleic acid vaccine candidates against VL and have had variable or unreproducible success in inducing protection. As demonstrated for both leishmaniasis and tuberculosis, combining multiple antigens in recombinant fusion proteins such as LEISH-F2/LEISH-111f, Mtb72f, ID93, H56, Ag85B-ESAT6, ESAT6-TB10.4, CSU-F36 and Ag85B-TB10 leads to increased vaccine efficacy. Of these, the multicomponent vaccine LEISH-F2/LEISH-111f+MPL-SE was the first defined vaccine candidate to progress to human clinical trials in healthy volunteers in CL and ML patients in Brazil and Peru and healthy subjects in India. Because of their low intrinsic immunogenicity, protein-based vaccines need suitable adjuvant

![Clinical trial CONSORT diagram](image-url)
systems for the induction of strong in vivo immune responses. Several of the most advanced subunit vaccine candidates against non-viral disease targets currently in human clinical trials, RTS, S (malaria vaccine)\(^\text{72}\), ID93, Mb72F, H56 (TB vaccine\(^\text{16,57}\)), are based on liposomal (AS01, CAF01) or oil-in-water (GLA-SE, MPL-SE) formulations, some of which contain the TLR-4 agonist GLA or MPL. Our selection of GLA-SE was based on its ability to activate dendritic cells in a TLR-4-dependent manner and potentiate TH1-type CD4

T-cell responses,\(^\text{22,26,61}\) which are critical for controlling intracellular infections.

Three doses of the LEISH-F3+GLA-SE vaccine induced antigen-specific TH1 responses in mice and humans. Increased production of IFN\(\gamma\)-IL-5 ratios were observed in BALB/c mice. IFN\(\gamma\), TNF and IL-2 are involved in protection against VL\(^\text{39,74,76}\) and TNF has been shown to synergize with IFN\(\gamma\) to kill Leishmania parasites.\(^\text{77}\) Therefore, induction of LEISH-F3-specific T cells capable of producing multiple cytokines upon antigen recall might be beneficial for control of Leishmania infection, and such induction may be a good indicator of whether a vaccine will be protective against leishmaniasis. In mice, the ability of LEISH-F3+GLA-SE to induce polyfunctional T-cell responses was associated with a statistically significant reduction in liver parasite burden after infection with L. donovani, in both BALB/c and C57BL/6 mouse models, or L. infantum in BALB/c mice. Given that the two antigens comprising LEISH-F3 each provide robust protection against challenge with multiple Leishmania species,\(^\text{15-20}\) lack homology with mammalian proteins and are conserved among multiple Old and New World species, we suggest that LEISH-F3 may prove to be a superior vaccine candidate for the spectrum of leishmaniasis diseases. The LEISH-F3-specific cytokine response in vaccinated clinical trial subjects was a complex mixture of TH1- and TH2-type responses, a phenotype consistent with observed T-cell memory responses in successful vaccine regimens, such as for yellow fever.\(^\text{78}\) Without a definitive correlate of protective immunity against leishmaniasis, vaccine development has had to rely on screening a range of cytokines to estimate the balance between TH1 and TH2 responses and this balance has been shown to be involved in the outcome of human leishmaniasis.\(^\text{79}\) We showed that, in addition to its TH1 response-inducing activity, LEISH-F3 induces IL-10 in vaccinated individuals. Although a balanced response that includes IL-10 may prevent the development of pathology associated with an excessively robust proinflammatory immune response, the induction of IL-10 is a potential concern that will be monitored in future leishmaniasis clinical trials.\(^\text{79}\) In conclusion, these results support the advancement of the LEISH-F3+GLA-SE leishmaniasis vaccine and plans for additional phase 1 clinical trials in endemic countries are underway.

**METHODS**

**Recombinant proteins**

Recombinant proteins were cloned and expressed in E. coli as described previously.\(^\text{58,80}\) The LEISH-F3 fusion protein was constructed by aligning the individual gene sequences for NH and SMT as a single product. Proteins were quantified using the BCA protein assay (Pierce, Rockford, IL, USA) and were all <100 endotoxin units per mg protein as measured by Limulus Amebocyte Lysate QCL-1000 assay (Lonza Inc., Basel, Switzerland). LEISH-F3 (2 \(\mu\)g per lane) was analyzed by reducing and nonreducing sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). LEISH-F3, NH, SMT (100 ng per lane) was analyzed by reducing and nonreducing sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). LEISH-F3, NH, SMT (100 ng per lane) or L. donovani whole-cell lysate (27 \(\mu\)g per lane) were immunoblotted with rabbit polyclonal antibody to LEISH-F3 and with mouse monoclonal antibody against NH or SMT. To demonstrate the absence of host cell proteins, LEISH-F3 (2 \(\mu\)g per lane) was immunoblotted with rabbit polyclonal antibody to HMS174 E. coli, with HMS174 E. coli lysate as a positive control.

**Recognition of NH and SMT in samples from individuals from a Leishmania-endemic area**

Sera were obtained from residents of a Leishmania-endemic area of Bangladesh (VL patients and asymptomatics) or the United States (non-endemic normal). Written informed consent for study participation was obtained from each participant before screening. Subjects were defined according to parameters set by the Kala-Azar Elimination Program as follows: VL, a person with clinical symptoms of VL (fever for more than 2 weeks duration and splenomegaly) and

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**Table 2 Safety evaluations for the phase 1 clinical trial**

| AE                                      | LEISH-F3 |
|-----------------------------------------|----------|
|                                         | 20 \(\mu\)g | 20 \(\mu\)g | 20 \(\mu\)g |
| GLA-SE                                  | 5 \(\mu\)g | -           |             |
| (n = 12)                                | (n = 12)  | (n = 12)    |             |
| (A) Any local reaction                  |          |            |             |
| Postinjection period 1                  | 11 (91.7%) | 12 (100%)  | 6 (50.0%)   |
| Postinjection period 2                  | 10 (83.3%) | 12 (100%)  | 5 (41.7%)   |
| Postinjection period 3                  | 7 (58.3%) | 9 (75.0%)  | 3 (25.0%)   |
| Injection site tenderness/pain          | 4 (33.3%) | 7 (58.3%)  | 1 (8.3%)    |
| Injection site induration/swelling      | 11 (91.7%) | 12 (100%)  | 5 (41.7%)   |
| Injection site erythema/redness         | 3 (25.0%) | 1 (8.3%)   | 1 (8.3%)    |
| Injection site ecchymosis               | 1 (8.3%) | 0 (0%)     | 0 (0%)      |
| (B) Any systemic reaction               |          |            |             |
| Postinjection period 1                  | 5 (41.7%) | 7 (58.3%)  | 5 (41.7%)   |
| Postinjection period 2                  | 4 (33.3%) | 4 (33.3%)  | 4 (33.3%)   |
| Postinjection period 3                  | 1 (8.3%) | 5 (41.7%)  | 1 (8.3%)    |
| Anorexia                                | 1 (8.3%) | 1 (8.3%)   | 0 (0%)      |
| Chills                                  | 0 (0%)   | 1 (8.3%)   | 0 (0%)      |
| Fatigue                                 | 5 (41.7%) | 7 (58.3%)  | 4 (33.3%)   |
| Headache                                | 1 (8.3%) | 1 (8.3%)   | 0 (0%)      |
| Myalgia                                 | 1 (8.3%) | 2 (16.7%)  | 0 (0%)      |
| (C) Injection-related reactions         |          |            |             |
| Anorexia                                | 3 (25.0%) | 2 (16.7%)  | 1 (8.3%)    |
| Fatigue                                 | 5 (41.7%) | 7 (58.3%)  | 4 (33.3%)   |
| Injection site induration/swelling      | 3 (25.0%) | 1 (8.3%)   | 1 (8.3%)    |
| Injection site tenderness/pain          | 11 (91.7%) | 12 (100%)  | 5 (41.7%)   |
| Laboratory Investigations              |          |            |             |
| Hemoglobin decreased                    | 2 (16.7%) | 4 (33.3%)  | 5 (41.7%)   |
| White blood cell count decreased        | 2 (16.7%) | 1 (8.3%)   | 3 (25.0%)   |
| General AEs                             |          |            |             |

Abbreviations: AE, adverse event; GLA-SE, glucopyranosyl lipid A-stable oil-in-water nanoemulsion.
Figure 6 LEISH-F3 formulated with GLA-SE is immunogenic in humans. Thirty-six healthy adult subjects were immunized on days 0, 28 and 56 with 20μg LEISH-F3+2μg GLA-SE (n=12), 20μg LEISH-F3+5μg GLA-SE (n=12) and 20μg LEISH-F3 protein alone (n=12). The immunogenicity of the vaccine was evaluated by assessing antibody and T-cell responses at days 0, 35, 63, 84 and 168 and days 0, 7, 35, 63, 84 and 168, respectively. (a and b) ELISAs for titers of LEISH-F3-specific antibodies (total and IgG subclasses and total IgE) in volunteer serum were conducted for the indicated time points. (c) Quantitative T-cell responses to LEISH-F3 was measured by IL-2, IL-5, IL-10, IFN-γ and TNF cytokine production in whole blood Luminex assay (WBA). Nine subjects were excluded from the per-protocol population immunology summaries resulting in evaluable groups such as: 2μg GLA-SE (n=8), 5μg GLA-SE (n=10) and LEISH-F3 alone (n=9). P-value for comparison was performed for various treatment groups: between 2 and 5μg GLA-SE vaccine groups and between vaccine (2 and 5μg GLA-SE vaccine groups combined) and 20μg LEISH-F3 alone. P-values were considered significant at the 0.05 significance level. *P-values significant at 0.05 significance level when compared between vaccine (2 and 5μg GLA-SE vaccine groups combined) and 20μg LEISH-F3 alone. **P-values significant at 0.05 significance level when compared between vaccine (2 and 5μg GLA-SE vaccine groups separately).
a positive rK39 RDT result; asymptomatic individual, a person from an endemic area with no present or past clinical symptoms of VL but with a DAT titer >600; non-endemic, healthy individuals living out with endemic regions reporting no history of VL. ELISA was conducted as previously described with the sera at a 1:400 dilution.82

Intracellular cytokine staining: PBMCs were isolated from peripheral blood of the subjects following the standard Ficoll procedure. PBMCs were stimulated with NH and SMT, phosphate-buffered saline, with costimulatory antibodies CD28 and CD49d (BD Biosciences, San Jose, CA, USA), in complete RPMI at 37 °C. Cells were then stained for membrane expression of CD35 (clone UCHT1; Beckman Coulter, Pasadena, CA, USA), CD4 (OKT4) and CD8 (HI108a) (BD Bioscience). Cells were fixed and permeabilized in Cytofix/Cytoperm (BD Biosciences) according to the manufacturer’s instructions and stained for intracellular expression of IFN-γ (4S.B3), TNF (MAB11) and IL-2 (MQ1-17H12) (BD Bioscience). Events were acquired using a BD LSRFortessa (BD Biosciences) and the data were analyzed with FlowJo (Tree Star Inc., Ashland, OR, USA). For analyses, the cells were first gated at CD3 to identify CD4 and CD8 T cells, and then gated for IFNγ, TNF and IL-2. Responses are shown as background (phosphate-buffered saline)-subtracted percent antigen-specific T cells producing the indicated cytokines.

Mouse treatments
All animal procedures were approved by the IDRI institutional animal care and use committee. Female C57BL/6 and BALB/c mice were purchased from Charles River Laboratories (Wilmington, MA, USA). Mice were maintained in specific pathogen-free conditions in the animal facilities of the IDRI and entered experiments at 6–8 weeks of age. Mice were immunized three times at 3-week intervals by injection at the base of the tail. Immunizations were made in a volume of 100 μl per dose as follows: 5 μg per dose of LEISH-F3; 5 μg of GLA in stable nanoemulsion (SE) of 2% oil (GLA-SE). CD4 or CD8 T cells were depleted by intraperitoneal injection of 0.5 mg of either CD4 (clone GK1.5), CD8 (clone 53–6.72) or a rat IgG2a isotype control (Bio X Cell, West Lebanon, NH, USA) on 3 consecutive days. Depletion was confirmed by flow cytometry analysis of blood. Three weeks after depletion, mice were infected with L. donovani.

Antibody analyses
Antigen-specific serum antibodies were measured by ELISA. End-point titers were calculated using the optical density >0.1 as a cutoff value with GraphPad Prism 6 (GraphPad Software Inc., La Jolla, CA, USA). End-point titers of samples were recorded as <100 if optical density values of the samples were lower than the cutoff value at 1:100 dilution.

Cell preparations and antigen stimulation assays
Four weeks after the final immunization, spleens were removed and single-cell suspensions were prepared. Splenocytes were enumerated using a ViaCount assay with a PCA system (Guava Technologies, Darmstadt, Germany), and then incubated with media only or 10 μg ml⁻¹ antigen.

ELISPOT assays: IFNγ and IL-5 ELISPOTs (R&D Systems, Minneapolis, MN, USA) were conducted according to the manufacturer’s instructions. Spot images were collected using ImmunoCapture 6.4 and analyzed with ImmunoSpot 5.0 on an automated ELISPOT plate reader (C.T.L. Seri3A Analyzer; Cellular Technology, Shaker Heights, OH, USA).

Secreted cytokines detection. Culture supernatants were collected after 72 h, and IFNγ, TNF, IL-2, IL-5 and IL-10 were determined using bead-based multiplex assay with BD Biosciences’ cytometric bead array (CBA) flex set according to the manufacturer’s instructions (BD Biosciences).

Intracellular cytokine staining and flow cytometry. Splenocytes were cultured with antigen plus 1 μg ml⁻¹ each of the costimulatory antibodies anti-mouse CD28 and CD49d (BD Biosciences), and then stained for membrane expression of CD90.2 (30-H12), CD4 (GK1.5) and CD8 (53–6.7) (BD Biosciences). Cells were then fixed and permeabilized in Cytofix/Cytoperm (BD Biosciences) according to the manufacturer’s instructions and stained for intracellular expression of CD154 (MR1; eBioscience), IFNγ (XMG1.2), TNF (MP6-XT22) and IL-2 (JES6-5H4) (BD Bioscience). Events were acquired using a BD LSRFortessa (BD Biosciences) and the data were analyzed with FlowJo (Tree Star Inc., Piscataway, NJ, USA). Cells were gated through CD90.2⁺ T cells to analyze CD4 and CD8 T cells. Intracellular molecules were acquired after gating through CD4⁺CD8⁻ T cells. For CD4 epitope mapping analysis, cells were gated and analyzed for CD154 expression.

Infection and parasite quantification. L. donovani (MHOM/SD/00/1S-2D) and L. infantum (MHOM/BR/82/BA-2) were routinely passed through Syrian golden hamsters to generate virulent amastigotes and promastigote stocks in the M199 medium. Mice were challenged by retro-orbital intravenous injection with either 1 × 10⁶ L. donovani or 5 × 10⁵ L. infantum parasites. Three or four weeks after infection, mice were euthanized, DNA was extracted from homogenate using QiAmp DNA Mini Kits (Qiagen, Hilden, Germany). DNA was quantified using Nanodrop UV-Vis spectrophotometer (ND-1000). L. donovani DNA was detected using primers for L42486 (forward, 5'-GC GACGTCCTGGAAAGAAA-3'; and reverse, 5'-GGCGGTACACATTAGCAG AA-3') with a FAM reporter sequence (5'-CAACCGGATCCTCC-3'), which detects a 203-bp genomic repeat region specific to Leishmania species (NCBI Blastn). L. infantum primers were used previously,83 Mouse Gapdh FAM (Life Technologies, Carlsbad, CA, USA) was used as an internal reference control. Cp's of samples were fitted to a standard curve to determine number of parasite per μl of DNA. Final parasite burdens are expressed in the number of Leishmania parasites per organ.

Statistical methods
Data for human ELISA were compared by analyses of non-endemic normals vs VL patients or asymptomatics. Data for intracellular cytokine staining were compared by analyses of non-endemic vs asymptomatics. Statistics were calculated by a Kolmogorov-Smirnov test using GraphPad Prism Version 6.01 (GraphPad Software, San Diego, CA, USA). For data generated using mouse samples, statistics was determined by one- or two-way ANOVA (GraphPad Prism). Significance was considered when the P-values were <0.05.

Clinical methods
Trial design: The safety, tolerability and immunogenicity of LEISH-F3+GLA-SE was evaluated in a first-in-man phase 1, randomized, dose-escalation study. A total of 36 healthy adults (male and female adults with no history of travel to Leishmania-endemic areas) were recruited after signing the informed consent form, and then randomly assigned to receive injections with 20 μg LEISH-F3 in combination with either nothing, 2 μg GLA-SE or 5 μg GLA-SE (12 per group) at 28-day intervals (days 0, 28 and 56). The treatment randomization list was generated using SAS software (SAS Institute Inc., Cary, NC, USA). The study biostatistician filled a set of individual, opaque, sealed envelopes, each labeled with a unique subject/participant identification number and containing the corresponding treatment assignment. Only the designated pharmacist(s) responsible for study injection preparation had access to the randomization assignment envelopes. All study injections were 0.5 ml in volume and were prepared no more than 2 h before administration. The investigators who evaluated study subjects for reactogenicity and AEs were blinded to study treatment assignments (only the study pharmacists were unblinded).

Twenty-four subjects were granted Medical Monitor approval for laboratory eligibility criteria deemed not clinically significant. In addition, the day 421 follow-up telephone visit for subjects was performed – 4 months early due to closure of the study site.

Out of 36 subjects who were randomized to injection, seven subjects withdrew consent or were lost to follow-up before receiving all three injections (three in the 2 μg GLA-SE group, two in the 5 μg GLA-SE group and two in the LEISH-F3-alone group), one subject (in the LEISH-F3-alone group) missed the day 56 evaluation and did not receive a third injection, and one subject (in the 2 μg GLA-SE group) was lost to follow-up before day 84. Thus, nine subjects were excluded from the per-protocol population immunology summaries resulting in evaluable groups for the ELISA (against LEISH-F3, NH and
SMT) and WBA (against LEISH-F3) as follows: 2 μg GLA-SE (n = 8), 5 μg GLA-SE (n = 10) and LEISH-F3 alone (n = 9). Final resulting evaluable groups for PBMC luminex (against NH and SMT) were: 2 μg GLA-SE (n = 6), 5 μg GLA-SE (n = 9) and LEISH-F3 alone (n = 7).

Safety evaluations. The primary safety end points were the proportion of subjects with AEs, serious AEs, AEs of special interest, local injection site reactions and specific systemic reactions to the study injections from days 0 to 84. The occurrence of AEs, SAEs and AEs of special interest was monitored from the time of the first study injection at day 0 through day 421, and the investigator assessed their relation to study injections and severity based on the Food and Drug Administration Guidance for Industry Toxicity Grading Scale for Healthy Adults Enrolled in Preventive Vaccine Clinical Trials, 2007. AE grades for laboratory parameters were set based on normal laboratory values. For non-laboratory conditions, AEs that were mild and did not limit activity were considered grade 1; those that were moderate and interfered with function, but not with activities of daily living, grade 2; those that were severe and interfered with activities of daily living, grade 3; and those that were life-threatening or disabling, grade 4. Serious AEs were those that resulted in death, were considered life-threatening, required hospitalization or prolongation of hospitalization, or resulted in persistent or significant disability, or resulted in a congenital anomaly or birth defect in the offspring of a study patient. AEs of special interest were those that met the Food and Drug Administration list for specific AEs. The occurrence of local injection site reactions was assessed at 30 min, 2 days and 7 days following each study injection. The occurrence of the specific systemic reactions headache, anorexia, malaise/fatigue, muscle pain, fever, hives, rash and chills was evaluated in the 7 days following each study injection. Blood samples were collected for measurement of hematology and serum chemistry parameters at screening and on days 7, 35 and 63. Vital signs were collected at every study visit and before and 30 min after study injection on days 0, 28, and 56.

Total IgG antibody assay. High-binding 384-well ELISA plates (Corning, Tewksbury, MA, USA) were coated with recombinant protein, and then serial dilutions of the plasma samples and positive control and a single dilution of negative control were prepared in serum diluent. After incubation, peroxidase-labeled anti-human IgG, IgG1, IgG2, IgG3, IgG4 and IgE (Life Technologies, Carlsbad, CA, USA) in serum diluent was added. End-point titer was determined by Graph Prism Nonlinear regression (curve-fitting) software (MiraiBio, South San Francisco, CA, USA).

Conflict of interest
SR is a coinventor on a patent for leishmaniasis vaccine development. All the other authors declare no conflict of interest.

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1 Kedzierski L, Sakthianandeswaran A, Curtis JM, Andrews PC, Junk PC, Kedzierska K. Leishmaniasis: current treatment and prospects for new drugs and vaccines. Curr Med Chem 2009; 16: 599–614.

2 World Health Organization. Control of the leishmaniases. World Health Organ Tech Rep Ser 2010; 793: 1–158.

3 Alvar J, Velez ID, Bern C, Herrero M, Desjeux P, Cano J et al. Leishmaniasis worldwide and global estimates of its incidence. PLoS ONE 2012; 7: e36571.

4 Joshi AB, Banjara MR, Pikhrel S, Jimba M, Singhhasivanon P, Ashford RW. Elimination of visceral leishmaniasis in Nepal: pipe-dreams and possibilities. Kalmimandu Univ Med J (KUMJ) 2006; 4: 488–496.

5 Kumar V, Kesari S, Dinesh DS, Tiwari AK, Kumar AJ, Kumar R et al. A report on the indoor residual spraying (IRS) in the control of Phlebotomus argentipes, the vector of visceral leishmaniasis in Bihar (India): an initiative towards total elimination targeting 2015 (Series-1). J Vector Borne Dis 2009; 46: 225–229.

6 Bhattacharya SK, Sur D, Sinha PK, Karbawang J. Elimination of leishmaniasis (kala-azar) from the Indian subcontinent is technically feasible & operationally achievable. Indian J Med Res 2006; 123: 195–196.

7 Das VN, Dinesh DS, Verma N, Kar SK. A case report on self-cure of visceral leishmaniasis. J Commun Dis 2002; 34: 302–303.

8 Lee BY, Bacon KM, Shah M, Kitchen SB, Connor DL, Slayton RB. The economic value of a visceral leishmaniasis vaccine in the Indian state. Am J Trop Med Hyg 2012; 86: 417–425.

9 Carvalho EM, Badaro R, Reed SG, Jones TC, Johnson WD Jr. Absence of gamma interferon and interleukin 2 production during active visceral leishmaniasis. J Clin Invest 1985; 76: 2066–2069.

10 Carvalho EM, Johnson WD, Barreto E, Marsden PD, Costa JL, Reed S et al. Cell-mediated immunity in American cutaneous and mucosal leishmaniasis. J Immunol 1985; 135: 4144–4148.

11 Kaye PM, Aeberscher T. Visceral leishmaniasis: immunology and prospects for a vaccine. Clin Microbial Infect 2011; 17: 1462–1470.

12 Kumar R, Nylen S. Immunobiology of visceral leishmaniasis. Front Immunol 2012; 3: 1–10.

13 Nylen S, Mauery R, Eidurmo L, Manandhar KD, Sundar S, Sacks D. Splenic accumulation of IL-10 mRNA in T cells distinct from CD4+CD25+ (Foxp3) regulatory T cells in human visceral leishmaniasis. J Exp Med 2007; 204: 805–817.

14 Goto Y, Coyer RN, Guderian J, Mohamath R, Reed SG. Cloning, characterization, and serodiagnostic evaluation of Leishmania infantum tandem repeat proteins. Infect Immun 2006; 74: 3939–3945.

15 Paraguai de Souza E, Bernardo RR, Palatinik M, Palatinik de Sousa CB. Vaccination of Balb/c mice against experimental visceral leishmaniasis with the GP36 glycoprotein antigen of Leishmania donovani. Vaccine 2001; 19: 3104–3115.

16 Aguilar-Be I, da Silva Zardo R, Paraguai de Souza E, Borja-Cabrera GP, Rosado-Vallado M, Mut-Martín M et al. Cross-protective efficacy of a prophylactic Leishmania donovani DNA vaccine against visceral and cutaneous murine leishmaniasis. Infect Immun 2005; 73: 812–819.

17 Borja-Cabrera GP, Correa Pontes NN, da Silva VC, Paraguai de Souza E, Santos WR, Gomes EM et al. Long lasting protection against canine kala-azar using the FML-QuilA saponin vaccine in an endemic area of Brazil (Salao Goncalo do Amarante, RN). Vaccine 2002; 20: 3277–3284.

18 Goto Y, Bogatzki LY, Bertholet S, Coyer RN, Reed SG. Protective immunization against visceral leishmaniasis using Leishmania sterol 24-c-methyltransferase formulated with MPL-SE induces cross-protection against L. major infection. Vaccine 2009; 27: 2884–2890.

19 Golder RN, Bertholet S, Mouttaifi M, Guderian JA, Windish HP, Baldwin SJ et al. Development and characterization of synthetic glucopyranosyl lipid adjuvant system as a vaccine adjuvant. PLoS ONE 2011; e61333.

20 Anderson RC, Fox CB, Dutilh TS, Shaverdin N, Evers TL, Poshusta GR et al. Physicochemical characterization and biological activity of synthetic TLR4 agonist formulations. Colloids Surf B 2010; 75: 123–132.

21 Bertholet S, Ireton GC, Ordway DJ, Windish HP, Pine SO, Kahn M et al. Cross-protective efficacy of a prophylactic Leishmania donovani DNA vaccine against visceral and cutaneous murine leishmaniasis. Infect Immun 2005; 73: 812–819.

22 Bertholet S, Ireton GC, Ordway DJ, Windish HP, Pine SO, Kahn M et al. Cross-protective efficacy of a prophylactic Leishmania donovani DNA vaccine against visceral and cutaneous murine leishmaniasis. Infect Immun 2005; 73: 812–819.

23 Wiley SR, Raman VS, Desbiens A, Bailor HR, Bhardwaj R, Shaki AR et al. Targeting TLRs expands the antibody repertoire in response to a malaria vaccine. Sci Transl Med 2011; 3: 93ra69.
24 Baldwin SL, Fox CB, Pallansch MA, Coler RN, Reed SG, Friede M. Increased potency of an inactivated trivalent polo virus with oil-in-water emulsions. Vaccine 2011; 29: 644–649.

25 Berthelet S, Goto Y, Carter L, Bhafia A, Howard RF, Carter D et al. Optimized subunit vaccine protects against experimental visceral leishmaniasis. Vaccine 2009; 27: 7036–7045.

26 Fox CB, Anderson RC, Duttill TS, Goto Y, Reed SG, Vedvick TS. Monitoring the effects of component structure and source on formulation stability and adjuvant activity of oil-in-water emulsions. Vaccine 2009; 27: 96–99.

27 Fox CB, Baldwin SL, Duthie MS, Reed SG, Vedvick TS. Immunomodulatory and physical effects of oil composition in vaccine adjuvant emulsions. Vaccine 2011; 29: 9563–9572.

28 Fox CB, Friede M, Reed SG, Ierton GC. Synthetic and natural TLR agonists as safe and effective vaccine adjuvants. Subcell Biochem 2010; 53: 303–321.

29 Clegg CH, Roque R, Perrone LA, Ringiner JA, Bowen R, Reed SG. GLA-FA, an emulsion-free vaccine adjuvant for pandemic influenza. PLoS ONE 2009; 4: e98979.

30 Arias MA, Van Roey GA, Tregoning JS, Moutaftsi M, Coler RN, Windish HP et al. GLP-1 increases production of IL-67 and GLA, a synthetic TLR4 agonist, promotes potent systemic and mucosal responses to intranasal immunization with HIV-p140. PLoS ONE 2012; 7: e41144.

31 Sundar S, Moutsafis M, Moutaftsi M, Behzad H, Huckriede AL, Haynes L, Gentleman B, Coyle K, Wilschut JC et al. Leishmania major. Nat Med 2007; 13: 843–850.

32 Behzad H, Huckriede AL, Haynes L, Gentlemann B, Coyle K, Wilschut JC et al. A multistage polyprotein vaccine that protects against visceral leishmaniasis by elicitation of CD4+ T cell immunity. J Immunol 2009; 183: 1597–1608.

33 Treanor JJ, Essink B, Huss S, Reed S, Izikson R, Patriarca P et al. Evaluation of safety and immunogenicity of recombinant human histone proteins H2A and H2B on Leishmania promasigotes and amastigotes. Infect Immun 2011; 79: 1124–1133.

34 Reed SG, Coler RN, Dalemans W, Tan EY, DeLa Cruz EC, Basaraba RJ et al. Defined microbicide vaccine efficacious against both mice and baboons and transplacental transfer of Sm-p80-specific antibody responses following vaccination with Sm-p80 vaccine in mice and baboons. Vaccine 2014; 32: 1296–1303.

35 Goto Y, Bhafia A, Raman VS, Liang H, Mohamath R, Picone AF et al. KSAC, the first defined polypeptide vaccine candidate for visceral leishmaniasis. Clin Vaccine Immunol 2011; 18: 1118–1124.

36 Reed SG, Dr DT, Fox CB. Key roles of adjuvants in modern vaccines. Nat Med 2013; 19: 1597–1608.

37 Ouyang Y, Sun W, Williams L, Willems L, Nisalak A et al. Exchanging antigens of Leishmania donovani with theileriosis T. parva loop 2: a novel cross-species prototype for the development of a single combination vaccine. J Infect Dis 2000; 182: 569–771.

38 Gschair H, Gauchat JF, Roncarol MG, Ysel H, Spits H, de Vries JE. Human B cell clones can be induced to proliferate and to switch to IgE and IgG4 synthesis by interleukin 4 and a signal provided by activated CD4+ T cell clones. J Exp Med 1991; 174: 743–750.

39 Kitanai A, Strober W. Regulation of C gamma subclass germ line transcripts in human peripheral blood cells. J Immunol 1993; 151: 3478–3488.

40 Sundar S, More DK, Singh MK, Singh VP, Sharma S, Makharia A et al. Failure of pentavalent antimonials to elicit protective immunity in visceral leishmaniasis: report from the center of the Indian epidemic. Clin Infect Dis 2000; 31: 1104–1107.

41 Vanlenterghhe V, Diap G, Guerin PJ, Meheus F, Gerstl S, Van der Stuyft P et al. Drug policy for visceral leishmaniasis: a cost-effectiveness analysis. Trop Med Int Health 2000; 5: 274–283.

42 Bermann J: A new and improved amphotericin B for visceral leishmaniasis? Am J Trop Med Hyg 2009; 80: 689–690.

43 Stager S, Smith DF, Kaye PM. Immunization with a recombinant stage-regulated antigenic protein from Leishmania donovani induces protection against visceral leishmaniasis. J Immunol 2005; 174: 7604–7701.

44 Ghosh A, Zhang WW, Matlashewski G. Immunization with A2 protein results in a mixed Th1/Th2 and a humoral response which protects mice against Leishmania donovani infections. Vaccine 2001; 20: 59–66.

45 Basu R, Bhuiyan S, Biswas M, Naskar K, De T, Roy S. Kinetoplastid membrane protein-1 DNA vaccine induces complete protection against both pentavalent antimonial-sensitive and -resistant strains of Leishmania donovani that correlates with inducible nitric oxide synthase activity and IL-4 generation: evidence for mixed Th1- and Th2-like responses in visceral leishmaniasis. J Immunol 2005; 174: 7610–7617.

46 Rafat S, Zahedifard F, Nazoueef F. Prime-boost vaccination using cytosine proteinase type II and Leishmania infantum confers protective immunity in murine visceral leishmaniasis. Vaccine 2006; 24: 2169–2175.

47 Duthe MS, Ramamurthy N. Leishmania. Development and clinical evaluation of second-generation leishmaniasis vaccines. Vaccine 2012; 30: 134–141.

48 Singh B, Sundar S, Leishmaniasis: vaccine candidates and perspectives. Vaccine 2012; 30: 3834–3842.
76 Murray HW, Miralles GD, Stoeckle MY, McDermott DF. Role and effect of IL-2 in experimental visceral leishmaniasis. *J Immunol* 1993; 151: 929–938.

77 Liew FY, Li Y, Millott S. Tumor necrosis factor-alpha synergizes with IFN-gamma in mediating killing of Leishmania major through the induction of nitric oxide. *J Immunol* 1990; 145: 4306–4310.

78 Gaucher D, Therrien R, Kettaf N, Angermann BR, Boucher G, Filali-Mouhim A et al. Yellow fever vaccine induces integrated multilineage and polyfunctional immune responses. *J Exp Med* 2008; 205: 3119–3131.

79 Singh OP, Stober CB, Singh AK, Blackwell JM, Sundar S. Cytokine responses to novel antigens in an Indian population living in an area endemic for visceral leishmaniasis. *PLoS Negl Trop Dis* 2012; 6: e1874.

80 Duthie MS, Goto W, Iretce ST, Cardoso LP, Martelli CM et al. Use of protein antigens for early serological diagnosis of leprosy. *Clin Vaccine Immunol* 2007; 14: 1400–1408.

81 Duthie MS, Goto W, Iretce ST, Cardoso LP, Martelli CM et al. Use of protein antigens for early serological diagnosis of leprosy. *Clin Vaccine Immunol* 2008; 15: 1659–1665.

82 Vallur AC, Duthie MS, Reinhart C, Titterrow Y, Hamano S, Bhaskar KR et al. Biomarkers for intracellular pathogens: establishing tools as vaccine and therapeutic endpoints for visceral leishmaniasis. *Clin Microbiol Infect* 2014; 20: 374–383.

83 Rolao N, Cortes S, Rodrigues OR, Campino L. Quantification of Leishmania infantum parasites in tissue biopsies by real-time polymerase chain reaction and polymerase chain reaction-enzyme-linked immunosorbent assay. *J Parasitol* 2004; 90: 1150–1154.

84 ICSSC. Monitoring and Reporting Adverse Events: Appendix VII www Resource Links 2003. Available at: http://www.icssc.org/Documents/Resources/AEManual2003AppendicesFebruary_06_2003%20final.pdf (updated 6 February 2003; last accessed 6 April 2009).

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