Development of nanocapsules bearing doxorubicin for macrophage targeting through the phosphatidylserine ligand: a system for intervention in visceral leishmaniasis

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Objectives: The purpose of this study was to explore the applicability, targeting potential and drug delivery to specialized phagocytes via phosphatidylserine (PS)-specific ligand-anchored nanocapsules (NCs) bearing doxorubicin.

Methods: The layer-by-layer method was utilized to prepare NCs having a nanoemulsion core loaded with doxorubicin (NCs-DOX), which was further grafted with PS. PS-coated NCs (PS-NCs-DOX) were compared with NCs-DOX for in vitro targeting ability by studying uptake by macrophages, intracellular localization, in vivo pharmacokinetics and organ distribution studies. The in vivo antileishmanial activity of free doxorubicin, NCs-DOX and PS-NCs-DOX was tested against visceral leishmaniasis in Leishmania donovani-infected hamsters.

Results: Flow cytometric data revealed 1.75-fold enhanced uptake of PS-NCs-DOX in J774A.1 macrophage cell lines compared with NCs-DOX. In vivo organ distribution studies in Wistar rats demonstrated a significantly higher extent of accumulation of PS-NCs-DOX compared with NCs-DOX in macrophage-rich organs, particularly in liver and spleen. Highly significant antileishmanial activity (P<0.05 compared with NCs) was observed with PS-NCs-DOX, causing 85.23%±4.49% inhibition of splenic parasitic burden. NCs-DOX and free doxorubicin caused only 72.88%±3.87% and 42.85%±2.11% parasite inhibition, respectively, in Leishmania-infected hamsters (P<0.01 for PS-NCs-DOX versus free doxorubicin and NCs-DOX versus free doxorubicin).

Conclusions: We conclude that the PS targeting moiety can provide a new insight for efficient drug delivery to specialized macrophages and thus may be developed for effective use in macrophage-specific delivery systems, especially for leishmaniasis.

Keywords: layer-by-layer assembly, polyelectrolytes, J774A.1 cell lines, L. donovani, amastigotes

Introduction

Leishmaniasis is a worldwide public health problem and is caused by various species of the genus Leishmania and is mainly associated with cutaneous, mucosal and visceral infections. It is characterized by fever and inflammation in the spleen and liver in humans.1,2 The disease is also associated with several pathophysiological lethal-cascading events without specific chemotherapy. Visceral leishmaniasis is caused by Leishmania donovani, which resides and multiplies within macrophages of the mononuclear phagocytic system (MPS). The MPS is of emerging importance both scientifically and therapeutically because of its crucial role in inflammation. For many diseases, such as asthma, atherosclerosis and cancer, and for pathogenic infections, including tuberculosis, filariasis and leishmaniasis, the inflammatory process is a key driver of both disease progression and pathogenesis.3–5 Therefore, the development of delivery systems that can target monocytes/macrophages intracellularly is crucial and could potentially open up new treatment paradigms for a range of diseases.

Macrophages express various engulfment receptors that are able to bind modified lipoproteins, senescent and apoptotic cells, proteins, polysaccharides and a range of polyanionic molecules.6 The clearance of apoptotic cells by macrophages is an
integral component of normal life. Several researchers have reported that phosphatidylserine (PS) is exposed on the outer surface of apoptotic cells and that recognition of PS is important for the resulting anti-inflammatory responses that are induced by macrophages. PS is a key ligand and is thought to exist in patches on apoptotic cells.

Recognition of PS involves multiple receptors, including some that can recognize naked PS and function together with other receptors that bind bridging protein bound to PS. Several receptors have been suggested to play a crucial role in mediating PS recognition, including scavenger receptors, CD68 [an oxidized low-density lipoprotein (oxLDL) receptor], CD14, annexins, β2 glycoprotein I and GAS6. Several groups of investigators have found that human and rodent macrophages, as well as insect phagocytes, preferentially take up negatively charged liposomes, particularly those containing PS. Currently available chemotherapeutic agents for leishmaniasis show potential toxicity and serious adverse effects, which ultimately lead to narrow therapeutic applications. Earlier antimonials were regarded as first-line therapy for leishmaniasis, but various pieces of evidence show resistance of Leishmania amastigotes to the antimonials. Doxorubicin, an anthracycline antibiotic used for the chemotherapy of various human cancers, was found to have potential antileishmanial activity, but its associated cardiac toxicity limits its clinical use. Aspects of the toxicity related to the delivery of doxorubicin can be reduced by its specific delivery to intracellular regions of macrophages. Various researchers have reported that incorporation of doxorubicin in liposomes and layer-by-layer (LBL)-assembled microparticles can increase the therapeutic index by reducing cardiotoxicity.

The above perspectives led us to explore the applicability, targeting potential and drug delivery to the macrophages via PS-specific ligand-anchored nanocapsules (NCs) bearing doxorubicin. We have developed NCs based on LBL technology having a nanoemulsion as a core, in which doxorubicin is loaded. The nanoemulsion was chosen as a core to improve the loading efficiency of doxorubicin, since free doxorubicin base has higher affinity for the internal oil phase, which restricts its entry into the external aqueous phase. Moreover, doxorubicin imposes steric hindrance, resulting in low entrapment in structured vehicles such as liposomes. The PS moiety provides an alternative means of targeting macrophages, and can easily be coated on cationic NCs due to its strongly anionic nature. Thus, in the present study we prepared and characterized PS-coated NCs loaded with doxorubicin (PS-NCs-DOX) and compared them with non-PS-coated NCs loaded with doxorubicin (NCs-DOX) for potential ability to target L. donovani parasites. Enhanced uptake can lead to smaller doses that are adequate for optimal therapeutic effect, thereby reducing the toxicity of the medication and providing a more efficient drug delivery system. PS-NCs-DOX are a candidate for macrophage-targeted drug delivery that can bind to macrophages in the liver and spleen and target the bioactive molecule inside them.

**Materials and methods**

**Materials**

PS (1,2-diacyl-sn-glycero-3-phospho-L-serine), sodium alginate (SA), proctamine sulphate (PRM) and poly(sodium 4-styrenesulphonate) (PSS), mol. wt 70000, were purchased from Sigma-Aldrich, USA. Tween 80 and Span 80 were obtained from HiMedia Laboratories Pvt Ltd, Mumbai, India. Soyabean oil (SBO) was purchased from Sunshine Oleochem Ltd, Gandhidham, India. Dialysis membrane (mol. wt cut-off 70 kDa) was obtained from Sigma-Aldrich, USA. The annexin V-FITC apoptosis detection kit was obtained from Calbiochem, CA, USA. For cell culture, Dulbecco’s modified Eagle’s medium (DMEM) with glutamate, fetal bovine serum (FBS) and antibiotic solution (penicillin/streptomycin, 0.1% v/v) were purchased from Sigma-Aldrich, USA. MT was also purchased from Sigma-Aldrich, USA. Well plates for cytotoxicity and uptake studies were from Greiner Bio-One (Frickenhausen, Germany). All materials were used without further purification. The water used in all experiments was prepared in a three-stage Millipore Milli-Q Plus 185 purification system (Bedford, MA, USA) and had a resistivity >18.2 mΩcm. All polyelectrolytes were used without further purification. All other chemicals and solvents were of analytical grade.

**Preparation of NCs containing nanoemulsion core**

The emulsion inversion point method, a low-energy emulsification method, was used to obtain a nanoemulsion core. It was obtained from a mixture of SBO (20 wt%) containing free doxorubicin base and weight ratio of Span 80/Tween 80 of ~0.43/0.57 by slowly adding aqueous phase containing PSS (0.025 wt%) at a rate of 1.0 mL/min with gentle agitation using a magnetic stirrer at a temperature of 60 °C. Free doxorubicin base was obtained using triethylamine-doxorubicin at a molar ratio of 3. Ten millilitres of an aqueous polyelectrolyte solution of PRM (0.125%, w/v) was injected into the pre-formed nanoemulsion core. Adsorption of the polyelectrolyte was allowed for 45 min, during which time the dispersion was occasionally stirred. The procedure was repeated to coat oppositely charged SA (0.150%, w/v). The dispersion was centrifuged for 15 min at 4000 rpm to separate the NCs from un-utilized polyelectrolyte aggregates. Finally, the layer of PRM was assembled on the SA layer.

**Coating of NCs with PS**

Coating of NCs with PS was initiated as described elsewhere with slight modification. Initially, PS was dissolved in chlorormethanol (19:1) and evaporated in a rotary evaporator at 50 °C and then hydrated with an aqueous dispersion of NCs for 20 min. The sample was then centrifuged at 40000 rpm for 30 min, after which the aqueous phase (lower layer) was withdrawn precisely to remove remaining lipid from the bulk. The PRM/PS weight ratio (process variable) was optimized by measuring the change in zeta-potential of the dispersion in triple-distilled water at 25 °C (Zetasizer 3000 HS; Malvern Instruments Co., UK).

**Characterization of NCs**

Developed formulations were characterized prior to and after surface ligand anchoring. For morphological characterization, transmission electron microscopy (TEM; 400T TEM, Philips, New Brunswick, Canada) studies were carried out using phosphotungstic acid as a negative stain. TEM was carried out to determine the surface characteristics of the optimized formulations in aqueous medium using a 3 mM formain (0.5% plastic powder in amyl acetate)-coated copper grid (300 mesh) at 60 kV using negative staining by 2% phosphotungstic acid at various magnifications. The presence of PS on the surface of NCs was confirmed by determining the change in zeta-potential and measuring the characteristic fluorescence of an FITC-conjugated PS-specific protein, annexin V, using a Cary Eclipse Fluorescence Spectrophotometer (Agilent Technologies, USA). Differential scanning calorimetry (DSC) was performed with a Diamond DSC (PerkinElmer, USA) to exclude the formation of PS vesicles during coating. Control measurement was performed by using a pure
dispersion of PS vesicles with a concentration of 1 mg/mL. The mean particle size and size distribution of NCs and PS-NCs were determined using a Zetasizer Nano-ZS (Malvern Instruments, Germany). Aliquots (100 µL) containing J774A.1 (1 × 10⁶) cells were suspended in 0.9 mL of fresh RPMI-1640 medium (Himedia) supplemented with 10 U/mL penicillin, 10% FBS, 100 µg/mL of streptomycin, 1 mM sodium pyruvate and 10 mM HEPES medium. These were transferred into 24-well plates (Sigma, Germany) containing fresh medium and suspended in a 37°C humidified incubator with a 5% CO₂ atmosphere. After 24 h the culture medium was replaced with fresh culture medium containing NCs-DOX and PS-NCs-DOX formulations. To separate the internalized and surface-bound NCs, the cells were washed three times with acetate buffer (pH 4.0). The cell-associated fluorescence was measured by FACS at an excitation wavelength of 480 nm and an emission wavelength of 550 nm. Macrophage uptake of PS-NCs-DOX was also assessed by confocal laser scanning microscopy (CLSM). Cells seeded at a density of 1 × 10⁶ cells/well on poly-l-lysine coated glass cover slips (in six-well plates) were left for 12 h, and then adherent macrophage monolayers were washed and incubated in fresh medium. PS-NCs-DOX were suspended in DMEM and added to macrophages and incubated for 4 h at room temperature in the dark. After incubation, monolayers were washed repeatedly to remove non-adherent PS-NCs-DOX and cells were fixed in 10% formalin (in PBS) and the cell nuclei were stained with 4',6-diamidino-2-phenylindole (DAPI). Macrophages were observed by CLSM (Olympus IX 81, Center Valley, PA, USA) equipped with a ×60 oil objective lens.

**In vitro evaluation against intra-macrophage amastigotes of L. donovani**

The J774A.1 macrophage cell line was used for the in vitro intracellular drug efficacy test. The macrophages were resuspended at 2.5 × 10⁵ cells/mL in serum-free RPMI-1640. Cell suspensions (200 µL/well) were plated on eight-chamber Lab-Tek tissue culture slides (Nunc, USA) and were allowed to adhere for 2 h in a CO₂ incubator with 5% CO₂ at 37°C. The wells were washed twice with serum-free medium, and the adherent macrophages were infected with metacyclic-stage parasites of L. donovani, maintaining a Leishmania:macrophage ratio of 10:1 in a 200 µL final solution of a complete medium, and incubated overnight. After 24 h, free promastigotes were washed with serum-free medium and infected macrophages were incubated with medium containing free doxorubicin, NCs-DOX and PS-NCs-DOX for 48 h at 37°C at various concentrations. Untreated infected macrophages served as control. The slides were fixed with methanol and stained with Giemsa. At least 100 macrophage nuclei were counted per well for calculating the percentage of infected macrophages and the number of amastigotes per 100 macrophages. Percentage parasite inhibition in treated wells was calculated using the formula 100 – (T × 100)/C, where T is the number of parasites in treated samples/100 macrophage nuclei, and C is the number of parasites in the control. The IC₅₀ (concentration of drug that inhibits 50% of L. donovani amastigotes) was obtained by plotting a graph of the percentage of inhibition at different concentrations of the formulations using Origin 6.1 version software, and was expressed in ng/mL.

**Cytotoxicity studies**

The in vitro cytotoxicity of blank formulations was measured with the MTT proliferation assay. The experiments were carried out on cells in the exponential growth phase. J774A.1 and RAW macrophage cells were seeded into 24-well plates at 5 × 10⁴ cells/well and were allowed to adhere overnight. The growth medium was replaced with fresh medium and then cells were incubated for 24 h with different blank formulations (NCs and PS-NCs). Cells were then washed twice with 1 mL of PBS. Cells were then incubated in a growth medium containing 1 mg/mL of MT1 for 4 h at 37°C, and then 500 µL of DMSO was added to each well to ensure solubilization of the formazan crystals. The optical density was measured using a multi-well scanning spectrophotometer (MRX Microplate Reader, Dynatech Laboratories Inc., Chantilly, VA, USA) at a wavelength of 570 nm.

**Animal host**

Laboratory bred male Syrian golden hamsters (Mesocricetus auratus, 45–50 g, for in vivo antileishmanial activity) and Wistar rats (150–200 g, for pharmacokinetic analysis) from the animal house facility of the Central Drug Research Institute (CDRI) were used as the experimental host after approval from the Institutional Animals Ethical Committee (IAEC) of CDRI (IAEC approval number 70/11/Pharmaceutics/IAEC). The study was carried out under the guidelines of the Council for the Purpose of Control and Supervision of Experiments on Animals (PCPSEA), Ministry of Social Justice and Empowerment, Government of India.

**Plasma and tissue sample analysis**

Wistar rats weighing ~150–200 g were divided into three groups of three rats each. Free doxorubicin, NCs-DOX and PS-NCs-DOX containing equivalent doses of doxorubicin (5 mg/kg of body weight) were administered intravenously to different groups. Animals from each group were sacrificed 0.5, 1, 2, 4, 8, 12 and 24 h after administration of the formulations. Blood samples were collected by cardiac puncture. The liver and spleen of the dissected rats were excised, isolated, washed with distilled water and blot dried using tissue paper. Doxorubicin was extracted from 200 µL of each plasma and tissue sample (liver and spleen, 100 mg homogenized in 1 mL of phosphate buffer) using 250 µL of a 50:50 (v/v) mixture of methanol and 40% ZnSO₄. After 5 min of vigorous stirring followed by centrifugation, the organic phases were collected and evaporated to dryness at 30°C under a flow of nitrogen. Dry residues from plasma and tissues were dissolved in 100 µL of mobile phase and 50 µL of the resulting solution was injected into the chromatograph. The samples were analysed by reverse-phase HPLC as reported previously with slight modification. In brief, the HPLC system (CLASS-VP, Shimadzu Corporation, Kyoto, Japan) consisted of a binary gradient pump (LC-20AT), fluorescence detector (RF-10AxL) and system controller (SCL-10A). The CLASS-VP workstation was used for data acquisition. The analytical C18 column (LChrompher, 5 µm, 250 × 4 mm) was used for the chromatographic separation of doxorubicin. The mobile phase consisted of a mixture of acetonitrile and water (35:65, v/v) with the pH adjusted to 2.05 using orthophosphoric acid, and detection was carried out at an excitation and emission wavelength of 480 and 550 nm, respectively, at a flow rate of 1 mL/min. The recovery of doxorubicin was determined by comparing the responses of the pre-extracted standard versus responses of post-extracted plasma and tissue standards at equivalent concentration.

**In vivo antileishmanial activity**

**Parasites**

Promastigotes of the WHO reference strain, L. donovani (M HO-M/IN/ 80/Dd8), were cultured in RPMI-1640 medium (Sigma-Aldrich, USA) supplemented with 10% FBS (Sigma-Aldrich, USA), penicillin (100 U/mL) and streptomycin (10 µg/mL) in a humidified incubator with a 5% CO₂ atmosphere at 37°C.

**Drug Research Institute (CDRI) were used as the experimental host after approval from the Institutional Animals Ethical Committee (IAEC), Ministry of Social Justice and Empowerment, Government of India.**
streptomycin (100 mg/mL) at 26°C. Parasites were also maintained through in vivo serial passages (amastigote to amastigote) in hamsters.²⁹,³⁰

Antileishmanial activity testing
The isolation of parasites and infection of naive hamsters were carried out as described previously.³¹ Briefly, infected animals with a 40–60-day-old infection were autopsied and the spleen was removed aseptically, homogenized in PBS and centrifuged at 900 rpm for 5 min at 4°C to sieve out tissue debris. Supernatant was centrifuged at 3500 rpm for 10 min. The pellet was washed twice with PBS and resuspended to obtain a concentration of 1×10⁷ amastigotes per 100 μL of PBS and this inoculum was injected into each hamster intracardially. After 25 days, infection was confirmed by Giemsa staining of tissue smears of spleen after the biopsy of two animals. Infected animals harbouring 38–40 amastigotes/100 macrophage nuclei were then distributed for drug treatment in the following manner: (i) infected controls (no therapy was given); (ii) free doxorubicin; (iii) NCs-DOX; (iv) PS-NCs-DOX; (v) blank NCs; and (vi) blank PS-NCs.

Infected hamsters were treated intraperitoneally with 250 μg/kg/day of doxorubicin in all formulations for four consecutive days. After 15 days post-treatment, splenic biopsies were performed and the parasite burden was determined by counting the number of amastigotes.³²

Measurement of haemolysis
The degree of haemolysis induced by NCs and PS-NCs on fresh citrated human blood was determined.³³ Red blood cells (RBCs) were obtained from male Wistar rats by centrifugation (1500 rpm, 10 min), washed three times with isotonic 120 mM phosphate sodium chloride buffer (pH 7.4) and suspended in the same buffer at a concentration equivalent to 5% of the normal haematocrit; 40 μL aliquots of this RBC suspension were added to 1 mL of NCs and PS-NCs, incubated at 37°C for 2 h and then centrifuged at 10000 rpm for 20 min. The haemoglobin released in the supernatant was measured spectrophotometrically at 550 nm. The absorbance (A) at this wavelength was directly proportional to the haemoglobin concentration, which was in turn proportional to the number of intact RBCs that were not destroyed by the added sample. For the control, blood was similarly incubated with 0.9% (w/v) saline. Haemolysis induced with hypertonic saline (A₂) was taken as 100%. The amount of haemoglobin released into the supernatant by the sample was recorded as A₁ and the percentage of haemolysis was calculated as A₂−A₁/A₂×100.

Statistical analysis
All results are given as means±SD (n=3). Differences between formulations were compared using one-way analysis of variance (ANOVA) followed by the Tukey–Kramer multiple comparison test, using InStat software (GraphPad Software Inc., CA, USA). P<0.05 denotes significance in all cases.

Results
Preparation of NCs
The low-energy emulsification method was used to prepare nanoemulsion cores and to deposit additional polyelectrolytes on the surface to form a three-layered shell. This procedure represents an innovative and preferable means for the production of polyelectrolyte NCs. In the first step, a nanoemulsion core (oil in water) was obtained from a mixture of SBO (20 wt%) and a ratio of surfactants (Span 80/Tween 80) of ≏0.43/0.57 by gradual addition of the aqueous phase containing PSS at the rate of 1 mL/min with gentle agitation using a magnetic stirrer. The temperature of the mixture was maintained at 60°C. In the second step, sequential addition of PRM and SA to nanoemulsion cores led to the formation of polyelectrolyte NCs in LBL fashion. The rigidity of the shell and the zeta-potential were also found to be altered at each layering of PRM and SA. Figure 1(a and b) shows the morphological characteristics of the nanoemulsion core and polyelectrolyte NCs by TEM. Polyelectrolyte layering was observed as a continuous opaque layer (Figure 1)(b) over a nanoemulsion core (Figure 1)(a).

Entrapment efficiency in NCs was found to be 81.37±3.76%. The PS coating on NCs did not significantly affect entrapment efficiency. The nanoemulsion core was negatively charged, initially with a zeta-potential of −34.12±2.41 mV due to PSS, and mean diameter of 322±4.31 nm (polydispersity index (PDI) 0.13±0.02). The first PRM layer reversed the charge to (+)42.11±3.22 mV, while the

Figure 1. Visualization of nanoemulsion core (a) and polyelectrolyte NCs (b) by TEM. The outer dark layer continuously covering the nanoemulsion core in polyelectrolyte NCs is clearly visible.
subsequent SA layer reversed the charge to \((-35.64 \pm 3.09 \text{ mV})\). The final PRM layer assembled on the SA layer changed the zeta-potential value finally to \((+36.42 \pm 2.48 \text{ mV})\). Depending on the type of polyelectrolyte added, the zeta-potential altered at every layering, confirming LBL assembly over the nanoemulsion core.

Table 1 shows the particle sizes and size distributions of different formulations measured by differential light scattering. The results indicate that the mean diameter of NCs was \(356 \pm 2.12 \text{ nm} \) (PDI \(0.22 \pm 0.03\)).

**NCs coated with PS-terminating ligand**

The NC formulation was finally coated with PS by hydrating the phospholipid film with an NC dispersion. The concluding PRM layer on the NCs facilitated spreading of PS uniformly over the NCs (Figure S1, available as Supplementary data at JAC Online), which was confirmed by a change in the zeta-potential to \((-13.23 \pm 2.16 \text{ mV})\). The process variable, i.e. the PS/PRM weight ratio, was optimized by measuring the change in zeta-potential of the dispersion in triple-distilled water at 25°C (Zetasizer 3000 HS; Malvern Instruments Co.) (Figure S2, available as Supplementary data at JAC Online). No change in zeta-potential was observed above a PS:PRM weight ratio of 0.05:0.1; this ratio was therefore selected as the optimum for preparing PS-NCs. Additionally, the presence of PS on the surface of NCs was verified by using an FITC-conjugated PS-specific protein, annexin V, by measuring its characteristic fluorescence. After treatment of PS-NCs with FITC-conjugated annexin V, robust fluorescence was visible. To rule out the formation of PS vesicles during coating on NCs, DSC was performed since phase transition is an important characteristic of lipid layers. It is evident from the thermogram (Figure S3, available as Supplementary data at JAC Online) that PS assembled on NCs observed a phase transition, which was 10°C below the phase transition of PS vesicles suspended in water. This lower value of phase transition temperature of PS assembled on NCs could be attributed to the disordering effect of polyelectrolyte molecules due to mutual interaction between polyelectrolyte and lipid headgroups. Table 1 shows the particle size and size distributions of NCs and PS-NCs measured by differential light scattering. The results indicate that the mean diameters of NCs and PS-NCs were \(356 \pm 2.12 \text{ nm} \) (PDI \(0.22 \pm 0.03\)) and \(371 \pm 1.02 \text{ nm} \) (PDI \(0.15 \pm 0.03\)), respectively.

**In vitro uptake studies**

Figure 2 shows the macrophagic uptake of various NC formulations in the J774A.1 macrophage cell line as shown by flow cytometry. This study represents the comparative uptake of free doxorubicin, NCs-DOX and PS-NCs-DOX. Figure 2 shows almost 3-fold \((>2.9)\) increased uptake of NCs-DOX compared with free doxorubicin, while PS-NCs-DOX showed almost 2-fold \((>1.75)\) increased uptake in J774A.1 cell lines compared with NCs. The typical confocal microscopic pictures shown in Figure 3 illustrate the distribution of internalized PS-NCs-DOX within cells. Most of the capsules are internalized within cells, as was manifest from the merged confocal image, showing the co-localization of red fluorescent signal arising from doxorubicin and blue fluorescence arising from DAPI, a nucleus-staining dye. This figure appears in colour in the online version of JAC and in black and white in the print version of JAC.

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**Table 1.** Characterization of NC and PS-NC formulations

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Charge (mV)</th>
<th>Size (nm)</th>
<th>Entrapment efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCs</td>
<td>(+36.42 \pm 2.48)</td>
<td>(356 \pm 2.12)</td>
<td>(81.37 \pm 3.76)</td>
</tr>
<tr>
<td>PS-NCs</td>
<td>((-13.23 \pm 2.16)</td>
<td>(371 \pm 1.02)</td>
<td>(80.13 \pm 6.43)</td>
</tr>
</tbody>
</table>

Values are expressed as means ± SD.
In vitro evaluation against intra-macrophage amastigotes of *L. donovani*

Inhibition of amastigote multiplication within macrophages by free doxorubicin, NCs-DOX and PS-NCs-DOX was recorded. All samples were stained and the number of infected macrophages was determined microscopically. Both NCs-DOX and PS-NCs-DOX showed significant improvement ($P<0.01$) in efficacy in comparison with free doxorubicin. Similarly, PS-NCs-DOX showed a significant improvement ($P<0.05$) in efficacy in comparison with NCs-DOX (Table 2).

**Pharmacokinetics of doxorubicin formulations in rats**

The plasma concentration–time profile of different doxorubicin formulations in rats at a dose of 5 mg/kg is presented in Figure 4. High mean residence time (MRT) and lower clearance was observed in PS-NCs-DOX compared with NCs-DOX in liver and spleen. The pharmacokinetic parameters of free doxorubicin, NCs-DOX and PS-NCs-DOX are given in Table 3. The extraction recovery of doxorubicin from plasma was 93.47% ± 2.89%.

**Table 2. In vitro activity of doxorubicin formulations against *L. donovani* intra-macrophagic amastigotes**

<table>
<thead>
<tr>
<th>Formulation</th>
<th>IC$_{50}$ (ng/mL)</th>
<th>Level of significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>A free doxorubicin</td>
<td>691.67 ± 10.41</td>
<td>$P&lt;0.01$ $P&lt;0.01$</td>
</tr>
<tr>
<td>B NCs-DOX</td>
<td>391.67 ± 20.40</td>
<td>$P&lt;0.05$</td>
</tr>
<tr>
<td>C PS-NCs-DOX</td>
<td>354.33 ± 5.13</td>
<td></td>
</tr>
</tbody>
</table>

Results are expressed as inhibition of parasite growth (IC$_{50}$) observed after 48 h of incubation; values are expressed as means ± SD ($n=3$).

Liver and spleen distribution of different doxorubicin formulations

The doxorubicin concentrations in liver and spleen after intravenous injection of different doxorubicin formulations are shown in Figure 5. PS-NCs-DOX and NCs-DOX showed significantly higher localization of doxorubicin at all timepoints compared with free doxorubicin ($P<0.05$). Similarly, PS-NCs-DOX showed enhanced drug distribution compared with NCs-DOX in liver and spleen. The extraction recoveries of doxorubicin from liver and spleen were 91.36% ± 3.12% and 88.34% ± 5.38%, respectively.

In vivo antileishmanial activity testing

There was significant improvement in activity with PS-NCs-DOX when compared with NCs-DOX ($P<0.05$). PS-NCs-DOX caused 85.23% ± 4.49% inhibition of the splenic parasitic burden, whereas NCs-DOX and free doxorubicin caused only 72.88% ± 3.87% and 42.85% ± 2.11% parasite inhibition, respectively, in *Leishmania*-infected hamsters ($P<0.01$ for PS-NCs-DOX versus free doxorubicin and NCs-DOX versus free doxorubicin). The parasite inhibition with blank PS-NCs and NCs was found to be 18.73% ± 2.79% and 13.8% ± 4.37%, respectively, as shown in Figure 6 ($P<0.05$ for PS-NCs versus NCs).

Cytotoxicity studies

The cytotoxicity of the different formulations was determined by the MTT assay. NCs showed 96% and 94% while PS-NCs showed 94% and 93% cell viability after 6 h of incubation in J774A.1 and RAW macrophagic cell lines, respectively.

Measurement of haemolysis

Blank NCs and PS-NCs showed 6.92% ± 3.34% and 5.82% ± 1.71% haemolytic activity, respectively, after 2 h of incubation.

**Discussion**

Chemotherapy of leishmaniasis primarily consists of the administration of antimonials [sodium stibogluconate (Pentostam) or N-methylglucamine antimoniate (Glucantime)], but responses to these drugs are abating due to the emergence of resistance, which causes frequent relapses. Pentamidine and amphotericin B have been the second-line treatment of leishmaniasis after antimony failure; however, their application is greatly limited due to their toxic manifestations.

There is a great requirement for new therapeutics due to the increasing financial burden arising from drug-resistant *Leishmania* strains. Doxorubicin, an anthracycline antibiotic used for the chemotherapy of various human cancers, is emerging as a potential antileishmanial agent with acceptable toxicity at limited doses. Attempts are therefore being made to develop a novel delivery system with a high payload of chemotherapeutic agent that is affordable and stable and reduces the obstacle of toxicity. Production of NCs by the LBL method has numerous therapeutic benefits, such as readily tailored physical properties (e.g. size, composition, porosity, stability, surface functionality and colloidal stability), high payload efficiency, controlled and targeted release at the site of infection, and...
increased stability, solubility and pharmacokinetic profile, over other nanoarchitectures. Apart from this, there is the possibility of numerous therapeutic benefits over liposomes, such as resistance to physiological stress and the possibility of oral delivery.

There are several reports of the use of inorganic core particles to fabricate LBL-based capsules. It has been observed by our group that preparing inorganic core particles, i.e. CaCO$_3$ and Ca(PO$_4$)$_2$, leads to the formation of several polymorphs, which hampers the formation of uniform layering of polyelectrolytes over the core. Similarly, the particle size of the core particle remains in the range of 3–5 μm, which may not be suitable for various applications. In the present investigation, we used a nanoemulsion with a core particle size range of 200–300 nm.

We used SA and PRM for the preparation of polyelectrolyte NCs. Both these polyelectrolytes are biocompatible and biodegradable. Drug incorporation in nanoparticles of biocompatible polymers has an advantage over other systems due to ease of preparation, longer shelf life and greater stability in biological fluids without any adverse effects. However, to prepare polyelectrolyte-based NCs in LBL fashion, the core particle must have some initial charge. We therefore incorporated PSS due to its surface adsorbing properties along with its ability to impart a highly negative charge to the nanoemulsion core. This initial negative charge facilitates the deposition of cationic PRM as a first layer. Subsequently, oppositely charged SA and PRM were used to develop NCs. The method of complex coacervation was used to form a capsule shell consisting of PRM-SA-PRM layers interlinked as a polyelectrolyte complex. The alternate coating of PRM and SA over the nanoemulsion core was confirmed by reversal of the zeta-potential at each layering. TEM images show

Table 3. Pharmacokinetic parameters following intravenous administration of various formulations of doxorubicin in rats

<table>
<thead>
<tr>
<th>Formulation</th>
<th>MRT (h)</th>
<th>$C_{\text{max}}$ (μg/mL)</th>
<th>Clearance (mL/h/kg)</th>
<th>AUC (h·μg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free doxorubicin</td>
<td>1.41 ± 0.29</td>
<td>33.16 ± 4.43</td>
<td>13.14 ± 1.24</td>
<td>74.55 ± 3.14</td>
</tr>
<tr>
<td>NCs-DOX</td>
<td>8.13 ± 0.13</td>
<td>75.26 ± 3.10</td>
<td>1.03 ± 0.03</td>
<td>779.24 ± 9.81*</td>
</tr>
<tr>
<td>PS-NCs-DOX</td>
<td>9.03 ± 3.14</td>
<td>108.37 ± 2.55</td>
<td>0.60 ± 0.04</td>
<td>1056.70 ± 4.74**</td>
</tr>
</tbody>
</table>

Values are expressed as means ± SD. Significance versus free doxorubicin: *$P<0.01$. Significance for PS-NCs-DOX versus NCs-DOX: **$P<0.01$. 

Figure 5. Percentage of dose recovered for different doxorubicin formulations in liver (a) and spleen (b). Data show the effect of NCs-DOX and PS-NCs-DOX on hepatic and splenic uptake of doxorubicin. PS-NCs-DOX show significantly enhanced uptake in comparison with NCs-DOX ($P<0.05$).
positively charged vesicles are formed in the presence of charged NCs (Figure S1). However, it has been reported that if on the surface. The phospholipid PS has a net negative charge, of PRM as an outermost layer, the NCs had a net positive charge over a plain nanoemulsion core (Figure 1)(a). Due to the presence that there was an opaque layer of polyelectrolytes (Figure 1)(b) of PS (beyond a PS:PRM weight ratio of 0.05:0.1) there a positive charge has been camouflaged by PS. On further add-

Figure 6. Antileishmanial activity of doxorubicin formulations against established infection of L. donovani in hamsters. Drug formulations (equivalent to 250 μg/kg/day for 4 consecutive days) and formulations without drug were injected intraperitoneally into each hamster on day 31 post-infection. The parasite burden was estimated by splenic biopsy on day 15 post-treatment and percentage parasite inhibition was calculated in comparison with the parasite burden of untreated infected animals (means ± SD) (n=3). The mean parasite burden in the spleen of untreated, infected control animals was 4.13 ± 45.90 amastigotes per 100 cell nuclei of macrophages. *P<0.05 (comparison of PS-NCs-DOX versus NCs-DOX and blank PS-NCs versus NCs), **P<0.01 (comparison of PS-NCs-DOX versus free doxorubicin and NCs-DOX versus free doxorubicin).

evident, indicating the presence of PS on the surface of NCs. It has been reported that annexin V specifically binds to PS. A flow cytometry study revealed almost doubled (≈1.75) uptake in J774A.1 macrophage cells by PS-NCs-DOX compared with NCs-DOX (Figure 2). The higher uptake of PS-NCs-DOX was confirmed by a CLSM study showing complete internalization of PS-NCs-DOX in macrophages (Figure 3). This significantly higher efficacy of PS-NCs-DOX in vitro (P<0.05) compared with NCs-DOX (Table 2) may be attributed to the facilitated uptake of NCs due to the presence of the specific ligand PS. It has been reported that macrophages have multiple receptors for the recognition of PS, which results in prolonged residence of doxorubicin inside macrophages.8 This higher uptake of PS-NCs-DOX compared with NCs-DOX in macrophages is of interest in the sense that there is scope for the enhanced localization of these modified NCs by intramacrophase amastigotes in liver and spleen, which may be significant for visceral leishmaniasis. Pharmacokinetic parameters showed that the AUC of PS-NCs-DOX was increased 14.17- and 1.35-fold, while the MRT was increased 6.39- and 1.11-fold compared with free doxorubicin and NCs-DOX, respectively, with a corresponding decrease in clearance. Similarly, significantly higher MRT and lower clearance with PS-NCs-DOX indicates effi-
cient macrophage targeting compared with NCs-DOX. The MRT of NCs-DOX and PS-NCs-DOX was found to be 8.13 ± 0.13 and 9.03 ± 3.14 h, respectively. The complex pharmacokinetics of PS-NCs-DOX might be attributed to its restricted distribution in the body due to the presence of the highly specific ligand PS on the surface of NCs. In our experimental conditions it seems there is non-linear clearance with saturation of the reticuloendothelial system, such as the liver and spleen, and redistribution of doxorubicin into the circulation. The data are consistent with the non-linear saturation-like kinetic profile, resulting in sustained plasma levels and decreased clearance. Apart from this, higher unspecific serum protein binding of the PS-NCs-DOX may also be responsible for its lower clearance. This complex pharmacokinetics may improve pharmacotherapy using doxorubicin due to reduction of possible adverse events because encapsulated doxorubicin will not be available directly to the other body tissues. Greater localization of doxorubicin by PS-NCs-DOX compared with NCs-DOX in both the liver and the spleen confirmed that most of the drug was rapidly localized in the liver and spleen due to enhanced uptake by macrophages (Figure 5).

Several receptors are involved in the recognition of PS, including some that can recognize naked PS and function together with additional receptors that bind bridging protein-bound PS.8 There are bridging molecules, termed GAS6 and Protein S, that can bind PS and in turn be recognized by engulfment receptors expressed by phagocytes.68,69 Similarly, there is another soluble bridging molecule, termed MFG8 (milk fat globule EGF factor 8 protein), secreted by macrophages, which then engulf MFG8-coated apoptotic cells (PS is exposed on the outer leaflet of the cells) through αvβ3-integrin.60 There is also another membrane protein expressed by macrophages, namely Ba1 (brain angiogenesis inhibitor 1), that can directly recognize PS exposed on apoptotic cells.80 Several receptors have been suggested to play a pivotal role in mediating PS recognition, including scavenger receptors (class B, CD36/ SR-BI; and class A, MARCO),82 CD68 (an oxLDL receptor), CD14, annexins, β2
glycoprotein I and GAS6.9 Tempone et al.53 described the macrophage-targeting ability of PS liposomes towards scavenger receptors and developed drug-loaded PS liposomes that were found to accumulate to a greater degree in macrophages compared with the free drug.

Macrophagic uptake of PS appended to NCs opens a new area for targeting drug-loaded carrier systems, which was confirmed by the significantly improved in vivo antileishmanial activity of PS-NCs-Dox when compared with NCs-Dox (P<0.05). Our data show that PS-NCs could be a useful carrier for targeting antileishmanial drugs into macrophages, and, interestingly, PS-NCs also show antileishmanial activity by themselves. Since PS-anchored NCs act as a signal for engulfment by phagocytes, they are specifically recognized by macrophages and are phagocytosed, which subsequently leads to macrophage activation. There are several reports that demonstrate that macrophage activation leads to rapid parasite killing and digestion.54 It has also been reported that up-regulation of inducible nitric oxide synthase (iNOS) and subsequent nitric oxide (NO) production are important for the parasiticidal activity of infected macrophages during experimental infection.54,55 The polyelectrolytes PRM and SA have been shown to induce the iNOS pathway, which results in subsequent production of NO.56,57

The primary objective of our work was to employ the naturally occurring ‘eat me’ signal, PS, as a specific ligand that is recognized by phagocytes. The superior efficacy of the encapsulated doxorubicin in eliminating intracellular amastigotes of L. donovani in both an in vitro macrophage model and an in vivo hamster model of visceral leishmaniasis demonstrates the effectiveness of this approach. Results clearly indicate a substantial enhancement of the efficacy of PS-NCs-Dox compared with NCs-Dox due to anchoring of PS.

To evaluate the safety profile of formulations on cells, a cell viability study was carried out in J774A.1 and RAW macrophage cells. The almost negligible cytotoxicity of both NCs and PS-NCs in J774A.1 and RAW macrophagic cells confirms that these excipients are safe to use. To evaluate the safety profile of the developed formulation, the haemolytic activity was measured with different NC formulations. Haemolysis was <6% when tested with both formulations, suggesting no deleterious effects on erythrocytes.

Conclusions

The present investigation provides evidence that the prototype formulation may be used for efficient treatment of leishmaniasis via PS-specific ligand-anchored NCs bearing doxorubicin. We report that PS-NCs provide an ‘eat me’ signal for specialized phagocytes because recognition of PS involves multiple receptors present on the phagocyte and causes enhanced internalization of the bioactive molecule into cells. Enhanced uptake directly reduces the dosage of the formulation, which is highly desirable for optimizing the therapeutic effect and reducing the side effects of doxorubicin.

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Transparency declarations

None to declare.

Supplementary data

Figures S1–S3 are available as Supplementary data at JAC Online (http://jac.oxfordjournals.org/).

References


