Combination Therapy Using Sodium Antimony Gluconate in Stearylamine-Bearing Liposomes against Established and Chronic *Leishmania donovani* Infection in BALB/c Mice

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In this work we report the activity seen with combination therapy using sodium antimony gluconate in liposomes composed of egg phosphatidyl choline and stearylamine for elimination of *Leishmania donovani* parasites from the liver and spleen of BALB/c mice with established and chronic infections.

Despite extended treatment regimens, parenteral administration, and toxic side effects, the pentavalent antimonial sodium antimony gluconate (SAG) has remained the first-line treatment for visceral leishmaniasis (VL) or kala-azar for decades (19, 33). Frequent therapeutic failures with SAG necessitate the use of more toxic second-line drugs, amphotericin B and pentamidine (22, 23, 25). Recently several lipid-based formulations of amphotericin B with reduced levels of toxicity have been used (9, 11, 32). However, no commercial lipid formulations of SAG are available even though liposome-encapsulated SAG is 200 to 700 times more active than free SAG (2, 5, 28). In most of these studies the efficacy demonstrated by neutral and negatively charged liposomal formulations of SAG was determined purely on the basis of liver parasite burdens (2, 5, 21). These formulations either failed to contain parasites in the spleen (6) or required multiple doses for a significant clearance in an acute-infection model (7, 13). The spleen is a site of persistent infection with *Leishmania donovani* (1, 12, 32), and parasite removal from this organ is more difficult than parasite removal from liver (6, 8, 31, 32). Earlier, Dey et al. reported on a multiple-dose treatment of drug-free, positively charged liposomes comprising egg phosphatidylcholine (PC) and stearylamine (SA) for the elimination of parasites from the liver and spleen of BALB/c mice with established *L. donovani* infection (10). Herein we evaluate a single-dose treatment with PC-SA liposome encapsulating a suboptimal concentration of SAG against an established and a chronic infection of *L. donovani* at both the sites of infection.

To optimize phospholipid composition the nonderivative L-α-PC was replaced with dimyristoyl (DMPC), distearoyl (DSPC), and dipalmitoyl (DPPC) derivatives of PC in combination with SA in a 7:2 molar ratio as detailed earlier (10). For drug encapsulation the lipid film was dispersed in 20 mM phosphate-buffered saline (PBS; pH 7.0) containing 1 mg of SAG (Glucanate Health Limited, Calcutta, India)/ml and sonicated for 30 s in an ultrasonicator. To remove free drug, liposomes with entrapped SAG were washed in PBS with centrifugation (10,000 × g, 30 min, 4°C) (15). Approximately 300 to 400 μg of SAG was associated with 22 mg of lipid, estimated as described previously (2, 17).

In vitro antileishmanial drug therapy was investigated as described earlier (10). Initially, four drug-free cationic liposomes, DSPC-SA, DMPC-SA, DPPC-SA, and PC-SA, each at 88 μg/ml, were tested against intracellular *L. donovani* amastigotes for optimum activity and minimum toxicity. Whereas DSPC-SA and DPPC-SA could not clear parasite infection by 96 h, DMPC-SA and PC-SA were almost equally effective at the time points studied (Fig. 1). However, while DMPC-SA induced 5% toxicity towards macrophages, as determined by lactate dehydrogenase release in supernatants (14) at 88 μg/ml, an identical dose of PC-SA exhibited no toxic effect. Hence,

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**FIG. 1.** Effect of different phospholipids with SA-bearing liposomes on the survival of *L. donovani* amastigotes in mouse peritoneal macrophages. Cells were infected with *L. donovani* promastigotes and incubated with 88 μg/ml concentrations of DPPC-SA (A), DSPC-SA (B), DMPC-SA (C), and PC-SA (D) or an equivalent amount of respective neutral liposomes or PBS for 0 to 96 h at 37°C. Data represent the means ± standard errors from triplicate coverglasses of one experiment representative of two performed.
PC-SA was selected for combination therapy with SAG. Subsequently, the antileishmanial activity of SAG-loaded PC-SA liposomes was compared with the results seen with empty liposomes and free SAG. Liposomal SAG therapy was found significantly more potent than either of the monotherapies (Fig. 2). The 50% effective doses for suppression of parasite infection in macrophages calculated using sigmoidal regression analysis (Microsoft Excel) for empty PC-SA and free SAG were 38.26 and 69.57 μg/ml, respectively. In contrast, 0.117 amastigotes per macrophage. The bars show the standard errors for three replicates and are representative of two independent experiments.

In vivo 8-week-infected BALB/c mice (10) were treated with a single dose of free SAG (16 mg/kg of body weight) or empty PC-SA (22 mg/mouse) or an equivalent dose of SAG entrapped in PC-SA (22 mg) (Fig. 3). Mice were sacrificed on days 1, 15, and 30 posttreatment, and levels of organ parasite burden were determined and expressed in Leishman Donovan units (1). Empty liposome or free SAG could suppress infection by 15 or 50%, respectively, only in the liver, with almost no reduction in the spleen 15 days posttreatment. On day 30, parasite suppression increased to 48% in liver and 60% in spleen (P < 0.05) with empty liposomes and to 76% in liver (P < 0.0001) and 65% in spleen (P < 0.05) with free SAG. Combination therapy was most effective, reducing the parasite burden by 89 and 98% in liver and 98% in spleen (P < 0.0001) after 1 month of treatment, and parasite survival (parasite per 100 cells) declined with less than 5% parasitemia in both organs.

Until recently splenectomy was the last recourse in cases of antimony resistance (25, 29). We thus examined the efficacy of combined treatment for mice after 3 months of infection, when splenic parasite load is well expressed (10). Interestingly, it was found that although the treatment efficiency declined with empty liposome and SAG to 16 and 50% (P < 0.05) in liver and 43 (P < 0.05) and 26% in spleen, respectively, at 30 days posttreatment, almost complete clearance of parasitemia from spleen (96%) and liver (98%), with a significant decrease in spleen weight (69%; P < 0.0001), was still achieved with the combination therapy in this model compared to the results seen with controls. Estimation of levels of serum alkaline phosphatase and glutamate pyruvate transaminase, specific enzymes related to liver dysfunction, and of levels of serum urea and creatinine, related to kidney dysfunction, 15 days postinjection of SAG entrapped in PC-SA liposome demonstrated levels within normal range, indicating no toxicity under the given conditions.

Herein we report a profound synergistic activity of SAG entrapped in PC-SA liposome in both in vitro and in vivo models of VL. To our knowledge this is the first report of a single dose of liposomal SAG treatment resulting in an almost complete elimination of parasites not only from the liver but also from the spleen of infected animals. An equivalent dose of free SAG or empty PC-SA liposome was only partially effective at clearing the intracellular amastigotes. Apart from a spin-
gomyelin-cholesterol-SA liposomal preparation of SAG, earlier formulations of SA-bearing liposomes demonstrated much less activity than negatively charged ones (2), resulting in follow-ups largely conducted with negatively charged (5, 21) and neutral (7, 28) vesicles. Although these formulations demonstrated spectacular success against liver parasites (2, 5, 6, 28), they failed to perform in spleens (6). There are considerable discrepancies in ideas as to how charges affect the interaction of liposomes with macrophages and their distribution in vivo. In contrast to previous observations (16, 20, 24), there are increasing reports of improved delivery of drug-entrapped positively charged liposomes to the liver as well as the spleen over the results seen with neutral and negatively charged vesicles (3, 18) and of better uptake by macrophages (27, 30) and effector cells infected with Leishmania donovani. Mol. Biochem. Parasitol. 124:59–60.

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