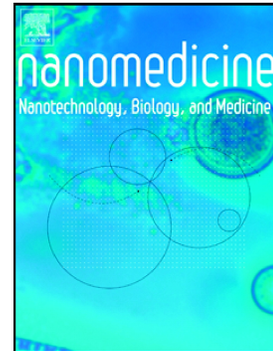


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ORIGINAL RESEARCH

Encapsulation in lipid-core nanocapsules improves topical treatment with the potent antileishmanial compound CH8.

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ABSTRACT

Cutaneous leishmaniasis (CL) is a neglected parasitic disease conventionally treated by multiple injections with systemically toxic drugs. Aiming at a more acceptable therapy, we developed lipid-core nanocapsules (LNC) entrapping the potent antileishmanial chalcone (CH8) for topical application. Rhodamine-labeled LNC (Rho-LNC-CH8) was produced for imaging studies. LNC-CH8 and Rho-LNC-CH8 had narrow size distributions (polydispersity index < 0.10), with similar mean sizes (~180 nm) by dynamic light scattering. *In vitro*, Rho-LNC-CH8 were rapidly internalized by extracellular *Leishmania amazonensis* parasites macrophages in less than 15 minutes. LNC-CH8 activated macrophage oxidative mechanisms more efficiently than CH8, and was more selectively toxic against the intracellular parasites. *In vivo*, topically applied Rho-LNC-CH8 efficiently permeated mouse skin. In *L. amazonensis*-infected mice, LNC-CH8 reduced the parasite load by 86% after three weeks of daily topical treatment, while free CH8 was ineffective. In conclusion, LNC-CH8 has strong potential as a novel topical formulation for CL treatment.

Keywords: Leishmaniasis, nitrochalcone, lipid-core nanocapsules, drug delivery, skin, topical treatment.

BACKGROUND

Cutaneous leishmaniasis (CL) is the most common clinical form of the disease, accounting for an estimated 1.2 million cases per year, mainly in developing countries¹. After inoculation of flagellated promastigotes into the skin by blood-feeding sandflies, successful parasites invade dermal macrophages, transform into amastigote forms, and survive and multiply intracellularly by downregulating host cell defense mechanisms such as the production of reactive oxygen species (ROS) and nitric oxide (NO), and proteolytic digestion inside the parasitophorous vacuole². Chronic and disfiguring ulcers develop at the infection site, and may spread to other parts of the skin.

Although the infection is limited to the skin, current CL chemotherapy involves long-term intravenous or intramuscular injections with pentavalent antimonials (Pentostam[®] or Glucantime[®]), pentamidine or amphotericin B. The injections are painful, systemically toxic and result in high hospital costs.³ The only oral antileishmanial drug – miltefosine – is used mostly for visceral leishmaniasis in the Old World, and is not approved for use in New World countries such as Brazil. To circumvent these difficulties, an effective topical treatment is the obvious choice for CL. However, creams containing paromomycin, miltefosine and amphotericin B have shown partial or no efficacy against CL, in particularly against New World *Leishmania* species⁴⁻⁶. Failure of topical formulations is typically linked to poor skin permeation, mainly to high drug molecular size and/or inappropriate lipophilicity⁷. The addition of strong permeation enhancers to topical formulations allows drugs to reach the deep dermis; however, most permeation enhancers cause unacceptable irritation at

the treatment site⁸. Thus, the present scenario justifies the search for new antileishmanial agents and formulations, particularly for topical CL treatment.

Due to the inadequacy of current CL chemotherapy, the Drugs for Neglected Disease initiative (DNDi) has supported the search for new topical or oral treatments for CL that are efficacious against all species, easy to administer, can be given in as a short treatment course (14-28 days), and are compatible with combination therapy and adapted to tropical climates⁹. We have previously reported the strong antileishmanial activity and selectivity of the synthetic chalcone CH8¹⁰ (3-nitro-2-hydroxy 4,6-dimethoxychalcone, Figure 1), a lipophilic and highly stable compound that fulfills most of the criteria specified by DNDi for a promising anti-leishmanial drug. *In vivo*, CH8 given to mice by the oral route is effective and safe against both cutaneous and visceral leishmaniasis¹¹. Importantly, local subcutaneous injections with CH8 either in the free form¹⁰ or loaded in slow-release systems^{12, 13} effectively treated mice with CL. However, effective topical formulations remain a challenge for this and other anti-leishmanial compounds.

Drug nanocarriers have been extensively applied to improve drug delivery and bioavailability of molecules with poor solubility in water, such as chalcones. These systems may protect the drug against degradation while promoting its intestinal uptake or skin permeation through the epidermis layer or hair follicle¹⁴. Lipid-core nanocapsules (LNCs) made with a biodegradable poly(ϵ -caprolactone) shell have emerged as appropriate nanocarriers to increase the bioavailability of lipophilic agents, since these can be entrapped into the carrier lipid core¹⁵. Previously, we showed that encapsulation in LNC promotes the intestinal absorption of the flavonoid quercetin, increasing by 40

fold the efficacy of this lipophilic molecule via the oral route, in a murine model of CL.¹⁶ In topical drug delivery assays, LNCs successfully promoted skin permeation of other lipophilic molecules such vitamin K1, resveratrol and curcumin^{17, 18}, without appreciable skin irritation.

Given the efficacy of CH8 against CL and the safety and potential of LNCs as nanocarriers of drugs to the skin, we describe here the development and testing – both *in vitro* and *in vivo* - of an LNC-CH8 formulation for the topical treatment of CL.

METHODS

Animals and ethics statement

BALB/c mice (female, eight-week-old, 23 g) used in the experiments were maintained under controlled conditions at the animal facilities of the Federal University of Rio de Janeiro (UFRJ, RJ, Brazil).

The animal protocols used in this study were approved by the local Animal Care and Use Committee (protocol number CAUAP118). All animal experiments were conducted in compliance with the principles stated in the *Guide for the Care and Use of Laboratory Animals, 8th Edition*.¹⁹

Production of lipid-core nanocapsules

The compound 3-nitro-2-hydroxy-4,6-dimethoxy chalcone (CH8, Figure 1) was synthesized as described previously¹⁰.

Lipid-core nanocapsules (LNC) were prepared by a self-assembly methodology²⁰. Briefly, 200 mg poly(ϵ -caprolactone) (80,000 g mol⁻¹ PCL;

Sigma-Aldrich Co., USA), 80 mg sorbitan monostearate (Sigma-Aldrich Co.), 310 mg capric/caprylic triglyceride (Alpha Quimica, Brazil) were dispersed in acetone (46 mL), and then injected into an aqueous phase composed of polysorbate 80 (185 mg; Gerbras, Brazil), ethanol (8 mL) and water (106 mL). The mixture was stirred for 10 minutes at room temperature, the solvent was evaporated under reduced pressure (rotatory evaporator, Büchi, Switzerland) at 40 °C and the aqueous suspension was concentrated for a final volume of 10 mL. Streptomycin (100 µg/mL) and penicillin (100 UI/mL) (Sigma-Aldrich Co.) were added. LNC-CH8 was prepared by adding 5 mg CH8 in the organic phase and Rho-LNC-CH8 was prepared as described above, with the exception that part of the unlabeled PCL was replaced with 10 mg of Rhodamine B-labeled PCL (Rho-PCL)²¹.

Characterization of lipid-core nanocapsules

Formulations were analyzed by laser diffraction, in a Mastersizer® 2000 particle size analyzer (Malvern, UK)¹⁵, to determine their volume-weighted mean diameter (D[4,3]). Size polydispersity was calculated considering the diameters at 10%, 50%, and 90% of the cumulative size distribution (SPAN). The Z-average diameter (method of cumulants) and polydispersity index were determined by photon correlation spectroscopy (PCS) using a Zetasizer Nano ZS particle analyzer (Malvern, UK) after dilution. The zeta potential of nanoparticles was calculated by examining the electrophoretic mobility of formulations dispersed in a 10mM NaCl aqueous solution (pH = 7.0) using a Zetasizer Nano ZS (Malvern, UK).

CH8 encapsulation efficiency

A sample of the formulation was dissolved in acetonitrile and filtered (0.45 μm , Merck Millipore, USA). The CH8 content was quantified by HPLC-UV [Perkin-Elmer® S200 pre-column, column LiChrospher 100 RP₁₈ (5 μm) (Merck®)], using a mobile phase of acetonitrile:aqueous 0.01% phosphoric acid (Tedia) (80:20 v/v), with a flow rate of 1 mL/min and a detection wavelength of 337 nm, as described previously.²²

Parasites

Leishmania amazonensis (MHOM/BR/75/Josefa strain) promastigotes were maintained in culture at 26°C in M199 medium (CultiLab, Brazil) supplemented with 10% heat-inactivated fetal calf serum (HIFCS) (CultiLab, Brazil), 0.2% hemin (Sigma-Aldrich Co., USA), 100 $\mu\text{g}/\text{mL}$ streptomycin and 100 UI/mL penicillin (Sigma-Aldrich Co., USA), herein referred to as 'complete medium'. Alternatively, GFP-expressing parasites were cultured as described previously²³ Parasites were always used at the stationary phase of growth.

Anti-amastigote activity

Bone marrow derived macrophages (BMDM) from BALB/c mice were prepared as described²³ and plated (2×10^5 / well) onto circular glass coverslips in 24-well culture plates, and incubated for 24 h at 37°C (and 5% CO₂). Then macrophage monolayers were kept uninfected or were infected with *L. amazonensis* promastigotes (5×10^6 cells/ mL) for 4 h at 34°C / 5% CO₂. After a further 24h of infection at 37°C, macrophages were incubated with 0.1, 1, 10 and 100 $\mu\text{g}/\text{mL}$ of CH8, LNC-CH8 or equivalent blank LNC concentrations, for

48h at 37°C. Then, cells were washed with PBS and stained with panoptic (Instant Prov kit, Newprov, Brazil) according to the manufacturer's instructions. The number of amastigotes in 200 cells was counted by light microscopy examination, in a Nikon Eclipse Ti microscope, at 1000 X magnification. The compound concentrations required to reduce amastigote numbers by 50% (IC₅₀ values) were calculated by linear regression analysis.

Cytotoxicity

The supernatants of the anti-amastigote assay were collected at the end of the treatment described above and the release of the cytosolic enzyme lactate dehydrogenase (LDH) by the macrophages was used as a cytotoxicity indicator. For total and spontaneous LDH releases, supernatants were collected from cells cultured with 1% triton and medium alone, respectively. LDH release was measured by a colorimetric assay using the detection kit DHL (Doles, Brazil), according to the manufacturer's instructions. Readings were taken at 492 nm in PowerWave XS spectrophotometer (BIO-TEK, USA), and cytotoxicity to BMDM calculated as the % of specific LDH release = $(\text{test} - \text{spontaneous} / \text{total} - \text{spontaneous}) \times 100$. The compound concentrations required to produce 50% of cytotoxicity (CC₅₀ values) were calculated by linear regression analysis. The selectivity index (SI) for each compound was calculated as the ratio between 50% cytotoxicity (CC₅₀) and 50% anti-amastigote activity (IC₅₀).

Rho-LNC-CH8 uptake assays

For Rho-LNC-CH8 uptake by promastigotes, parasites (10⁶ cells/mL), were plated in 24 well-plates in M199 medium containing 10% HIFCS, in the presence or absence of Rho-LNC-CH8 (10 µg/mL CH8). After 15 and 60 min of

incubation, cells were washed with phosphate-buffered saline (PBS) and transferred to glass slides for visualization in an Eclipse-Ti fluorescence microscope (Nikon) equipped with a CFI Plan Apo Lambda 100X Oil objective. Standard filters for FITC (B-2A) or rhodamine (G2 E/C) and DIC were used. Images were analyzed using the Adobe Photoshop CC software (version 14.2.1). Alternatively, fluorescent cells were analyzed by flow cytometry in a FACScan equipment (BD Biosciences), and the results were expressed as the percentage of cells (as gated by FSC vs. SSC) with fluorescence above 6×10^0 arbitrary units (10^5 events/sample).

For LNC-CH8 uptake by macrophages and intracellular amastigotes, BMDM (2×10^5 cells/well) were infected as described for the anti-amastigote activity assay. After 24 h, infected and non-infected macrophages were treated with Rho-LNC-CH8 (10 μ g/mL CH8) for up to 240 min and cells were imaged by fluorescence microscopy as described above. Alternatively, BMDM were infected and treated in 6-well plates (at 10^6 cells/well), and then subjected to flow cytometry analysis as described above.

Macrophage reactive oxygen species (ROS) production

BMDM (1×10^5 cells/well) were plated into black, clear-bottom 96-well plates, and then infected with *L. amazonensis* promastigotes and treated with CH8, LNC-CH8, or LNC as described above for the anti-amastigote assay. Then, cells were incubated with the oxidation-sensitive fluorescent dye 2',7'-dichlorodihydrofluorescein diacetate (H₂DCFDA, at 10 μ M; Invitrogen, USA) for 20 min at 37°C, and fluorescence was read at 485/528 nm (excitation/emission) in an FLX800 plate fluorimeter (BIO-TEK, USA). Uninfected and infected

macrophages incubated with 250 µg/mL Zymosan (Sigma–Aldrich Co.) were used as positive controls for ROS production.

Macrophage nitric oxide (NO) production

BMDM (1×10^5 cells/ well) were plated in 96-well plates, and then infected with *L. amazonensis* promastigotes and treated with CH8, LNC-CH8, or LNC as described above for the anti-amastigote assay. NO production in culture supernatants was measured by the Griess method, as described previously²⁴. Readings were taken at 570 nm in a PowerWave XS spectrophotometer (BIO-TEK, USA). The nitrite concentration was calculated using a standard curve of sodium nitrite (NaNO_2), from 0 to 50 µM. Uninfected and infected macrophages incubated with 1 µg/mL lipopolysaccharide (LPS; Sigma–Aldrich Co.) were used as positive controls for NO production.

Macrophage intracellular proteolysis

BMDM (1×10^5 cells/well) were plated in black clear bottom 96-well plates, maintained and infected as described above for the anti-amastigote activity assay. At 24 h after infection, cells were incubated with CH8 (1 µg/mL), LNC-CH8 or LNC for 20 min at 37°C. Then, cells were allowed to internalize the fluorogenic protein substrate DQ green-BSA (Thermo Fisher, UK) and human IgG (Sigma-Aldrich) - functionalized carboxylated silica beads (Kisker Biotech GmbH & Co., Germany) (5:1) for 5 minutes, as described previously²⁵. Alexa Fluor 594 succinimyl ester (Molecular Probes, USA) was used as a calibration fluorophore. After washing in PBS (to remove non-phagocytosed beads), the increase in fluorescence, indicative of proteolysis, was measured at different time points during a 2h period, at 37°C, using an FLX800 plate fluorimeter (BIO-

TEK, WI, USA) set at 594/620nm (for calibration fluorescence - CF) or 485/528nm (for substrate fluorescence - SF). The 'blank' (B) represented the cell fluorescence before treatment. The relative fluorescence (RF) was calculated as $RF = (SF - B) / (CF - B)$.

Topical drug absorption assay

Mice were anesthetized with 2% isoflurane inhalation, and Rho-LNC-CH8 (20 μ L, containing 11 μ g of CH8) were applied to a shaved dorsal area of approximately 1.5 cm^2 . At 0, 5, 15, 30, 60, 90, 120, 180, 240 and 300 min after application, fluorescence images of the whole animals were taken using an IVIS® Lumina Imaging System (Xenogen Corp, USA) with filters for rhodamine detection. A region of interest (ROI) was drawn 2 mm around the outer fluorescent margin, and the specific fluorescence was expressed as the difference between values for Rho-LNC-CH8 and LNC-CH8 surface radiance (photons/s/ cm^2) for each time point. As a fluorescence quenching/decay control, the same amount of Rho-LNC-CH8 was applied per cm^2 onto an impermeable plastic blade and the fluorescence was measured as performed for animal skin.

Mouse infection and topical treatment

Mice were infected in the ear with 2×10^6 promastigotes of *L. amazonensis*. On days 7 to 28 of infection, the ears were treated topically (5 times a week, using a disposable spatula) with one of the following formulations: (1) 10 μ g of CH8 in PBS/2% DMSO (CH8 group); (2) 10 μ g of CH8 in LNC-CH8 (LNC-CH8 group); blank LNC (LNC group); or 20 μ L of 2% DMSO alone (untreated group). At the end of treatment (day 30 post-infection), the animals were euthanized by terminal 5% isoflurane inhalation, the infected ears were excised aseptically at the base, and homogenized individually in 1

mL of complete M199 medium. The parasite burden was evaluated by a limiting dilution assay (LDA), as described previously²⁶.

Statistical analysis

Data were analyzed by one-way or two-way ANOVA followed by Bonferroni multiple comparison test. Data were expressed as arithmetic mean \pm standard deviation (SD) values and samples were considered significantly different when $p \leq 0.05$ in a series of at least three independent experiments.

RESULTS

3.1 LNC efficiently encapsulates CH8 into homogeneous nanoparticle suspensions.

All formulations were macroscopically homogeneous, and had unimodal narrow size dispersions, with mean diameters below 238 nm (laser diffraction) and 184 nm (PCS) (Table 1). All nanocapsules showed a slight negative zeta potential (ζ) close to zero (-6.1 to -7.5 mV). The drug content was close to 500 $\mu\text{g}/\text{mL}$ with an encapsulation efficiency close to 100% (99.9%) (Table 1). Overall, nanocapsule sizes and surface potential were slightly affected by the drug loading or rhodamine labeling. The high efficiency of drug encapsulation corroborate the previous results observed for other drugs, such as doxorubicin, tacrolimus, quercetin, resveratrol and curcumin^{16, 18, 27, 28}.

3.2 Rho-LNC-CH8 is internalized by *L. amazonensis* promastigotes

To evaluate whether LNC could be internalized by parasite promastigote forms, these cells were incubated with Rho-LNC-CH8 (at 10 $\mu\text{g}/\text{mL}$ of CH8) and the nanoparticle uptake was monitored over time, by flow cytometry and

fluorescence microscopy. Promastigotes internalized Rho-LNC-CH8 rapidly, and, within 15 minutes, a red fluorescence signal was observed in a specific sub-cellular compartment. Afterwards, at 60 min of internalization, the fluorescence signal appeared widespread in the cytoplasm. Quantification of the kinetics of Rho-LNC-CH8 internalization showed that the proportion of cells positive for Rho-LNC-CH8 increased with the incubation time (Figure 2), with ~80% of promastigotes having internalized Rho-LNC-CH8 by 60 min of incubation. These data showed the nanosystem is efficiently internalized by the *Leishmania* form that initiates infection in mammalian hosts.

3.3 CH8 encapsulation in LNCs did not decrease its anti-leishmanial activity.

In a previous study, we demonstrated that CH8 has potent and selective activity against *Leishmania* promastigotes¹⁰. Here, we tested both CH8 and LNC-CH8 against the intracellular forms of the parasite, the amastigotes. Infected macrophages were incubated with CH8, LNC-CH8 or LNC and the number of amastigotes inside host cells was assessed by direct counting under a light microscope, after 48 h of treatment (Table 2). Entrapping of CH8 in LNCs did not alter the anti-amastigote activity of the compound *in vitro*, as both CH8 and LNC-CH8 showed similar IC₅₀ values (2.2 ± 0.1 and 2.9 ± 0.1 $\mu\text{g/mL}$, respectively). LNC (blank formulation) was not able to kill intracellular amastigotes (IC₅₀ of 38.8 ± 0.3 $\mu\text{g/mL}$). However, LNC was not selective to the parasite, with encapsulation leading to a reduced selectivity index (SI) of CH8 towards the parasite, in comparison to the free drug.

3.4 LNC-CH8 delivers CH8 to *Leishmania* amastigotes inside host cells.

Since encapsulation on LNCs did not alter the activity of CH8 *in vitro*, we evaluated the internalization of Rho-LNC-CH8 by infected and uninfected macrophages, to verify if the nanocapsules could deliver CH8 effectively to parasites inside macrophages, where the parasite survives and multiplies. For this purpose, bone marrow-derived macrophages (BMDM) infected with *L. amazonensis* promastigotes expressing GFP²³ were incubated with Rho-LNC-CH8 (10 µg/mL CH8), and the internalization was evaluated at different time points (up to 240 min), by fluorescence microscopy and flow cytometry (Figure 3).

Similarly to that observed in promastigotes, Rho-LNC-CH8 was internalized quickly by infected and uninfected BMDMs, and we observed red fluorescence in the macrophage cytoplasm (but not in the nucleus) within 15 min of internalization (Figure 3A). In infected macrophages, the Rho-LNC-CH8 red fluorescence appeared faster (at 7.5 min) than in uninfected cells (Figure 3C), and internalization increased over time (Figure 3A and C). Importantly, the red fluorescence signal of Rho-LNC-CH8 was concentrated in the parasitophorous vacuole (PV) of infected cells (Figure 3A). Moreover, co-localization of Rho-LNC-CH8 with GFP-labeled amastigotes in the PV could be observed after 60 and 120 min of incubation. After 240 min of incubation, the Rho-LNC-CH8 fluorescence was found predominantly co-localized with GFP-labeled parasites in the PV, suggesting that Rho-LNC-CH8 had likely killed the amastigotes by that time point. In uninfected cells, the Rho-LNC-CH8 fluorescence signal had decreased by 240 min of internalization, with only low fluorescence being detected by fluorescence microscopy and a reduced

percentage of fluorescent cells detected by flow cytometry (Figure 3A, B and C). This decrease in fluorescence at later time points was not observed in infected cells (Figure 3C).

Together, these results confirm that LNC-CH8 can be internalized by infected macrophages and delivered to the PV, the intracellular compartment where the parasite is located.

3.5 LNC-CH8 triggers macrophage microbicidal mechanisms.

Phagocytes such as macrophages are capable of responding to infection by triggering microbicidal mechanisms, which include the production of reactive oxygen species (ROS) and nitric oxide (NO), and the activation of proteolytic enzymes and acidification of the PV. To study the influence of entrapment into LNCs on the activation of macrophage microbicidal mechanisms by CH8, we evaluated the stimulation of these mechanisms – which constitutes macrophage activation - in infected cells treated with LNC-CH8.

For ROS production, infected and uninfected macrophages were incubated with increasing concentrations of CH8, LNC-CH8 or LNC, and ROS production was assayed using the fluorescent dye H₂DCFDA (Figure 4, A and B). As expected, *Leishmania* infection decreased the basal ROS levels in untreated cells. CH8 was unable to activate ROS production in macrophages, even at high concentrations. In contrast, both LNC and LNC-CH8 induced ROS production at medium to high concentrations (Figure 4A and B).

In contrast, NO production (as estimated by NaNO₂ quantification in culture supernatants) increased only after treatment with high concentrations of

LNC-CH8; however, LNC and CH8 could not stimulate NO production, even at high concentrations (Figure 4C and D).

We assessed the proteolytic activity of macrophages by measuring the degradation of functionalized beads over time, in infected and uninfected cells treated with 1 $\mu\text{g}/\text{mL}$ CH8, or equivalent amounts of LNC-CH8 and LNC. The proteolytic activity of uninfected cells was similar for all treatments (Figure 4E), while proteolysis levels increased over time in infected cells treated with LNC-CH8 or LNC, but not CH8 alone (Figure 4F). Although the proteolytic response was delayed in infected compared with uninfected cells, treatment of the former with LNC or LNC-CH8 eventually increased proteolysis levels above those observed uninfected cells (Figure 4F). CH8 encapsulation in LNC did not change cell acidification significantly (as assessed using the LysoTracker Green dye), irrespective of the presence of infection (data not shown).

The combined analysis of different antimicrobial mechanisms showed that LNC-CH8 can modulate host macrophage activation positively, which contributes to the activity of the LNC-CH8 nanosystem against the intracellular parasite.

3.6 Rho-LNC-CH8 is absorbed efficiently by mouse skin.

After the promising *in vitro* results described above, we evaluated if the LNC-CH8 formulation could permeate mouse skin and deliver the drug to macrophages *in vivo*. The kinetics of Rho-LNC-CH8 skin absorption was evaluated for up to 300 min after application onto the lightly shaved mouse hump. Although Rho-LNC-CH8 fluorescence was stable for up to 120 min after application, we observed a clear decrease in fluorescence thereafter until 240 min, when almost complete Rho-LNC-CH8 absorption was achieved (Figure 5A

and B). Spontaneous fluorescence quenching by drying was discarded by surface radiance quantification demonstrating that the fluorescence intensity of Rho-LNC-CH8 applied onto an impermeable plastic surface remained stable throughout the experiment (Figure 5B).

3.7 LNC-CH8 increased the efficacy of CH8 in cutaneous leishmaniasis treatment.

Given that Rho-LNC-CH8 is absorbed by mouse skin, we tested the activity of LNC-CH8 *in vivo* as a topical treatment for experimental CL, in BALB/c mice infected in the ear with GFP-labeled *L. amazonensis*.

After 3 weeks of daily topical treatment (5 days / week) with LNC, LNC-CH8 or CH8 (10 μg CH8 per cm^2 / dose), the parasite burden was evaluated in infected mouse ears by a limiting dilution assay. Mice treated with LNC-CH8 had significantly fewer parasites in infected ears than animals either treated with CH8 alone, or kept untreated (Figure 6). Thus, encapsulation of CH8 in LNC-CH8 increased its efficacy by the topical route in a murine model of CL.

DISCUSSION

In recent years, various topical formulations containing paromomycin, pentavalent antimonials, β -lapachone and amphotericin B have been developed for CL treatment^{4-6, 29}, but none were sufficiently effective or safe for clinical approval. Here, we evaluated the efficacy of a new topical nanoformulation against CL that combined the potency of a promising antileishmanial compound – the nitro-chalcone CH8¹¹ - with the skin permeation ability of the versatile LNC³⁰.

The results demonstrated that PCL and the oily component used here were adequate to provide a high rate of CH8 incorporation into LNC-CH8 (99.9%). Most of the drug was probably retained into the lipid compartment, which is protected by the PCL shell, thus increasing its bioavailability, as demonstrated for oral quercetin and resveratrol^{16,31}. Despite their weak negative charge (-7.5 mV), LNC-CH8 stability was achieved by steric hindrance promoted by the nonionic polysorbate 80 surfactant³².

The resulting LNC-CH8 were readily taken up by both macrophages and promastigote forms of *Leishmania*. Light microscopy analysis of Rho-LNC-CH8 internalization suggests that the nanocapsules enter promastigotes through the flagellar pocket and are transported to organelles of the endocytic pathway (likely corresponding to lysosome multivesicular tubules³³) within 15 min, and then reach the cytoplasm. In macrophages, Rho-LNC-CH8 may reach the cytoplasm by fluid phase endocytosis, as described for LNC-doxorubicin²⁸. Therefore, our data suggest that LNC-CH8 are only degraded after reaching the PV³⁴, where *Leishmania* amastigotes reside.

Encapsulation in LNC did not impair the anti-amastigote activity of CH8, as observed previously with poly(lactide-co-glycolide) (PLGA) microspheres that delayed intracellular parasite killing²². LNC may produce faster drug delivery to intracellular parasites due to their smaller size and/or higher sensitivity to lysosomal enzymes, compared with PLGA microspheres. The higher cytotoxicity of LNC-CH8 and LNC to macrophages, compared with CH8, is likely caused by the presence of polysorbate 80 surfactant in both LNC formulations. Despite the increased cytotoxicity, LNC-CH8 selectivity remained high - at 61 fold - which is comfortably above the 10-fold selectivity index advocated by

DNDi as a 'cut-off' for promising new antiparasitic drugs³⁵. Additionally, the polysorbate 80 cytotoxicity observed here may not be relevant *in vivo*, especially for topical administration, since intradermal polysorbate 80-containing LNC systems are safe to rats³⁶.

The innate immune response to parasitic infection leads to the production of ROS and NO which together generate peroxynitrite (ONOO⁻), a strong oxidant considered essential for parasite killing by phagocytic cells³⁷. These compounds can also increase the expression of genes associated with pro-inflammatory responses, such as NF- κ B transcription factors³⁸. While nanocarriers are known to activate these mechanisms,^{39,40} the LNC nanoparticles used in this studies only led to an increase in ROS species. Moreover, in line with the antioxidant and anti-inflammatory effects of chalcones^{40,41}, CH8 alone could not activate macrophage antimicrobial mechanisms. In contrast, encapsulation into LNC led to ROS and NO production in both uninfected and infected cells. The former possibly accounting for the slightly increased macrophage cytotoxicity observed with LNC-CH8. Interestingly, intracellular *Leishmania* parasites downregulate these macrophage antimicrobial mechanisms² and the use of LNC nanocarriers may help to overcome this effect.

In addition to ROS and NO production, intracellular proteolysis is essential for microbial killing after endocytosis⁴². Interestingly, for safe therapeutic purposes, LNC-CH8 induced proteolytic activity in infected but not uninfected macrophages, suggesting that the presence of the parasites was necessary for the activation of cellular digestion mechanisms.

The lower efficacy of previous topical formulations for CL is possibly due to lack of proper drug permeation through the *stratum corneum* barrier of the skin, necessary to reach the infected cells in the lower dermis layer³⁰. Despite its low molecular weight (329Da) and high lipophilicity (LogP = 3.63), CH8 alone cannot permeate the skin barrier⁴³. However, the imaging studies shown here demonstrate that Rho-LNC-CH8 applied onto the mouse skin surface is no longer detectable 3 hours post-application, suggesting that the drug is absorbed efficiently by the skin. The time required for skin signal decrease suggests that drug absorption is not due to occlusion, as reported previously for solid lipid nanoparticles⁴⁴. Rho-LNC-CH8 and LNC-CH8 permeation through skin appendages (hair follicles or sweat glands)³⁰ appears more plausible. This is agreement with the studies of Raber et al⁴⁵ demonstrating that negatively charged nanoparticles are able to interact more efficiently with the skin surface, facilitating follicular uptake.

Although the mechanism by which LNC-CH8 promotes drug permeation through the skin remains unclear, the *in vivo* data in murine CL clearly show that topical application of LNC-CH8 onto mouse lesions effectively reduced the parasite load by ~80% in comparison with free CH8. Interestingly, LNC alone reduced the number of parasites in skin lesions by ~30%. Although blank LNC does not have intrinsic antiparasitic activity, as seen by the lack of *in vitro* activity, our data suggest that this blank nanodevice is an excellent adjuvant for topical CL treatment. LNC-CH8 was more effective than other nanosystems attempted for CL topical treatment^{29, 46}. Following the skin permeating potential demonstrated here, future LNC-CH8 manipulations may further improve its antiparasitic activity. One possibility is LNC co-entrapment with

superparamagnetic iron oxide nanoparticles (SPIONs) to allow heating and killing of the infected cells after local application of a magnetic field, as envisaged for cancer thermotherapy⁴⁷.

Overall, the results reported here show that LNC-CH8 is an appropriate nanosystem not only for intracellular drug delivery, but also for skin permeation and intracellular parasite killing. This study has clinical relevance for localized and safe treatment of CL, a skin pathology for which there is currently no adequate topical treatments.

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FIGURE LEGENDS

Figure 1. Chemical structure of the chalcone CH8.

Figure 2. Kinetics of Rho-LNC-CH8 internalization by promastigotes.

Leishmania amazonensis promastigotes were incubated with rhodamine-labeled LNC-CH8 (Rho-LNC-CH8, at 10 µg/mL CH8) and analyzed at different time points, to detect compound internalization. (A) Light microscopy analysis of live promastigotes, showing internalization of red-fluorescent Rho-LNC-CH8 from 0 to 60 min of incubation. DIC, differential interference contrast. Scale bar,

5 μm . (B, C) Flow cytometry analysis of the Rho-LNC-CH8 fluorescence intensity in promastigotes (10^5 events/sample) at different time points, displayed as histograms (B) and as the percentage of cells expressing $> 6 \times 10^0$ arbitrary units of fluorescence (C; corresponding to cells in gate M1, in B). AU, arbitrary units. (n=3).

Figure 3. Rho-LNC-CH8 is efficiently internalized by macrophages. Bone marrow derived macrophages (BMDM) were infected with *L. amazonensis* expressing GFP and incubated with Rho-LNC-CH8 (10 $\mu\text{g}/\text{mL}$ CH8). (A) Light microscopy analysis of uninfected and infected BMDM at different time points of incubation with Rho-LNC-CH8, showing co-localization between amastigotes (in green) and Rho-LNC-CH8 (in red), after 120 min of incubation. DIC, differential interference contrast. Scale bar, 20 μm . (B, C) Flow cytometry analysis of uninfected and infected BMDM after 240 min of Rho-LNC-CH8 internalization. The histograms in B show treated cells in blue, and untreated in grey, and the percentage of cells expressing $>3 \times 10^0$ arbitrary fluorescence units (gate M1 in B) is shown in C (mean \pm SD values, n=3 independent experiments). *p<0.05. AU, arbitrary units.

Figure 4. LNC-CH8 (but not CH8 alone) activates infected macrophages. Bone marrow derived macrophages (BMDM) were kept uninfected (A, C, E) or were infected with *L. amazonensis* (B, D, F), treated with LNC, CH8 or LNC-CH8 and then processed for the analysis of different markers of macrophage activation. (A, B) Reactive oxygen species (ROS) production was measured fluorimetrically using H_2DCFDA , after 30 min of treatment. FU, fluorescence units (C, D) Nitric oxide (NO) levels were estimated by quantifying NaNO_2 in culture supernatants (by the Griess method), after 48h of treatment. (E, F).

Intracellular proteolysis was assessed by using DQ green-BSA and IgG-functionalized beads, after 30 min of treatment. Results are expressed as relative fluorescence (RF). Data represent mean \pm SD values (n=3 independent experiments). *p < 0.05, **p < 0.01, relative to free CH8.

Figure 5. Topical Rho-LNC-CH8 is absorbed through living mouse skin.

Rhodamine-labeled CH8 (Rho-LNC-CH8, with 10 μ g/mL CH8) was applied onto the pre-shaved dorsum of BALB/c mice and the skin fluorescence kinetics was followed by *in vivo* fluorescent imaging using the IVIS® Lumina system (A). A region of interest (ROI) was drawn 2 mm around the outer fluorescent margin, and data expressed as average surface radiance (photons/s/cm²) after deduction of background fluorescence of skin to which unlabeled LNC-CH8 was applied (red line). As a control for fluorescence quenching/decay, the same amount of Rho-LNC-CH8 or LNC-CH8 was applied onto an impermeable surface (blue line) (B). Means \pm SD values (n=5).

Figure 6. Efficacy of topical LNC-CH8 against cutaneous leishmaniasis.

BALB/c mice were infected in the left ear with 2×10^6 promastigotes of *L. amazonensis*. Seven days after infection, the infected ears were treated topically once a day (five times a week), for 3 weeks, with 20 μ L of LNC, LNC-CH8 containing 10 μ g of CH8 (22.5 μ g CH8 per cm²) or the same amount of free CH8 in PBS/2% DMSO. The untreated group was administered only the drug-free vehicle. After 30 days of infection, animals were euthanized, the ears were removed and the parasite burden was quantified by a limiting dilution assay. Data represent mean \pm SD values of one out of two independent experiments (n=5 animals/group). ** p<0.01.

Formulation	D[4,3] (nm)	SPAN	ζ (mV)	z-average diameter (nm)	PDI	CH8 content ($\mu\text{g/mL}$)	EE%
LNC	167 \pm 0.009	1.20 \pm 0.1	-6.10 \pm 0.09	181 \pm 09	0.12 \pm 0.01	-	-
LNC-CH8	238 \pm 0.010	0.81 \pm 0.1	-7.50 \pm 0.07	174 \pm 07	0.10 \pm 0.01	501 \pm 0.02	99.9
Rho-LNC-CH8	145 \pm 0.013	1.12 \pm 0.1	-6.47 \pm 0.08	184 \pm 11	0.08 \pm 0.02	550 \pm 0.03	99.9

Table 1. Physicochemical features of the different nanocapsules.

LNC, “blank” LNC; LNC-CH8, LNC entrapping CH8; Rho-LNC-CH8, LNC-CH8 produced using a rhodamine(Rho)-labelled precursor. ζ , zeta potential; EE, Encapsulation Efficiency. (n=3).

Table 2. Anti-Leishmania activity of LNC-CH8 *in vitro*.

TREATMENT	ANTI-AMASTIGOTE IC₅₀ (µg/mL)	CYTOTOXICITY CC₅₀ (µg/mL)	SI
CH8	2.15 ± 0.14	393.72 ± 0.09	183.1
LNC	38.83 ± 0.29	247.90 ± 0.06	6.3
LNC-CH8	2.90 ± 0.15	178.60 ± 0.12	61.6

IC₅₀, anti-amastigote activity estimated by light microscopy. CC₅₀, cytotoxicity estimated by lactate dehydrogenase release. SI, selectivity index (SI = CC₅₀/IC₅₀). (n=3).

GRAPHICAL ABSTRACT

The antileishmanial drug candidate CH8 was encapsulated in lipid-core nanocapsules (LNC-CH8), to improve drug bioavailability for topical treatment of cutaneous leishmaniasis. LNC-CH8 were able to control the infection by penetrating mouse skin and delivering the drug to infected dermal macrophages.

Journal Pre-proof

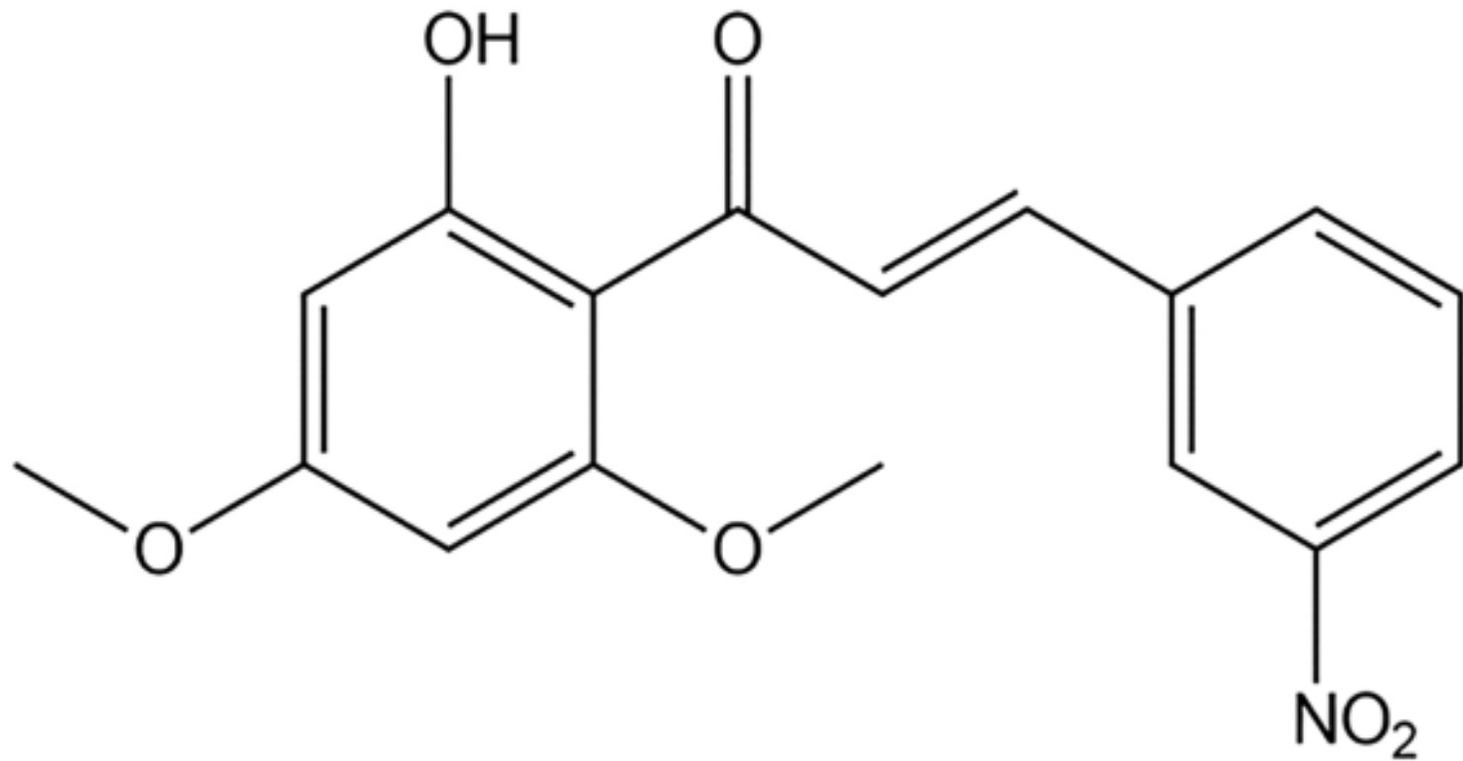


Figure 1

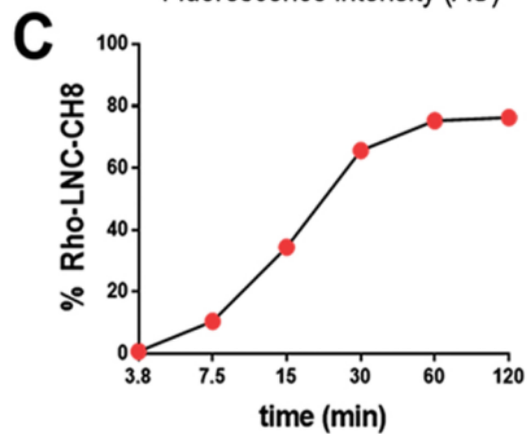
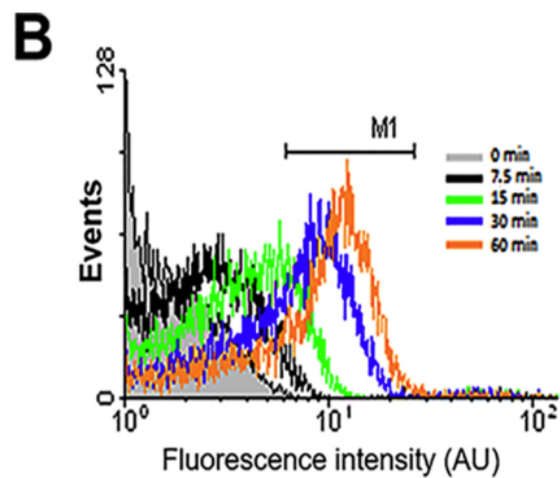
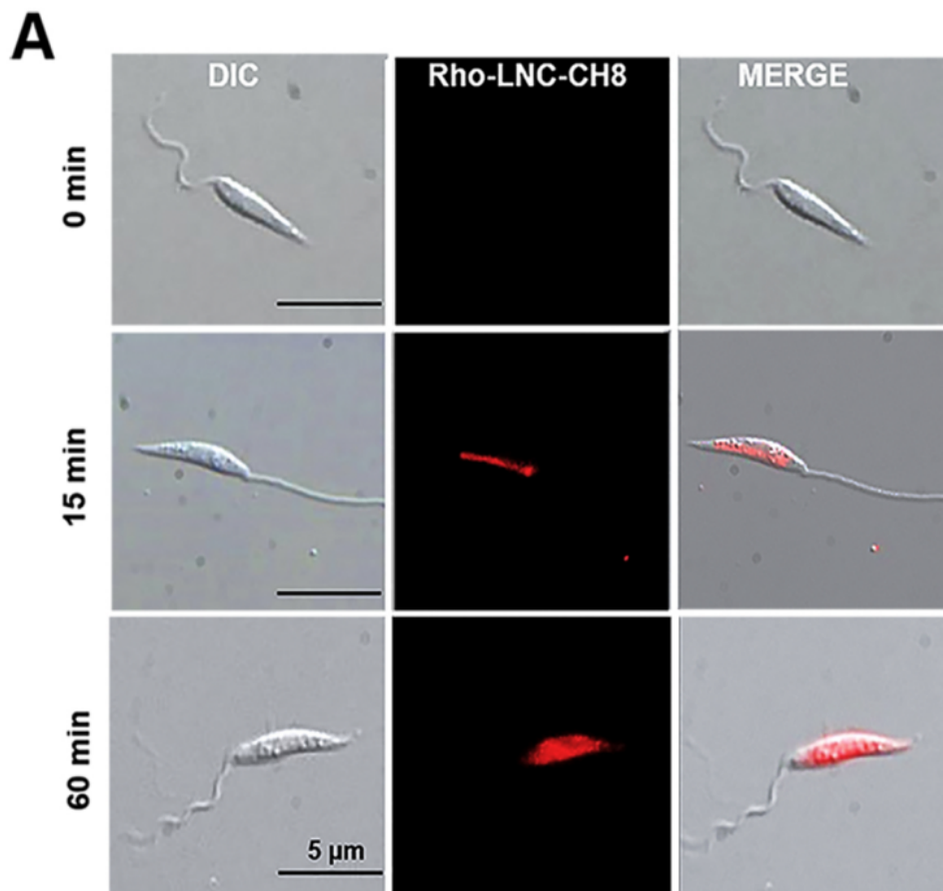


Figure 2

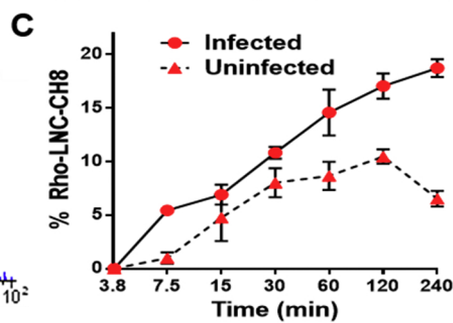
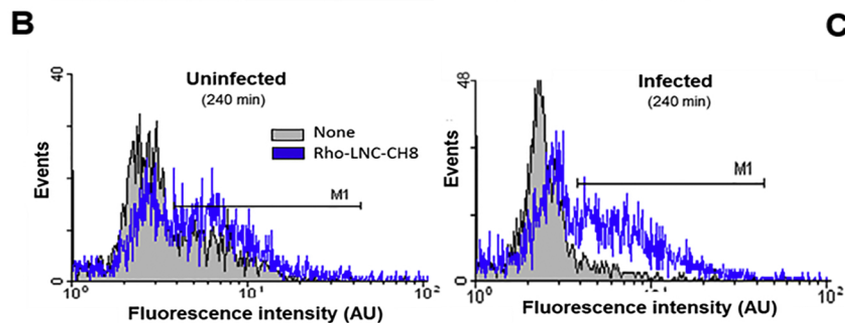
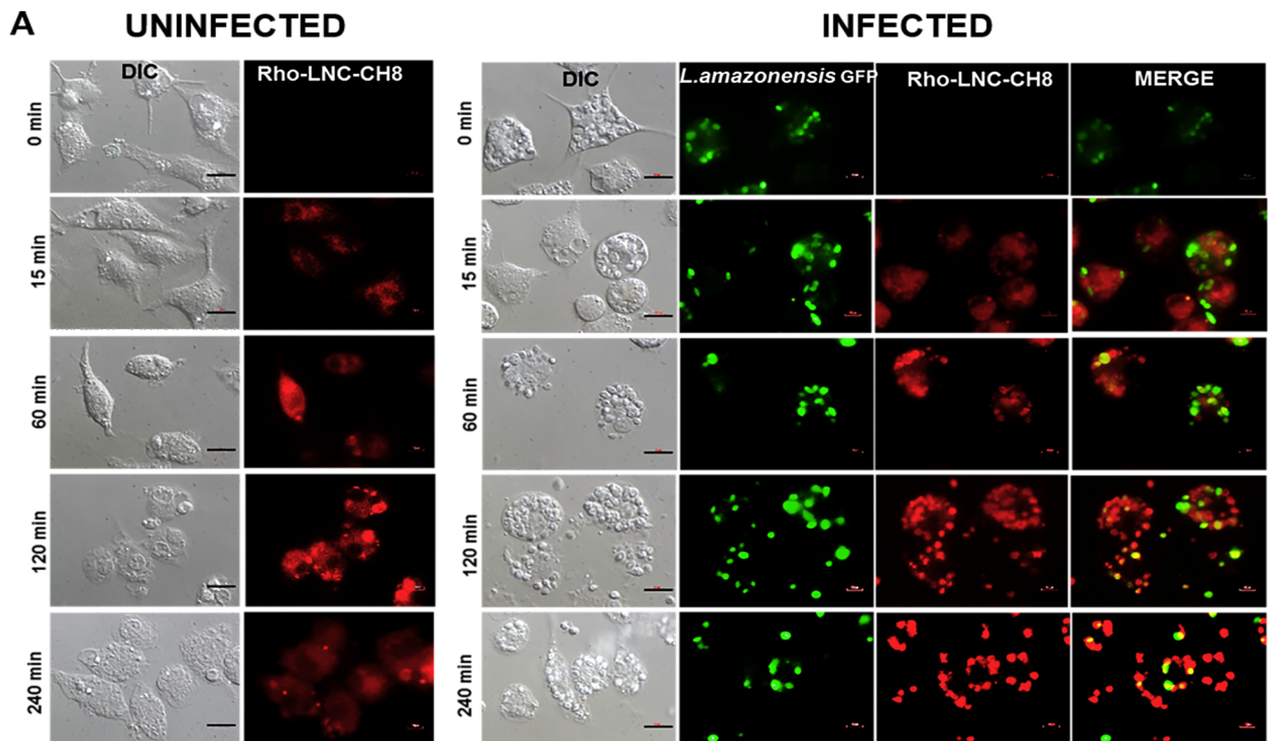


Figure 3

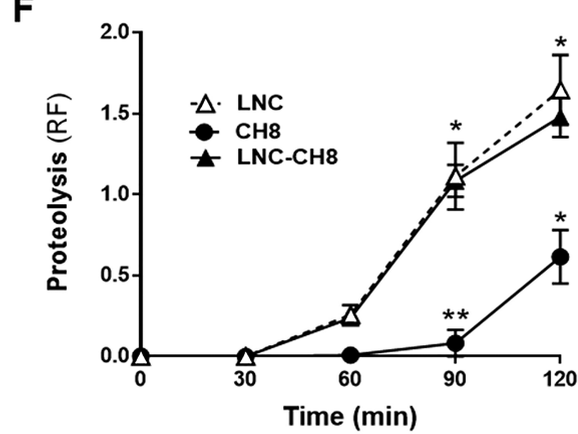
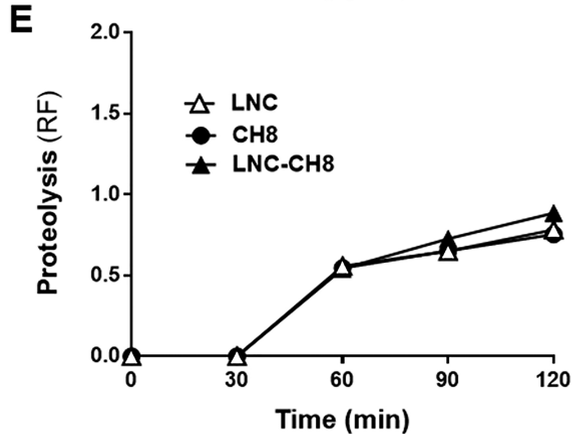
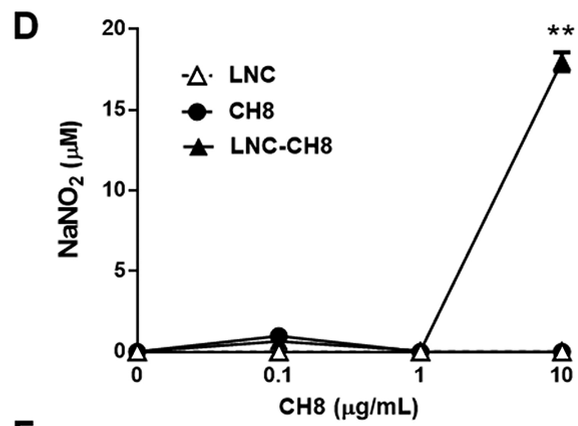
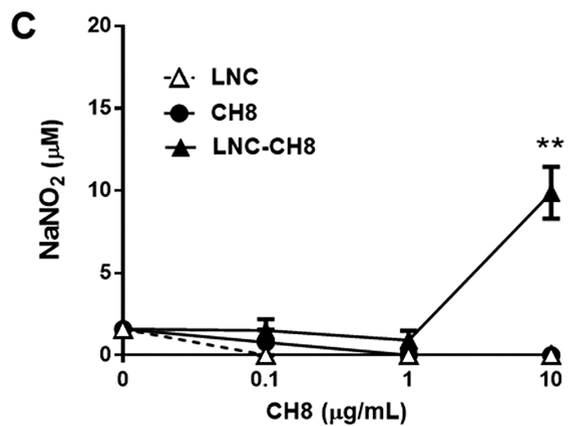
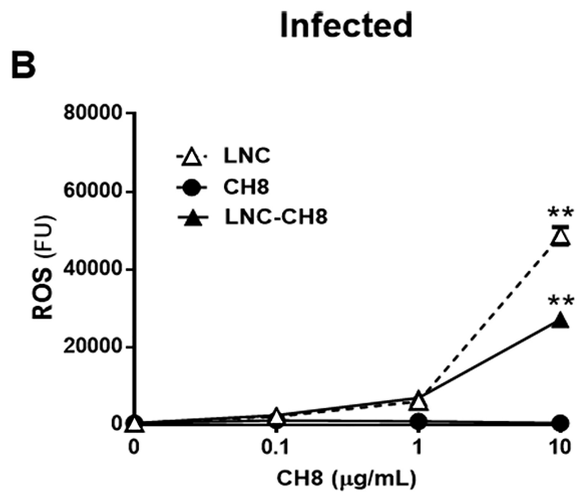
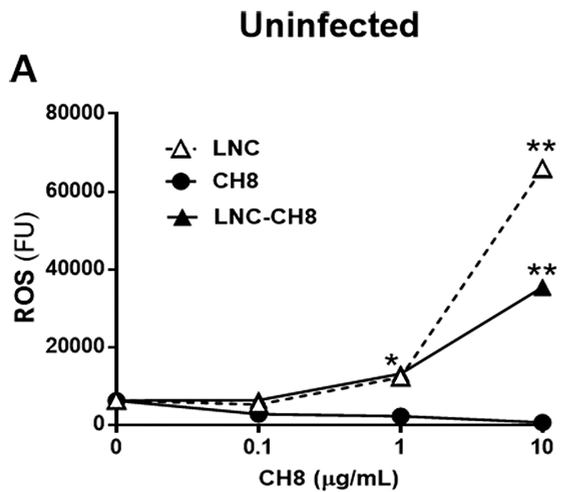


Figure 4

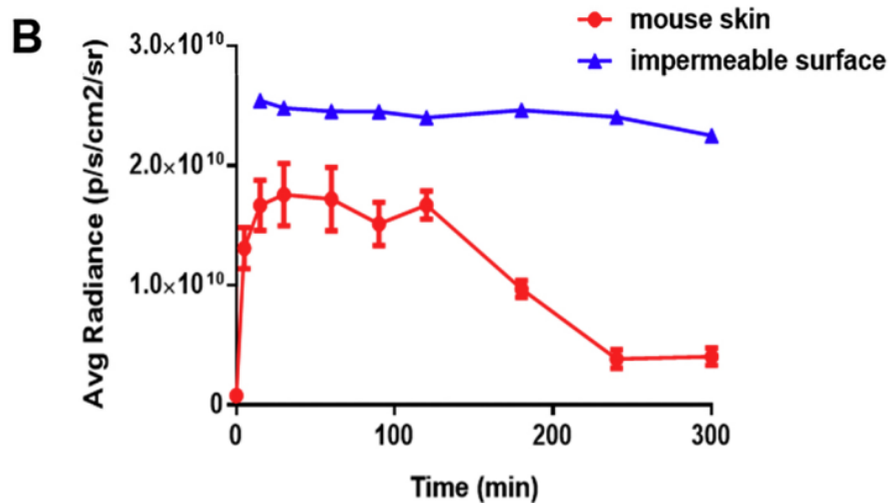
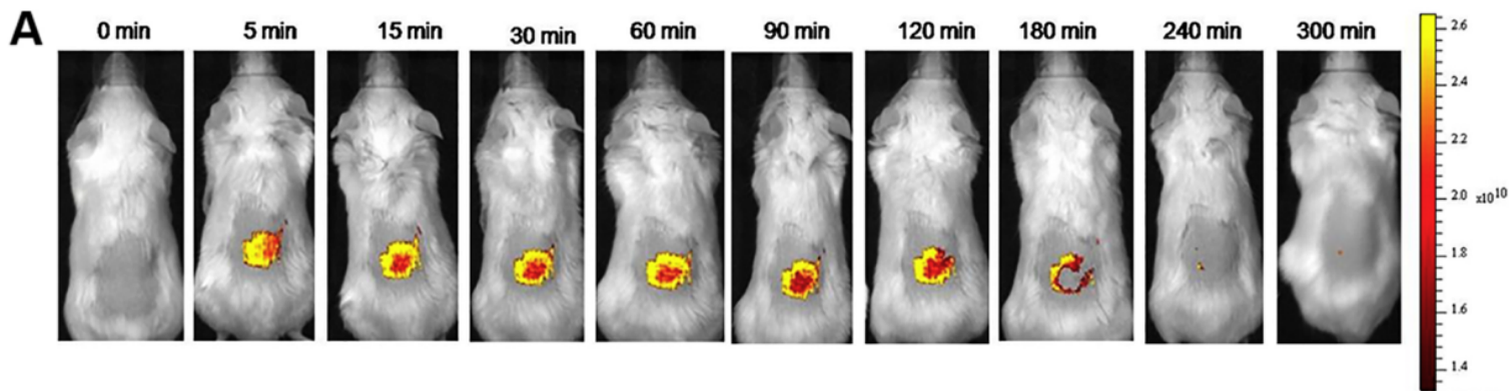


Figure 5

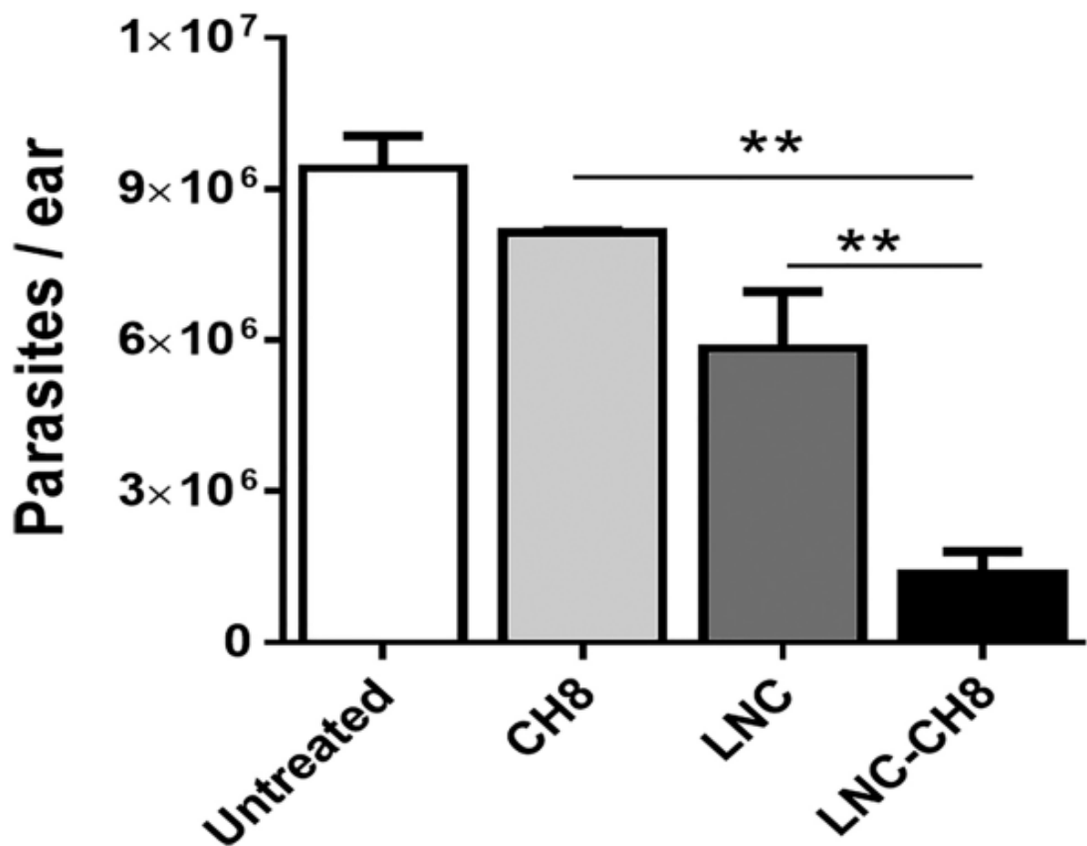


Figure 6