Insights into the application of chitosan as an anti-leishmanial compound.

Alaa Riezk

Thesis submitted in accordance with the requirements for the degree of
Doctor of Philosophy
of
the University of London

November 2019

Department of Infection Biology
Faculty of Infectious & Tropical Disease
London School of Hygiene & Tropical Medicine, University of London

Research supervisors: Dr Vanessa Yardley
Prof Simon L Croft
Dr Sudaxshina Murdan

Funded by the London School of Hygiene & Tropical Medicine (Prof S L Croft) and
CARA (the Council for At-Risk Academics
DECLARATION

I, Alaa Riezk, confirm that the work presented in this thesis is my own. Where information has been derived from other sources, I confirm that this has been indicated in the thesis.

Alaa Riezk

November-2019
Abstract

There is an urgent need for safe, efficacious, affordable and field-adapted drugs for the treatment of cutaneous leishmaniasis, a disease which affects around 1.5 million people worldwide every year. Chitosan, a biodegradable cationic polysaccharide, has previously been reported to have antimicrobial, anti-leishmanial and immunostimulatory activities. The work described here found that chitosan and its derivatives were approximately 7-20 times more active in vitro against Leishmania promastigotes and amastigotes at pH 6.5 than at pH 7.5, with high molecular weight chitosan being the most potent. Despite the in vitro activation of bone marrow macrophages by chitosan to produce nitric oxide and reactive oxygen species, this work showed that the anti-leishmanial activity of chitosan was not mediated by these metabolites. It was subsequently shown that rhodamine-labelled chitosan is taken up by pinocytosis and accumulates in the parasitophorous vacuole of Leishmania-infected macrophages. The application of chitosan in drug delivery systems was then studied by preparing two types of chitosan nanoparticles (positive (with tripolyphosphate sodium (TPP)) and negative (with dextran sulphate) surface charge with different sizes) and incorporation of amphotericin B within these nanoparticles. These amphotericin B-loaded nanoparticles demonstrated a good in vitro anti-leishmanial activity, similar to pure amphotericin B, and were also significantly less toxic than pure amphotericin B. The positive amphotericin B-loaded chitosan-TPP nanoparticles showed promising in vivo efficacy against cutaneous leishmaniasis caused by L. major in the BALB/c mouse model, via the intravenous route, and they were more active than AmBisome®. The impact of an in vitro media perfusion system on host cell phagocytosis and macropinocytosis was evaluated as well as the anti-leishmanial activity of chitosan solution and blank or amphotericin B-loaded chitosan-TPP nanoparticles. There was a significant difference between in vitro static and flow culture systems in the cell uptake and anti-leishmanial activity of the studied compounds.
Acknowledgements

I would like to express my sincere gratitude to my supervisors, Dr Vanessa Yardley and Prof Simon L Croft for the continuous support of my PhD study, for their enthusiasm, guidance, motivation, and immense knowledge. How can I ever thank you enough for all you have done? I’m forever grateful. You helped shape my research and professional life and showed me how to transform my mistakes into skills. I really appreciate everything you have taught me!

Many thanks to my supervisor Dr Sudax Murdan (Department of Pharmaceutics, School of Pharmacy, UCL) for providing invaluable guidance, helping me improve my skills, and for all the advice, ideas, support and patience in guiding me through this project. My supervisors, three of you, have been role models over the last years and was a great pleasure to meet you and have you in my life.

Special thanks should also be given to my colleague Katrien Van Bocxlaer. Kat, your passion for the research is contagious, your ability to repair things keeps astonishing me. I am sure that, your thoughtfulness is a gift I will always treasure, a huge thanks.

And thanks to the rest of the Croft group for all the good times in and out of the lab: Alec O’Keefee, Gert-Jan Wijnant and Markella Koniordou.

I wish to acknowledge Dr John Raynes for the technical support and guidance.

I acknowledge the facilities and the scientific and technical assistance of the LSHTM Wolfson Cell Biology Facility, with specific thanks to Dr Elizabeth McCarthy

Thanks to everyone at School of Pharmacy who helped me through my project, Dr S. Somavarapu, Dr Andrew Weston, Satinder Sembi.

Many thanks to CARA (Zeid, Laura, Riman, Lucia and all the team), LSHTM and Goodenough college, my PhD would not have been possible without their support.

I would also like to take this opportunity to thank Associate Prof Philippe M Loiseau and Dr Mohamed Alhnan—my viva examiners, for their very helpful comments and suggestions.

Thanks to all of my friends who have supported me throughout the process.

I would like to dedicate this PhD thesis to Simon L Croft, the symbol of love and giving. To my beloved wife Rana, never forget that I love you and this love will never fade for anything in the world, Rana I have faith in you. Carlos, you are one of the most amazing gifts I have ever received. Carlos, I thank Jesus for you every single day. You are a great blessing in my life, my always and forever.

Whatever I am today is because of my parents. It is their values and upbringing that have held me together despite all the struggles and adversities I have faced in such a hard time. Thanks to them, I have not crumbled under pressure, I have not stopped believing in myself and I am still fighting, Fadaa and Essam

Finally, my brothers and their families, despite the loneliness that comes with being so far from you, I hold on knowing that you truly love me, that our bond can’t be broken, Nazier, Subee, Elie, Hala, Hala, Maram, Wisam, Tala, Lana, Rita, Essam, Jad, Ameer and Marcos with much love.
3.2. Material and methods

3.2.1. Preparation of blank chitosan nanoparticles

3.2.2. Preparation of AmB loaded chitosan nanoparticles

3.2.3. Physicochemical properties of the nanoparticles (size, charge and morphology)

3.2.4. Stability of nanoparticles regarding size and zeta potential

3.2.5. Determination of drug encapsulation efficiency and AmB loading and release

3.2.6. In vitro release of AmB

3.2.6.1. Quantification of AmB by HPLC

3.3. Results
3.3.1. Effects of the initial concentration of chitosan and sonication time on
the quality of the nanoparticles ................................................. 108
3.3.1.1. Conditions that resulted in poor quality nanoparticles ............ 108
3.3.1.2. Conditions that resulted in good quality nanoparticles .......... 111
3.3.2. Effects of pH of chitosan solution and the mass ratio on the size and
charge of good quality nanoparticles ........................................... 112
3.3.3. Effects of AmB loading and freeze-drying with and without
cryoprotectants on physicochemical properties and the morphology of the
nanoparticles............................................................................ 115
3.3.4. Stability of physicochemical properties of AmB loaded chitosan
nanoparticles............................................................................ 119
3.3.5. Nanoparticles loading and encapsulation properties ............... 123
3.3.6. In vitro release of AmB from the nanoparticles ....................... 124
3.4. Discussion............................................................................. 129
3.4.1. Conclusion ......................................................................... 133
4. In vitro and in vivo activity of chitosan formulations in experimental
cutaneous leishmaniasis .............................................................. 134
4.1. Introduction ........................................................................... 134
4.2. Material and methods ......................................................... 140
4.2.1. Preparation of blank and AmB loaded chitosan nanoparticles ... 140
4.2.2. Red blood cells haemolysis.................................................. 140
4.2.3. In vitro cytotoxicity assays.................................................. 141
4.2.4. In vitro 72-hour activity of chitosan and its derivatives against
extracellular L. major and L. mexicana promastigotes ...................... 142
4.2.5. In vitro 72-hour activity of chitosan and its derivatives against
intracellular amastigotes of L. major and L. mexicana .................... 142
4.2.6. Evaluation of the in vivo anti-leishmanial activity of chitosan
formulations .............................................................................. 143
4.2.6.1. In vivo L. major model of CL .............................................. 143
4.2.6.2. Measurement of lesion size .............................................. 146
4.2.6.3. Measurement of the bioluminescence signal ....................... 147
4.2.6.4. Quantification of AmB in skin samples .............................. 147
4.2.7. Skin permeation study by Franz diffusion cell (FDC) assay ....... 148
4.2.7.1. Quantification of AmB by HPLC ....................................... 149
4.2.7.2. Fluorescence microscopy of skin sections post formulation
application .................................................................................. 150
4.2.8. Statistical analysis................................................................ 151
4.3. Results.................................................................................. 152
4.3.1. Haemolysis activity of chitosan nanoparticles ....................... 152
4.3.2. Cytotoxicity of blank and AmB loaded chitosan nanoparticles against
KB cells in RPMI (pH 7.5 and pH 6.5) ........................................... 154
4.3.3. Activity of blank and AmB loaded chitosan nanoparticles against L.
major and L. mexicana promastigotes in RPMI (pH 7.5 and pH 6.5) .... 157
4.3.4. Activity of blank and AmB loaded chitosan nanoparticles against L.
major and L. mexicana amastigotes infecting PEMs ....................... 160
4.3.5. Host cell dependence of the anti-leishmanial activity of chitosan
nanoparticles at pH of 6.5 ........................................................... 163
4.3.6. In vivo anti-leishmanial activity (intravenous route route) .................. 165
4.3.6.1. In vivo experiment 1 ........................................................................ 165
4.3.6.1.1. Evaluation of the lesion size progression ........................................ 165
4.3.6.1.2. Evaluation of the parasite load (bioluminescent signal) ............... 168
4.3.6.1.3. Intrallesional amphotericin B levels ............................................. 171
4.3.6.2. In vivo experiment 2 (dose-response effect) ..................................... 172
4.3.6.2.1. Evaluation of the lesion size progression ........................................ 172
4.3.6.2.2. Evaluation of the parasite load (bioluminescent signal) ............... 175
4.3.6.2.3. Intrallesional amphotericin B levels ............................................. 178
4.3.6.2.4. Dose concentration-response of AmB loaded chitosan-TPP nanoparticles in L. major-infected mice .............................................. 180
4.3.7. Ex vivo permeability of Leishmania-infected skin in Franz diffusion cells ................................................................. 182
4.4. Discussion ......................................................................................... 187
5. Comparison of in vitro static and dynamic culture systems to evaluate the macrophages functions and the anti-leishmanial efficacy of chitosan formulations ** ........................................... 196
5.1. Media perfusion system: an introduction ............................................ 196
5.2. Material and methods ....................................................................... 201
5.2.1. Preparation of chitosan solution and blank and AmB loaded chitosan nanoparticles ................................................................. 201
5.2.2. QV900 and media perfusion system ................................................. 202
5.2.3. Macrophages ................................................................................... 202
5.2.4. Infection of macrophages by L. major promastigotes ....................... 203
5.2.5. Measurement of macrophage functions ............................................ 203
5.2.5.1. Phagocytosis ................................................................................. 203
5.2.5.2. Macropinocytosis ......................................................................... 204
5.2.6. Evaluation of the anti-leishmanial activity of chitosan solutions, blank and AmB loaded chitosan TPP nanoparticles in the media perfusion system at pH 6.5 .................................................. 205
5.3. Results .............................................................................................. 206
5.3.1. Macrophage functions ..................................................................... 206
5.3.1.1. Phagocytosis ................................................................................. 206
5.3.1.2. Macropinocytosis ......................................................................... 210
5.3.2. Effects of media perfusion system on the anti-leishmanial activity of chitosan formulations .......................................................... 216
5.4. Discussion ......................................................................................... 219
6. General discussion .............................................................................. 224
6.1. Discussion and conclusion ............................................................... 224
6.2. Future work ...................................................................................... 229
7. References .......................................................................................... 231
8. Appendix ............................................................................................. 253
8.1. Appendix 1: Validation of HPLC methods ........................................... 253
8.2. Appendix 2: Paper 1 ......................................................................... 255
8.3. Appendix 3: Paper 2 ......................................................................... 275
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>3D</td>
<td>Three Dimensional</td>
</tr>
<tr>
<td>ADME</td>
<td>absorption, distribution, metabolism and excretion</td>
</tr>
<tr>
<td>AmB</td>
<td>Amphotericin B</td>
</tr>
<tr>
<td>AUC</td>
<td>Area under curve</td>
</tr>
<tr>
<td>BCG</td>
<td>Bacille Calmette-Guérin</td>
</tr>
<tr>
<td>BMMs</td>
<td>Murine bone-marrow macrophages</td>
</tr>
<tr>
<td>BMMs</td>
<td>Bone marrow-derived macrophages</td>
</tr>
<tr>
<td>C&lt;sub&gt;max&lt;/sub&gt;</td>
<td>Maximum concentration</td>
</tr>
<tr>
<td>CL</td>
<td>Cutaneous leishmaniasis</td>
</tr>
<tr>
<td>Clr</td>
<td>Clearance</td>
</tr>
<tr>
<td>CME</td>
<td>Clathrin-mediated endocytosis</td>
</tr>
<tr>
<td>C&lt;sub&gt;ss&lt;/sub&gt;</td>
<td>Steady-state concentration</td>
</tr>
<tr>
<td>C&lt;sub&gt;trough&lt;/sub&gt;</td>
<td>Trough plasma concentration 24 h after dose</td>
</tr>
<tr>
<td>DAPI</td>
<td>4′,6-diamidino-2-phenylindole</td>
</tr>
<tr>
<td>DCFDA</td>
<td>2′,7′–dichlorofluorescein diacetate</td>
</tr>
<tr>
<td>DCL</td>
<td>Diffuse cutaneous leishmaniasis</td>
</tr>
<tr>
<td>DCs</td>
<td>Dendritic cells</td>
</tr>
<tr>
<td>DDs</td>
<td>Drug delivery systems</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl sulfoxide</td>
</tr>
<tr>
<td>DNDi</td>
<td>Drugs for Neglected Diseases initiative</td>
</tr>
<tr>
<td>DsCL</td>
<td>Disseminated cutaneous leishmaniasis</td>
</tr>
<tr>
<td>EC&lt;sub&gt;50&lt;/sub&gt;</td>
<td>50% effective concentration</td>
</tr>
<tr>
<td>EC&lt;sub&gt;90&lt;/sub&gt;</td>
<td>90% effective concentration</td>
</tr>
<tr>
<td>ED&lt;sub&gt;50&lt;/sub&gt;</td>
<td>The required dose to achieve 50% of maximum effect</td>
</tr>
<tr>
<td>ED&lt;sub&gt;90&lt;/sub&gt;</td>
<td>The required dose to achieve 90% of maximum effect</td>
</tr>
<tr>
<td>EE</td>
<td>Encapsulation efficiency</td>
</tr>
<tr>
<td>EM</td>
<td>Emission wavelength</td>
</tr>
<tr>
<td>EX</td>
<td>Excitation wavelength</td>
</tr>
<tr>
<td>FDA</td>
<td>Federal drug agency</td>
</tr>
<tr>
<td>FDC</td>
<td>Franz diffusion cell</td>
</tr>
<tr>
<td>GFP</td>
<td>Green fluorescent protein</td>
</tr>
<tr>
<td>GM-CSF</td>
<td>Granulocyte-macrophage colony stimulating factor</td>
</tr>
<tr>
<td>GRAS</td>
<td>Generally Recognized As Safe</td>
</tr>
<tr>
<td>HiFCS</td>
<td>Heat-inactivated fetal calf serum</td>
</tr>
<tr>
<td>HMW</td>
<td>High molecular weight</td>
</tr>
<tr>
<td>HPLC</td>
<td>High Performance Liquid Chromatography</td>
</tr>
<tr>
<td>HTS</td>
<td>High throughput screening</td>
</tr>
<tr>
<td>i.p.</td>
<td>Intraperitoneal</td>
</tr>
<tr>
<td>ISC</td>
<td>Indian subcontinent</td>
</tr>
<tr>
<td>i.v.</td>
<td>Intravenous</td>
</tr>
<tr>
<td>KB cells</td>
<td>Human squamous carcinoma cells</td>
</tr>
<tr>
<td>KDa</td>
<td>Kilodaltons</td>
</tr>
<tr>
<td>kp</td>
<td>Permeability coefficient</td>
</tr>
<tr>
<td>LD&lt;sub&gt;50&lt;/sub&gt;</td>
<td>50% lethal dose</td>
</tr>
<tr>
<td>LMW</td>
<td>Low molecular weight</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Term</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------</td>
</tr>
<tr>
<td>LPS</td>
<td>Lipopolysaccharides</td>
</tr>
<tr>
<td>LR</td>
<td>Leishmaniasis recidivans</td>
</tr>
<tr>
<td>MBCL</td>
<td>Methyl-benzethonium chloride</td>
</tr>
<tr>
<td>MCL</td>
<td>Mucocutaneous leishmaniasis</td>
</tr>
<tr>
<td>MES</td>
<td>2-N-morpholino ethanesulfonic acid</td>
</tr>
<tr>
<td>MF</td>
<td>Miltefosine</td>
</tr>
<tr>
<td>MIC</td>
<td>Minimum inhibitory concentration</td>
</tr>
<tr>
<td>MMW</td>
<td>Medium molecular weight</td>
</tr>
<tr>
<td>MW</td>
<td>Molecular weight</td>
</tr>
<tr>
<td>NAC</td>
<td>N-acetyl-L-cysteine</td>
</tr>
<tr>
<td>NaOH</td>
<td>Sodium hydroxide</td>
</tr>
<tr>
<td>NCEs</td>
<td>New chemical entities</td>
</tr>
<tr>
<td>nMDP</td>
<td>Normalized mean deviation product</td>
</tr>
<tr>
<td>NMMA</td>
<td>NG-methyl-L-arginine acetate salt</td>
</tr>
<tr>
<td>NO</td>
<td>Nitric oxide</td>
</tr>
<tr>
<td>NPs</td>
<td>Nanoparticles</td>
</tr>
<tr>
<td>NTD</td>
<td>Neglected tropical disease</td>
</tr>
<tr>
<td>PBMCs</td>
<td>Peripheral blood mononuclear cells</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate-buffered saline</td>
</tr>
<tr>
<td>PC1-CH</td>
<td>Phosphorylcholine substituted chitosan</td>
</tr>
<tr>
<td>PD</td>
<td>Pharmacodynamic</td>
</tr>
<tr>
<td>PDI</td>
<td>Polydispersity index</td>
</tr>
<tr>
<td>PDMS</td>
<td>Polydimethylsiloxane</td>
</tr>
<tr>
<td>PEMs</td>
<td>Murine peritoneal macrophages</td>
</tr>
<tr>
<td>PK</td>
<td>Pharmacokinetic</td>
</tr>
<tr>
<td>PLGA</td>
<td>Polylactic-co-glycolic acid</td>
</tr>
<tr>
<td>PM</td>
<td>Paromomycin</td>
</tr>
<tr>
<td>PMN</td>
<td>Polymorphonuclear leukocytes</td>
</tr>
<tr>
<td>PV</td>
<td>Parasitophorous vacuole</td>
</tr>
<tr>
<td>QAD</td>
<td>Every other day</td>
</tr>
<tr>
<td>QD</td>
<td>Every day</td>
</tr>
<tr>
<td>QV</td>
<td>Quasi Vivo</td>
</tr>
<tr>
<td>R&amp;D</td>
<td>Research and development</td>
</tr>
<tr>
<td>RBC50</td>
<td>50% haemolytic concentration</td>
</tr>
<tr>
<td>RBCs</td>
<td>Red blood cells</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive Oxygen Species</td>
</tr>
<tr>
<td>SbV</td>
<td>Pentavalent antimony</td>
</tr>
<tr>
<td>SC</td>
<td>Stratum corneum</td>
</tr>
<tr>
<td>SD</td>
<td>Standard deviation</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning electron microscopy</td>
</tr>
<tr>
<td>t1/2</td>
<td>Half-life</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission electron microscopy</td>
</tr>
<tr>
<td>THP-1</td>
<td>Human leukemic monocyte-like derived cell line</td>
</tr>
<tr>
<td>tmax</td>
<td>Time-point corresponding with maximum concentration</td>
</tr>
<tr>
<td>TNF-α</td>
<td>Tumour necrosis factor alpha</td>
</tr>
<tr>
<td>TPP</td>
<td>Tripolyphosphate sodium</td>
</tr>
<tr>
<td>Trans-epidermal water loss</td>
<td>TEWL</td>
</tr>
<tr>
<td>----------------------------</td>
<td>--------------</td>
</tr>
<tr>
<td>VL</td>
<td>Visceral leishmaniasis</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
</tbody>
</table>
List of figures

Figure 1.1. The distribution of cutaneous leishmaniasis WHO .................... 3
Figure 1.2. *Leishmania* species and related clinical manifestations .......... 3
Figure 1.3. The life cycle of *Leishmania* parasites ................................ 7
Figure 1.4. Immune response against leishmaniasis. A: neutrophils play an important role during the early stage of infections. B: the essential role of monocytes in killing *Leishmania* and promoting the differentiation of Th-1, which leads to the elimination of parasites.............................................. 8
Figure 1.5. Strategies for treatment of CL and the related limitations. Syst=systemic. Tx= treatment. ACL=asymptomatic CL .................................. 10
Figure 1.6. Proposed structural formula for 364 Da and 365 Da ions identified by ESI(-)-MS in aqueous solutions of meglumine antimoniate and stibogluconate, respectively, copied from ........................................... 12
Figure 1.7. Chemical structure of miltefosine ........................................ 13
Figure 1.8. Chemical structure of amphotericin B (A), Fungizone (B) and AmBisome® (C) ................................................................. 15
Figure 1.9. Chemical structure of pentamidine ...................................... 16
Figure 1.10. Chemical structure of some azoles ...................................... 17
Figure 1.11. Chemical structure of paromomycin ................................... 18
Figure 1.12. The process of drug discovery and drug development. a) drug discovery stages b) drug development (107) .................................. 26
Figure 1.13. The Drug Development Pipeline - potential drugs for NTDs are frequently stuck in the early stage of development as a result of pipeline gaps ............................................................................................................. 27
Figure 1.14. New treatment candidates for leishmaniasis ....................... 28
Figure 1.15. Route a drug must take to access intracellular *Leishmania* amastigotes within macrophages (A) and DDs to intracellular *Leishmania* amastigotes(B) ........................................... 29
Figure 1.16. Administration routes of DDs and anatomical barriers .......... 32
Figure 1.17. Factors to be considered in topical delivery ......................... 35
Figure 1.18. Pathways of skin nanoparticles penetration. 1) via hair follicles, 2) intracellularly through corneocytes and 3) intercellularly around corneocytes .................................................................................................... 36
Figure 1.19. Interaction of nanoparticles with lesions of CL .................... 38
Figure 1.20. The passive diffusion of drug through a membrane ............ 39
Figure 2.1. Structure of chitin and chitosan and method of preparation chitosan from raw materials ................................................................. 48
Figure 2.2. Schematic illustration of chitosan’s versatility ....................... 49
Figure 2.3. The structure of chitosan and its derivatives ......................... 57
Figure 2.4. Dose-response curves of the cytotoxicity of chitosan and its derivatives against KB cells at pH=7.5 (A) and 6.5 (B) ...................... 67
Figure 2.5. Dose-response curves of the activity of chitosan and its derivatives against *Leishmania* promastigotes at two pH values. A: *L. major* at pH=7.5; B: *L. mexicana* at pH = 7.5; C: *L. major* at pH = 6.5; D: *L.*
mexicana at pH = 6.5. Promastigotes were cultured in the presence of different concentrations of chitosan and its derivatives. .......................... 70

Figure 2.6. Dose-response curves of the activity of chitosan and its derivatives against Leishmania amastigotes at two pH values. A: L. major at pH=7.5; B: L. mexicana at pH = 7.5; C: L. major at pH = 6.5; D: L. mexicana at pH = 6.5. ........................................................................................................... 73

Figure 2.7. Morphology of infected (PEMs) with L. major and L. mexicana after treatment with HMW chitosan ........................................................................................................... 74

Figure 2.8. Influence of incubation duration on the chitosan and Fungizone activity against L. major intracellular amastigotes in BMMs. .............................. 76

Figure 2.9. TNF-α production in uninfected and L. major infected BMMs, PEMs and THP-1 macrophages* after 24 h of exposure to 1.64, 4.9,14.8, 44.4, 133.3 and 400 µg/ml of chitosan at pH = 6.5 ........................ ........................................ 78

Figure 2.10. ROS production in uninfected and L. major infected BMMs, PEMs and THP-1 macrophages * after 4 h of exposure to 1.64, 4.9,14.8, 44.4, 133.3 and 400 µg/ml of HMW chitosan at pH=6.5 .............................. 80

Figure 2.11. Activity of HMW chitosan against L. major amastigotes in BMMs* after 4 h, with and without ROS scavenger at pH = 6.5...................... 81

Figure 2.12. NO production in uninfected and L. major infected BMMs, PEMs and THP-1 macrophages * after 24 h of exposure to 1.64, 4.9,14.8, 44.4, 133.3 and 400 µg/ml of chitosan at pH = 6.5 .......................... 84

Figure 2.13. Activity of HMW chitosan against L. major -infected BMMs* after 24 h in the presence or absence of an NO inhibitor at pH = 6.5 ......... 85

Figure 2.14. Activity of HMW chitosan against L. major infected BMMs* after 4 h, pH=6.5 (A), 24 h, pH=6.5 (B) and at 24 h, pH=7.5 with or without phagocytosis inhibitor or pinocytosis (CME) inhibitor. .............................. 87

Figure 2.15. Fluorescence microscopy images of the cellular uptake of rhodamine-labelled chitosan over 4 and 24 h at pH=6.5 by BMMs infected with L. major (XA) or with L. mexicana (XB)............................................ 89

Figure 3.1. Preparation methods of chitosan nanoparticles .......................... 98

Figure 3.2. Chemical structure of TPP and dextran sulphate ................. 100

Figure 3.3. Mechanisms of drug release from chitosan nanoparticles. .. 100

Figure 3.4. Schematic representation for Blank nanoparticles and AmB loaded chitosan nanoparticles with TPP or dextran sulphate .......... 104

Figure 3.5. TEM micrographs of unloaded and amphotericin B loaded chitosan nanoparticles .............................................. 110

Figure 3.6. TEM micrographs of unloaded and amphotericin B loaded chitosan nanoparticles .............................................. 110

Figure 3.7. TEM micrographs of unloaded and amphotericin B loaded chitosan nanoparticles .............................................. 111

Figure 3.8. Effectiveness of sucrose 5% and D-mannitol 5% as a cryoprotectant for freeze drying of blank and AmB loaded chitosan nanoparticles .......................................................... 117

Figure 3.9. TEM micrographs of unloaded and amphotericin B loaded chitosan nanoparticles .............................................. 118
Figure 3.10. SEM micrographs of unloaded and amphotericin B loaded chitosan nanoparticles

Figure 3.11. Size of AmB loaded chitosan-TPP nanoparticle (A) and AmB loaded chitosan-dextran sulphate nanoparticle (B) in different media over time.

Figure 3.12. Comparison of AmB encapsulation, loading and yield of the two types of nanoparticles.

Figure 3.13. In vitro release profile of AmB loaded chitosan nanoparticles at 37 °C.

Figure 5.1. Microfluidic system (A) The integrated perfusion culture micro-chamber array chip. (B) Enlarged view of a micro-chamber array unit.

Figure 5.2. Kirkstall LTD. Quasi Vivo 900 media perfusion system in use circulating RPMI 1640 media.

Figure 5.3. Schematic overview of evaluation of the anti-leishmanial activity in static and flow culture systems.

Figure 5.4. Phagocytosis of fluorescent latex beads (2 μm) by uninfected and infected PEMs (A), BMMs (B) and THP-1 (C) in static culture system.

Figure 5.5. Phagocytosis of fluorescent latex beads (2 μm) by infected PEMs (A), BMMs (B) and THP-1 (C) in the three culture systems (static, slow flow rate 1.45 x 10^{-9} m/s and fast flow rate 1.23 x 10^{-7} m/s).

Figure 5.6. Macropinocytosis of pHrodo Red dextran by uninfected and infected PEMs (A), BMMs (B) and THP-1 (C) in static culture system.

Figure 5.7. Macropinocytosis of pHrodo Red dextran by infected PEMs (A), BMMs (B) and THP-1 (C) at the three culture systems (static, slow flow rate 1.45 x 10^{-9} m/s and fast flow rate 1.23 x 10^{-7} m/s).

Figure 5.8. Dose-response curve of the activity of chitosan solution (A), blank chitosan-TPP nanoparticles (B), AmB loaded chitosan-TPP nanoparticles (C) and AmB solution (pure) (D) against L. major amastigotes infecting PEMs in pH=6.5 under different flow conditions.

Figure 5.9. Sedimentation under a) static conditions, b) homogeneous distribution of drugs under flow conditions.

Figure 5.10. Accumulation of amphotericin B (left) and miltefosine (right) in peritoneal macrophages at three culture systems using the QV900 over time. Static and two flow rates (1.33 x 10^{-9} at the base of the chamber or 1.17 x 10^{-7} m/s) on an insert.

Figure 5.11. A Hollow-Fibre model.
List of tables

Table 1.1. Clinical and epidemiological characteristics of the main Leishmania species copied from (4) ................................................................. 4
Table 1.2. Clinical features of New World CL that might modify management copied from (23) ......................................................................................... 9
Table 1.3. In vitro screening models with positive and negative drawbacks copied from (97). .................................................................................................. 22
Table 1.4. In vivo models for leishmaniasis copied from (97) ....................... 24
Table 1.5. Experimental studies using nanosystems for CL treatment copied from (115). ............................................................................................................ 30
Table 1.6. Chitosan-based drug delivery systems ........................................ 34
Table 1.7. Basic PK parameters copied from (71) ........................................ 41
Table 1.8. Pharmacokinetic profile of leishmaniasis drugs (70, 170) ............ 43
Table 2.1. The antimicrobial activities of chitosan and its derivatives (173, 185, 186, 187) ............................................................................................. 51
Table 2.2. The anti-leishmanial activity of chitosan .................................... 53
Table 2.3. Details of chitosan and its derivatives used in the study ............. 55
Table 2.4. In vitro cytotoxicity of chitosan and its derivatives against KB cells at two pH values after 72h of incubation ...................................................................... 66
Table 2.5. In vitro activity of chitosan and its derivatives against promastigotes at two pH values after 72h of incubation ........................................... 69
Table 2.6. In vitro activity of chitosan and its derivatives against amastigotes infecting PEMs after 72h of incubation ...................................................... 72
Table 2.7. In vitro activity of chitosan against promastigotes based on molarity ...................................................................................................................... 74
Table 2.8. In vitro activity of chitosans against amastigotes based on molarity ...................................................................................................................... 74
Table 2.9. HMW chitosan activity against L. major amastigotes in three different macrophage cultures after 72 h ............................................................ 75
Table 2.10. ROS production in uninfected and L. major infected BMMs after 8 h of exposure to different concentrations of HMW chitosan at pH=6.5 ...... 79
Table 2.11. ROS production in uninfected and L. major infected BMMs after exposure to chitosan in the presence of ROS scavenger ............................ 82
Table 2.12. NO production in uninfected and L. major-infected BMMs after 4h of exposure to different concentrations of HMW chitosan at pH=6.5 ...... 83
Table 2.13. NO production in uninfected and L. major infected BMMs after exposure to chitosan in the presence of NO inhibitor at pH=6.5 ............ 85
Table 2.14. Phagocytosis and pinocytosis by L. major infected BMMs in the presence of the uptake inhibitors ................................................................. 88
Table 3.1. HPLC parameters for AmB quantification .................................. 108
Table 3.2. Conditions which did not produce good quality nanoparticles .. 109
Table 3.3. Effect of pH and concentration of chitosan and mass ratio of the reactants on the physicochemical properties of blank chitosan-TPP nanoparticles .................................................................................................. 113
Table 3.4. Effect of pH and concentration of chitosan and mass ratio of the reactants on the physicochemical properties of blank chitosan-dextran sulphate nanoparticles ......................................................... 114

Table 3.5. Effect of cryoprotectants used during freeze drying on the physicochemical properties of unloaded and AmB loaded chitosan nanoparticles .............................................................................. 116

Table 3.6. Variations of physicochemical properties of AmB loaded chitosan-TPP nanoparticles in different media upon storage at different temperatures ........................................................................ 121

Table 3.7. Variations of physicochemical properties of AmB loaded-chitosan dextran sulphate nanoparticles in different media upon storage at different temperatures ....................................................................... 122

Table 3.8. Percentage of AmB loading, encapsulation and yield ........................................ 123

Table 3.9. In vitro cumulative release of AmB from the two formulations at different conditions .......................................................................................................................... 125

Table 4.1. Disadvantages of different DDs (67, 112, 279, 280, 281, 282, 283, 284) .............................................................................................................................. 135

Table 4.2. Antimicrobial activity of chitosan nanoparticles (285) .................................... 137

Table 4.3. Anti-leishmanial activity of AmB loaded chitosan nanoparticles ........................ 138

Table 4.4. In vitro haemolytic activity of chitosan formulations after 1h of incubation ................................................................. 152

Table 4.5. In vitro cytotoxicity of chitosan formulations against KB cells at two pH values after 72h of incubation ................................. 154

Table 4.6. In vitro activity of chitosan formulations against promastigotes at two pH values .................................................................................................................. 158

Table 4.7. In vitro activity of chitosan formulations against intracellular amastigotes at two pH values .................................................. 161

Table 4.8. Activity of chitosan formulations against L. major amastigotes in three different macrophage cultures after 72 h at pH of 6.5 ..................................... 164

Table 4.9. Flux, lag time and the permeability coefficient (kp) for AmB loaded chitosan nanoparticles ................................................................. 184

Table 4.10. Disposition of topically applied AmB loaded chitosan nanoparticles on healthy and L. major infected BALB/c mice skin using Franz diffusion cells ............................................. 185

Table 5.1. Specifications of QV500 and QV900 media perfusion system (340, 341, 342) .................................................................................................................. 201

Table 5.2. Phagocytosis of fluorescent latex beads (2 μm) by uninfected and infected PEMs, BMMs and THP-1 in static culture system .................................. 207

Table 5.3. Phagocytosis of fluorescent latex beads (2 μm) by infected PEMs, BMMs and THP-1 in the three culture systems (static, slow flow rate 1.45 x 10^{-9} m/s and fast flow rate 1.23 x 10^{-7} m/s ) ........................................ 209

Table 5.4. Macropinocytosis of pHrodo™ Red dextran by uninfected and infected PEMs, BMMs and THP-1 in static culture system .................................. 212

Table 5.5. Macropinocytosis of pHrodo™ Red dextran by infected PEMs, BMMs and THP-1 at the three culture systems (static, slow flow rate 1.45 x 10^{-9} m/s and fast flow rate 1.23 x 10^{-7} m/s ) ........................................ 215
Table 5.6. *In vitro* activity of chitosan solution and nanoparticles against *L. major* amastigotes in RPMI medium (pH=6.5) at different flow rates ........ 217
Table 8.1. HPLC validation parameters.................................................. 253
Table 8.2. The precision of AmB HPLC assay ................................. 254
1. General introduction

1.1. Leishmaniasis and *Leishmania* species

Leishmaniasis is an infectious disease caused by parasites belonging to the genus *Leishmania* in the family Trypanosomatidae. *Leishmania* parasites are transmitted to mammals through the bite of sandflies that belong to the genus *Phlebotomus* (Old World) or *Lutzomyia* (New World). *Leishmania* species cause two main clinical forms, cutaneous leishmaniasis (CL) and visceral leishmaniasis (VL) (1). CL is the most common type of leishmaniasis and in addition to “simple” CL, there are other complex cutaneous leishmaniasis manifestations including mucocutaneous leishmaniasis (MCL), diffuse cutaneous leishmaniasis (DCL), disseminated cutaneous leishmaniasis (DsCL) and leishmaniasis recidivans (LR) (1, 2). CL is caused by *Leishmania* species that are classified into Old World species, for instance *Leishmania major* (*L. major*), *L. tropica*, and *L. aethiopica* and New World species, such as *L. amazonensis*, *L. mexicana*, *L. braziliensis* and *L. guyanensis* (Fig 1.1) (3, 4). CL occurs in 88 countries and 90% of the cases are reported in Afghanistan, Brazil, Iran, Peru, Saudi Arabia and Syria (Fig 1.2) (1). Recently, a recrudescence has been noticed in Syria as a result of the destruction of the public health system and the lack of sanitation caused by the current conflict (5). Because of the displacement of Syrian people and the millions forced to flee the country, with the majority of them residing in Lebanon, Jordan, Egypt and Iraq, reporting of CL has increased across the region (6).

The clinical features of leishmaniasis depend on the parasite, the host and the vectors – Fig 1.2 shows an overview of the taxonomy of *Leishmania* species and the related clinical manifestations (Fig 1.1 and Table 1.1) (7).

- VL, also known as kala-azar (black fever), a potentially fatal illness which is characterised by irregular fever lasting for 14 days, the enlargements of spleen and liver, pancytopenia and weight loss. The incubation time for VL is between 2 weeks and 8 months and without treatment, the disease is typically fatal. One of the big challenges for
VL is the co-infection with HIV. VL is caused mainly by *L. donovani*, *L. infantum* and rarely by *L. tropica* (8).

- LCL is associated with an erythemic papule at the bitten site (1-10 mm diameter) and then can lead to rounded ulcers combined with nodal or thick edges. These ulcers or lesions can stay from 5 months to 20 years. Lesions caused by *L. mexicana* are typically self-healing within 3-9 months, 6-15 months in the case of *L. braziliensis*, *L. tropica* or *L. panamensis* and within 2-6 months for *L. major* infections (8).

- DCL is uncommon anergic dissemination form of CL caused by *L. aethiopica*, *L. amazonensis* or *L. mexicana*. It begins with erythematous nodules resembling lepromatous leprosy and infiltrative plaques and then might ulcerate. DCL starts firstly on the face and subsequently affects other parts of the body and could affect the complete skin surfaces in some cases (8).

- MCL is caused by *L. braziliensis*, *L. guyanensis*, or *L. panamensis*. MCL is identified by invasive and destructive lesions of the mucosal membrane of the face, mouth and throat cavities. MCL is more frequent in immunocompromised patients (4).

- DsCL is caused by *L. aethiopica*, *L. guyanensis* and *L. mexicana*, spotted in Latin America and characterised by ten or more lesions (mixed type) located in two or more parts of the body.

- LR is caused by *L. tropica* and *L. braziliensis* and usually identified as new lesions around the old scar that has been cured and infiltrated with lymphocytes.
Figure 1.1. The distribution of cutaneous leishmaniasis WHO (9)

Figure 1.2. Leishmania species and related clinical manifestations (7).
**Table 1.1. Clinical and epidemiological characteristics of the main *Leishmania* species copied from (4)**

<table>
<thead>
<tr>
<th>Subgenus</th>
<th>Clinical form</th>
<th>Main clinical features</th>
<th>Natural progression</th>
<th>Risk groups</th>
<th>Main reservoir</th>
<th>High-burden countries or regions</th>
<th>Estimated annual worldwide incidence</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Leishmania</em></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>donovani</em></td>
<td>VL and PKDL</td>
<td>Persistent fever, splenomegaly, weight loss, and anaemia in VL; multiple painless macular, papular, or nodular lesions in PKDL</td>
<td>VL is fatal within 2 years; PKDL lesions self-heal in up to 85% of cases in Africa but rarely in Asia</td>
<td>Predominantly adolescents and young adults for VL; young children in Sudan and no clearly established risk factors for PKDL</td>
<td>Humans</td>
<td>India, Bangladesh, Ethiopia, Sudan, and South Sudan</td>
<td>50 000–90 000 VL cases; unknown number of PKDL cases</td>
</tr>
<tr>
<td>tropica</td>
<td>CL, LR, and rarely VL</td>
<td>Ulcerating dry lesions, painless, and frequently multiple</td>
<td>CL lesions often self-heal within 1 year</td>
<td>No well defined risk groups</td>
<td>Humans but zoonotic foci exist</td>
<td>Eastern Mediterranean, the Middle East, and northeastern and southern Africa</td>
<td>200 000–400 000 CL</td>
</tr>
<tr>
<td>aethiopica</td>
<td>CL, DCL, DsCL, and oronasal CL</td>
<td>Localised cutaneous nodular lesions; occasionally oronasal; rarely ulcerates</td>
<td>Self-healing, except for DCL, within 2–5 years</td>
<td>Limited evidence; adolescents</td>
<td>Hyraxes</td>
<td>Ethiopia and Kenya</td>
<td>20 000–40 000 CL</td>
</tr>
<tr>
<td>major</td>
<td>CL</td>
<td>Rapid necrosis, multiple wet sores, and severe inflammation</td>
<td>Self-healing in &gt;50% of cases within 2–8 months; multiple lesions slow to heal, and severe scarring</td>
<td>No well defined risk groups</td>
<td>Rodents</td>
<td>Iran, Saudi Arabia, north Africa, the Middle East, central Asia, and west Africa</td>
<td>230 000–430 000 CL</td>
</tr>
<tr>
<td><em>infantum</em></td>
<td>VL and CL</td>
<td>Persistent fever and splenomegaly in VL; typically single nodules and minimal inflammation in CL</td>
<td>VL is fatal within 2 years; CL lesions self-heal within 1 year and confers individual immunity</td>
<td>Children under 5 years and immunocompromised adults for VL; older children and young adults for CL</td>
<td>Dogs, hares, and humans</td>
<td>China, southern Europe, Brazil, and South America for VL and CL; Central America for CL</td>
<td>6200–12 000 cases of Old World VL and 4500–6800 cases of New World VL; unknown number of CL cases</td>
</tr>
<tr>
<td>Subgenus</td>
<td>Clinical form</td>
<td>Main clinical features</td>
<td>Natural progression</td>
<td>Risk group</td>
<td>Main reservoir</td>
<td>High-burden countries or regions</td>
<td>Estimated annual worldwide incidence</td>
</tr>
<tr>
<td>------------------</td>
<td>---------------</td>
<td>------------------------------------------------------------</td>
<td>------------------------------------------</td>
<td>-----------------------------------------</td>
<td>-------------------------</td>
<td>--------------------------------------</td>
<td>--------------------------------------</td>
</tr>
<tr>
<td><em>Leishmania</em></td>
<td><em>Leishmania</em></td>
<td>CL, DCL, and DsCL</td>
<td>Ulcerating lesions, single or multiple</td>
<td>Often self-healing within 3-4 months</td>
<td>No well defined risk groups</td>
<td>Rodents and marsupials</td>
<td>Limited number of cases</td>
</tr>
<tr>
<td><em>Leishmania</em></td>
<td><em>Leishmania</em></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Possums</td>
<td>Limited number of cases</td>
</tr>
<tr>
<td><em>amazonensis</em>†</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>South America</td>
<td></td>
</tr>
<tr>
<td><em>Leishmania</em></td>
<td><em>Viannia</em></td>
<td>CL, MCL, DCL, and LR</td>
<td>Ulcerating lesions can progress to mucocutaneous form; local lymph nodes are palpable before and early on in the onset of the lesions</td>
<td>Might self-heal within 6 months; 2-5% of cases progress to MCL</td>
<td>No well defined risk groups</td>
<td>Dogs, humans, rodents, and horses</td>
<td>Majority of the 187 200–300 000 total cases of New World CL‡</td>
</tr>
<tr>
<td><em>braziliensis</em>†</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>South America</td>
<td></td>
</tr>
<tr>
<td><em>Leishmania</em></td>
<td><em>Viannia</em></td>
<td>CL, DsCL, and MCL</td>
<td>Ulcerating lesions, single or multiple that can progress to mucocutaneous form; palpable lymph nodes.</td>
<td>Might self-heal within 6 months²</td>
<td>No well defined risk groups</td>
<td>Possums, sloths, and anteaters</td>
<td>Limited number of cases, included in the 187 200–300 000 total cases of New World CL‡</td>
</tr>
<tr>
<td><em>guyanensis</em>†</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>South America</td>
<td></td>
</tr>
</tbody>
</table>

VL=visceral leishmaniasis. PKDL=post-kala-azar dermal leishmaniasis. CL=cutaneous leishmaniasis. LR=leishmaniasis recidivans. DCL=diffuse cutaneous leishmaniasis. DsCL=disseminated cutaneous leishmaniasis. MCL=mucocutaneous leishmaniasis. *Old World leishmaniasis. †=New World leishmaniasis. ‡Estimates are of all New World leishmaniases, with Leishmania braziliensis comprising the vast majority of these cases.
1.2. Life cycle

Many causative species for CL, have a zoonotic cycle\(^1\) (*L. major, L. aethiopica*, and all the New World species), whilst few have an anthroponotic cycle\(^2\) (*L. tropica*). Regarding the VL, humans are the main reservoir for *L. donovani* while dogs form the primary reservoir for *L. infantum*. The *Leishmania* life cycle starts when infected female sandflies (*Phlebotomus* species in the Old World, *Lutzomyia* species in the New World) bite their hosts and inject parasites (the infective metacyclic promastigote form) into the skin of a mammalian host (a sand fly injects 100-1000 promastigotes). Sandflies salivary chemoattractants enhance the flow of macrophages, dendritic cells (DCs) and neutrophils to the biting site. These promastigotes are then phagocytised by resident phagocytes. After which, promastigotes transform in these cells into amastigotes which replicate by simple division in the parasitophorous vacuole and infect other macrophages, either locally or in remote tissues (1, 4). Neutrophils play a critical role in leishmaniasis by acting as Trojan horses for *Leishmania* promastigotes before entering their target cells (macrophages). *Leishmania* survive in the neutrophils by inhibiting the phagosome acidification. *Leishmania* promastigotes directly infect DCs and reside within parasitophorous vacuoles. In macrophages, promastigotes are interlined into phagolysosome like compartment, named the *Leishmania* parasitophorous vacuole. The maturation of parasitophorous vacuole is regulated by *Leishmania* parasites to protect them from destruction by the macrophage microbicidal activity and to avoid the host immune defence responses (9, 10, 11).

Female sandflies become infected when they feed on an infected host and amastigotes transform into promastigotes in the midgut of the sandfly and then migrate to salivary glands and transform into infectious metacyclic promastigotes (Fig 1.3) (1, 12).

---

\(^1\) In zoonotic cycles: animals are main reservoirs  
\(^2\) In anthroponotic cycles: humans are main reservoirs
1.3. Immune response in CL

The cellular immune responses play a critical role in the control or progress of cutaneous leishmaniasis and have been widely studied in mouse models, often using *L. major*. Progressive lesions have been developed in susceptible mice (BALB/c mice) with a dominance of the Th2 response, leading to the production of anti-inflammatory cytokines, such as IL-4, IL-5, and IL-13, which suggests that Th2 cells are associated with developing progressive lesions. On the other hand, resistant mice (C57BL/6 and C3H/HeJ mice), infected by *L. major*, present small lesions with few parasites and a dominance of the Th1 response, with the production of IFN-γ, TNF-α and IL-12. These cytokines activate macrophages to produce reactive oxygen species (ROS) and nitric oxide (NO), which are responsible for killing intracellular parasites as seen in Fig 1.4 (14, 15, 16). In humans, resolution from cutaneous leishmaniasis is recognized by induction of specific IFN-γ releasing CD4+ T cells (17, 18). The response in individuals with moderate CL caused by *L. major* is a mixture of Th1 and Th2. There is an absence of a Th1 response in individuals with severe CL (17, 18).
To conclude, the control of CL is linked with *Leishmania*-specific T lymphocytes producing TNF-α and IFN-γ and this enhances macrophages in the skin to produce microbicidal materials (NO and ROS). It is obvious from Fig 1.5 that, the balance between pro- and anti-inflammatory factors controls the consequence of CL infection (19).

The functions of B cells are still a matter of debate. Several studies suggest that these cells enhance the *Leishmania* infection while some state that B cells have a protective function against *L. amazonensis* (20).

![Figure 1.4. Immune response against leishmaniasis. A: neutrophils play an important role during the early stage of infections. B: the essential role of monocytes in killing *Leishmania* and promoting the differentiation of Th-1, which leads to the elimination of parasites (16)](image)

### 1.4. Current treatment of cutaneous leishmaniasis

CL lesions can heal spontaneously in most cases within 2-18 months. Infection is not usually fatal but can cause considerable cosmetic morbidity, psychological disorders, social stigma leading to changes in individual self-esteem (4, 8). The important goal of making the decision to treat CL is to eradicate the *Leishmania* parasites and enhance the lesion healing process. This will reduce the risk of scarring and help to lower the risk of dissemination or progression other forms of more severe CL.
Other criteria to commence treatment includes the presence of many lesions (more than 5), large size (>4 cm), location over sensitive body areas such as the face, or lasting for more than 6 months and/or in Immunosuppressed patients (Table 1.2) (4, 21). The Infectious Diseases Society of America recently published comprehensive treatment guidelines for the management of CL according to the clinical characteristics, summarised in Table 1.2 (22).

**Table 1.2.** Clinical features of New World CL that might modify management copied from (23)

<table>
<thead>
<tr>
<th>Simple CL</th>
<th>Complex CL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caused by a <em>Leishmania</em> species unlikely to be associated with mucosal leishmaniasis</td>
<td>Caused by a <em>Leishmania</em> species that can be associated with increased risk for ML, particularly <em>Viannia</em> spp in the “mucosal belt” of Bolivia, Peru, and Brazil $^{a,b,c}$</td>
</tr>
<tr>
<td>No mucosal involvement noted</td>
<td>Local subcutaneous nodules $^{d}$</td>
</tr>
<tr>
<td>Absence of characteristics of complex CL</td>
<td>Large regional adenopathy $^{d}$</td>
</tr>
<tr>
<td>Only a single or a few skin lesions</td>
<td>&gt;4 skin lesions of substantial size (eg, &gt;1 cm)</td>
</tr>
<tr>
<td>Small lesion size (diameter)</td>
<td>Large individual skin lesion (diameter ≥5 cm)</td>
</tr>
<tr>
<td>Location of lesion feasible for local treatment</td>
<td>Size or location of lesion such that local treatment is not feasible</td>
</tr>
<tr>
<td>Nonexposed skin (ie, not cosmetically important)</td>
<td>Lesion on face, including ears, eyelids, or lips; fingers, toes, or other joints; or genitalia</td>
</tr>
<tr>
<td>Immunocompetent host</td>
<td>Immunocompromised host (especially with respect to cell-mediated immunity)</td>
</tr>
<tr>
<td>Lesion(s) resolving without prior therapy</td>
<td>Clinical failure of local therapy</td>
</tr>
<tr>
<td></td>
<td>Unusual syndromes: leishmaniasis recidivans, diffuse CL, or disseminated CL</td>
</tr>
</tbody>
</table>

Abbreviation: CL, cutaneous leishmaniasis.

$a$ The highest risk areas for mucosal leishmaniasis (ML) are south of the Amazon basin in parts of Bolivia, Peru, and Brazil (defined here as the “mucosal belt”). Moderate-risk areas are south of Nicaragua to the Amazon basin. Low-risk areas for ML are in New World CL (*Viannia*)–endemic regions north of Costa Rica. Amazonian basin regions up to an altitude of approximately 2000 meters are referred to as increased ML-risk regions.

$b$ *Leishmania* species with an increased risk of causing ML include *L. (V.) braziliensis* mainly, but also *L. (V.) guyanensis* and *L. (V.) panamensis*. There are other species that can be associated with ML less frequently. In this document, we refer to these 3 species as “increased ML-risk species.” Geographic regions in which there is an increased risk for ML are defined above.

$c$ High therapeutic failure rates after treatment with pentavalent antimonial drugs have been observed in CL acquired in Amazonian Bolivia (eg, Madidi National
Park) and southeastern Peru (eg, Manu National Park and Puerto Maldonado). Poor efficacy after using miltefosine in the treatment of *L. (V.) braziliensis* was reported in Guatemala.

d It is somewhat controversial whether the presence of small subcutaneous nodules is always associated with complex CL, but certainly complex CL applies if bubonic-like adenopathy is present in regional drainage area of lesions. These findings have been linked to complications or treatment failure when only local treatment is administered. Some experts would not consider systemic therapy needed for a few, small subcutaneous nodules in Old World CL.

Treating CL can include (i) chemotherapy (anti-leishmanial drugs that kill the parasites directly) (ii) local physical methods (cryo- or thermotherapy), (iii) immunotherapy (by immune modulators for stimulating effective immune response against *Leishmania* parasites) (Fig 1.5) (24).

![Diagram of CL treatment options and gaps](image)

**Figure 1.5.** Strategies for treatment of CL and the related limitations. Syst=systemic. Tx= treatment. ACL=asymptomatic CL (25).
1.4.1. Systemic therapies

1.4.1.1. Pentavalent Antimony

Pentavalent antimony (SbV) compounds like sodium stibogluconate (SSG, Pentostam®, GSK, contains 100 mg/ml of SbV) and meglumine antimoniate (Fig 1.6) (MA, Glucantime®, Sanofi, contains 85 mg/ml) have been the standard therapy for CL since they were developed in the 1940s (26, 27). The severity of CL can determine the routes of administration (locally or systemically). In local treatment, SbV (1-5 ml) is administrated by injection (1-5 times every 3-7 days for up to 5 sessions) in lesions edges with or without cryotherapy (application of liquid nitrogen after the injection) (28, 29). The parenteral route includes intravenous or intramuscular administration of 20 mg/kg/day of SbV, typically in the case of complex CL (28). Intralesional administration benefits include making a high enough concentration of the drug at the site of infection, reduced costs, limiting the systemic side effects and faster healing time (30). However, the problems with this route includes the difficulty of administration, pain of these injections, sensations of burning, itching and sometimes the appearance of inflammation in the location of the injections (31). On the other hand, parenteral injections can lead to adverse side effects (hepatotoxicity and cardiotoxicity) (32). There is a lack of placebo-controlled randomized clinical trials to compare the activity of SbV therapy against specific species of CL (31). Variability of the sensitivity of *Leishmania* (promastigotes and intracellular amastigotes) species to SbV has been confirmed *in vitro* (33, 34).

There is still no clear definition of the mechanism of the action of SbV, in spite of these drugs being used for several decades. One of the suggested mechanisms is that SbV is converted after administration to the trivalent form (SbIII) which is the active but more toxic form. This trivalent antimony (SbIII) intervenes with the trypanothione reductase system that protects the *Leishmania* amastigotes from the harm caused by the oxidation and toxicity of heavy metals (35, 36). Others suggested that SbIII can cause *Leishmania* apoptosis by fragmenting DNA of amastigotes (37, 38). A third mechanism has
suggested that SbV interacts with adenine ribonucleoside and produces a complex that causes a depletion of intracellular ATP and the prevention of macromolecule synthesis in amastigotes by inhibiting type I DNA topoisomerase (32, 39).

![Proposed structural formula for 364 Da and 365 Da ions identified by ESI(−)-MS in aqueous solutions of meglumine antimoniate and stibogluconate, respectively, copied from (40)](image)

**Figure 1.6.** Proposed structural formula for 364 Da and 365 Da ions identified by ESI(−)-MS in aqueous solutions of meglumine antimoniate and stibogluconate, respectively, copied from (40)

1.4.1.2. Miltefosine

Miltefosine (MF), an alkylphospholipid, was developed as an antineoplastic agent (for cutaneous cancers). Croft *et al.* in 1987 showed the anti-leishmanial activity of miltefosine and other phospholipid compounds (1). MF is recommended for VL and complex cases of CL and considered as the only effective drug that can be given orally for leishmaniasis treatment. The effective dose for CL is a daily oral dose 2.5 mg/kg for 28 days (1, 41).

However, different *Leishmania* species show significantly different sensitivity to MF (42). Randomized clinical trials have been conducted in different regions against different species with various clinical responses. For instance, in Colombia the cure rates against *L. panamensis* were 91% in comparison to 38% for placebo group (43). While the cure rates were just 53% in Guatemala
against *L. mexicana* and *L. braziliensis* compared to 32% in placebo treatment (43). *In vitro* studies confirmed the species variation in MF sensitivity (44). The two major concerns about this drug is that, (i) MF is a teratogenic agent and so cannot be given to women who are pregnant and (ii) the presence of resistance development *in vitro* (42). Some common side effects of MF treatment are gastrointestinal discomfort, renal disorders, headache and raised liver enzymes (45).

The mechanism of action of MF remains unclear. Different mechanisms have been suggested such as the inhibition of synthesis of phospholipids, interaction with the parasite membrane, dysfunction of mitochondria or stimulation of apoptosis-like cell death (46, 47, 48, 49).

**Figure 1.7.** Chemical structure of miltefosine (50)

**1.4.1.3. Amphotericin B (AmB)**

The second most common treatment for leishmaniasis is amphotericin B, which is a polyene antibiotic (Fig 1.8), mainly used for VL and MCL (51). The therapeutic dose of AmB deoxycholate (Fungizone) is 0.7 mg/kg/day by slow intravenous infusion for 25-30 days or 2-3 mg/kg/day of liposomal formulations for 10-15 days (28). In 1950s, AmB was firstly noted and derived from *Streptomyces nodosus*. Sodium deoxycholate solution of AmB (DAmB, Fungizone) was brought to the market in 1959. Fungizone has been used intravenously as a standard treatment for invasive fungal infections for several decades. Fungizone has serious side effects such as nephrotoxicity and fever, anaemia, malaise and abdominal pain (52).

Several lipid formulations, including liposomal amphotericin B (AmBisome®), amphotericin B lipid complex (Abelcet®), and amphotericin B colloidal
dispersion (Amphocil™) have been developed and used in treatment of VL to reduce the previous toxicities since 1990s (53). AmBisome® (liposomal amphotericin B, LAmB; Gilead Sciences, Dimas, CA, USA) has been approved by the Food and Drug Administration (FDA) for treatment of VL in 1997 with 7 intravenously doses of 3 mg/kg/day over 21 days (54). Yardley and Croft (2000) found that AmBisome® (liposomal amphotericin B) was also successful in reducing the size of lesions in CL caused by L. major in BALB/c mouse model (55). The high cost of these formulations (up to 250 USD$ per vial) prevents more widespread use (56, 57). Recently there is an agreement between WHO and Gilead Sciences for the donation through WHO of AmBisome® vials for VL treatment (58). Other problems related to AmBisome® were reported in a study with a low positive outcome of 63% among travellers infected with CL and MCL coming back from both Old- and New-World countries and 53% of these treated patients showed renal toxicity and infusion-related reactions (59) and higher rates of relapse were noticed in immunocompetent patients with VL treated with AmBisome® (60, 61).

Amphotericin B acts by forming a complex between its hydrophobic polyene region and the ergosterol in the plasma membrane of Leishmania or fungi which causes transmembrane channels, after which a death of the microbe is induced by the collapsing of ion gradient (62, 63). Recently, an alternative mode of action has been suggested, that AmB primarily exists as large, extra-membranous aggregates that results in the removal of ergosterol from the lipid bilayer leading to microbe death (64). Additionally, reports claim that AmB has immunomodulatory effects and stimulates oxidative stress in immune cells (52, 65, 66).
1.4.1.4. Pentamidine

Pentamidine is an aromatic diamidine, as effective as antimonial drugs for healing CL caused by *L. panamensis* or *L. guyanensis* (30, 57). The cure rates of parenteral pentamidine with 7 doses of 2 mg/kg for 14 days vary from 35% to 95% (1). This drug offered significant advantages such as shorter duration
of the therapy and lower costs in comparison with other drugs for CL (68) but is rarely used due to low cure rates and significant side effects of diabetes, myocarditis and nephrotoxicity (69).

The mode of action is not completely clear but interference with *Leishmania* DNA and disruption of mitochondrial membrane have been suggested (70).

![Chemical structure of pentamidine](image)

**Figure 1.9.** Chemical structure of pentamidine (71)

### 1.4.1.5. Azoles

Azoles are antifungal agents, which also have an anti-leishmanial activity because they inhibit the 14 a-demethylation of lanosterol and this inhibition leads to an accumulation of 14 a-methyl sterols and blocks ergosterol synthesis of *Leishmania* parasites (72). The most important azoles that are active against *Leishmania* parasites are fluconazole, ketoconazole and itraconazole which have been used orally with different results against CL. The effectivity of ketoconazole, with the oral dose 8 mg/kg/day for 4 to 6 weeks, was 76–90% in CL caused by *L. panamensis* and *L. mexicana* in Guatemala and Panama (28). However, in a clinical trial in Colombia, itraconazole (oral dose 200 mg twice daily for 28 days) was ineffective against CL caused by *L. panamensis* (73).

Fluconazole has important properties including a longer half-life and increased concentrations in cutaneous tissues. In *L. major* infections, there was a good evidence of the benefit for the use of 200 mg oral fluconazole for 6 weeks (31). A study in Saudi Arabia showed cure rates 79% in CL caused by *L. major* after 6-weeks of 200 mg daily of oral fluconazole (74). An important advantage of
azoles is the ease of administration via the oral route. However, these drugs have many side effects such as the low cure rates, hepatotoxicity and gastrointestinal symptoms (72).

Figure 1.10. Chemical structure of some azoles(75)

1.4.2. Local therapy

1.4.2.1. Paromomycin

Paromomycin (PM) is an aminoglycoside antibiotic (Fig 1.11) and was identified as an anti-leishmanial drug in the 1960s. The sulphate salt of PM is given parenterally to treat VL, e.g. 11 mg/kg/day intramuscularly for 21 days. A topical formulation of paromomycin sulphate 15% plus 12% methyl-benzethonium chloride (MBCL) ointment has been used for LCL by applying twice daily for 20 days (28, 76). Topical 15% PM + 12% MBCL was active in BALB/c mice infected with New World species (L. mexicana) but did not show activity against L. panamensis and L. amazonensis (77).

Different formulations with a lower skin irritancy including one containing 15% paromomycin with 0.5% gentamicin gave cure rates of 81-82% for CL caused by L. major and 80% in Panama for CL caused by L. braziliensis and L.
*panamensis* in phase 3 studies. However, these results compared with a placebo cure rate of 58%, and almost no difference between formulations combining paromomycin and gentamicin or paromomycin alone (4). Paromomycin has low cure rates against certain *Leishmania* species and in many cases, relapse can be found during the first year (30, 46).

The exact mechanism of PM against *Leishmania* parasites is not fully known, studies suggest PM inhibits protein synthesis by blocking the dissociation of ribosomal subunits (78), others suggest that PM alters leishmanial lipid metabolism leading to the arresting of growth (79).

![Figure 1.11. Chemical structure of paromomycin(71)](image)

### 1.4.2.2. Physical treatments

Physical methods such as, localised heat or cryotherapy have been used in the treatment of CL. Localized heat is performed by using a device (e.g. Thermomed®) which provides a focused heat on the lesion (50°C for 30 seconds once per week for one month) and this method demonstrates about 69% overall efficacy against CL (1, 80). Cryotherapy is the use of liquid nitrogen to freeze lesions, repeated on three separate days. The efficacy of this procedure is about 57% against *L. major*. The benefits of localised heat or
cryotherapy methods are the ease of use and the safety. The problems with these methods include the low cure rates, the need for expensive equipment, and availability of electricity in rural areas (1). A comparison of the effects of three different therapies for CL was done in Iran: intralesional meglumine antimoniate or cryotherapy (liquid nitrogen (−195 °C)) or a combination of these two methods. They found that combining both MA and cryotherapy gave a significant higher activity than the two monotherapies (81, 82).

1.4.3. Immunomodulatory treatment

The immune response plays an important role in the control of CL - cure depends upon the activation of macrophages to produce toxic nitrogen and oxygen metabolites to kill the intracellular amastigotes (83). Consequently, immunomodulators for CL and VL have been studied widely for many years either alone or in combination with other drugs (84). For example, 11532 Venezuelan patients with American cutaneous leishmaniasis were treated with a combination of an immunomodulator (heat killed Leishmania promastigotes and bacille Calmette-Guérin (BCG)) and chemotherapy (meglumine antimoniate). Cure rates of 91.2 to 98.7% were achieved (85).

Examples of other clinically used immunomodulators include:
- Imiquimod: an antiviral compound [1-(2- methylpropyl)-1H-imidazo (4, 5-c) quinolin-4-amine] used topically for the treatment of genital warts, caused by the human papillomavirus, via the stimulation of localised immune response. Macrophages are activated to produce cytokines and nitric oxide at the site of application (76). Many studies have shown that imiquimod has anti-leishmanial activity. A randomized, double-blind clinical trial in Peru showed that patients with CL treated with 5% Imiquimod cream in combination with meglumine antimonate therapy showed faster lesion cure in comparison with those received meglumine antimonate with placebo vehicle cream therapy
- Pentoxifylline: Pentoxifylline is a methylxanthine derivative that inhibits TNF-α and decreases tissue inflammation. A clinical study showed that patients with CL caused by L. braziliensis who received a combination of
pentoxifylline plus SbV had higher cure rates than in those receiving antimony plus placebo (86, 87).

- A topical immunomodulator which is cytokine granulocyte-macrophage colony stimulating factor (GM-CSF) was found to accelerate the lesion healing in CL patients (88).

1.5. Challenges for CL treatment

CL is classified as a neglected tropical disease (NTD). NTDs have been described by the WHO as a varied group of diseases that have an impact on more than one billion people and dominate in 149 countries in tropical and subtropical conditions. These diseases are commonly associated with poverty and cause a huge economic and health burden in low- and middle-income countries (89, 90). Many factors form a challenge for CL treatments. CL happens in tropical areas with high temperatures, humidity and without cold chains and these conditions affect the stability of drug formulations for CL and even for other diseases (91). For example, AmBisome® requires a cold chain to protect its activity and a consistent supply of electricity is often difficult in rural regions. Moreover, some patients live in remote areas and are unable to access treatment easily. Availability of medicine(s) is also a challenge (92). Besides that, WHO estimated the cost of CL treatment to be between 12-40 USD$ per patient (28), this cost is prohibitive for many as the monthly income in many CL-endemic areas is only 7-17 USD$ /per person (28). Additionally, a delay between recognition of CL and starting the treatment increases the possibly of lesion progression to an ulcer with subsequent treatment complications and scarring (93, 94).

1.6. Assays to test the anti-leishmanial activity of drugs

The existing predictive models to study the anti-leishmanial activity of compounds are classified into *in vitro* and *in vivo* assays.
1.6.1. *In vitro* assays

These models are classified as either promastigote, axenic (extracellular) amastigote or intracellular amastigote assays. The advantages of using promastigote and axenic (extracellular) amastigotes are higher-throughput, cheaper, quicker and more straightforward screening. However, the drawbacks encompass that promastigotes are significantly different from intracellular amastigotes (target form in mammalians resides within the macrophages of the dermal skin layer) in terms of metabolism and ecology. Moreover, these promastigotes grow at 26°C and this could affect the anti-leishmanial action of drugs while *in vivo* temperature of 37 °C (34 °C skin temperature) (95, 96). The axenic amastigotes test is semi–predictive as it does not examine the penetration of the compound into the host cell and does not reflect the activity of the compound in the host environment and accordingly, is prone to false positive and negative results (95, 97).

On the other hand, the intracellular amastigote test (infected macrophages) is the gold standard model. In this model, macrophages can be derived from a range of sources, for example murine peritoneal macrophages (PEMs) or murine bone-marrow macrophages (BMMs), or chemically differentiated from human cancer cell lines (THP-1) (71). The activity of tested drugs is evaluated by exposing infected macrophages to particular concentrations of the drug for a specific period (such as 2, 3 or 5 days), and then stained with Giemsa after fixation with methanol. Activity is measured by either microscopical counting of number of amastigotes per macrophage or the percentage of infected macrophages (containing at least one parasite) (% infection). The selections of new compounds as anti-leishmanial depend on the 50 % and 90 % effective concentrations (EC\textsubscript{50}, EC\textsubscript{90}) after comparison with an untreated control and a positive control drug (95, 97).

In addition, there are more *in vitro* methods used to test the anti-leishmanial activity of drugs are summarised in Table 1.3 with positive and negative points for each assay.
Table 1.3. *In vitro* screening models with positive and negative drawbacks copied from (97).

<table>
<thead>
<tr>
<th><em>In vitro</em> models</th>
<th>Merits</th>
<th>Demerits</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Promastigote</strong></td>
<td>Rapid method and very little amount of test compounds are required for screening.</td>
<td>Not relevant life cycle stage for mammalian leishmanial infection. Data correlation with amastigote screening is unreliable.</td>
</tr>
<tr>
<td><strong>Axenic amastigotes</strong></td>
<td>Test is direct on relevant stage of the parasite.</td>
<td>The assay is semi-predictive.</td>
</tr>
<tr>
<td></td>
<td>This stage is as easy to manipulate as the promastigotes.</td>
<td>It neither tests for penetration of compound into host cell nor for activity in peculiar environment of the macrophage phagolysosome.</td>
</tr>
<tr>
<td></td>
<td>Quantification of drug activity is simple and often inexpensive.</td>
<td>Different metabolic processes than intracellular amastigotes. Screening of axenic amastigotes from clinical isolates is not possible as they require time to get adapted in the cultures.</td>
</tr>
<tr>
<td><strong>Intracellular amastigotes</strong></td>
<td>Effective screening method.</td>
<td>Labour intensive and subjective.</td>
</tr>
<tr>
<td></td>
<td>Mimic the environment encountered by the target cell.</td>
<td>Provide an approximation of the macrophages that are counted. Rendered difficult the screening of several drugs at a time and incompatible with HTS.</td>
</tr>
<tr>
<td></td>
<td>Shows the effect of drug mediated toxicity on host cell.</td>
<td></td>
</tr>
<tr>
<td><strong>Reporter gene assays:</strong> (GFP) Green fluorescent protein</td>
<td>Simple</td>
<td>Fluorescence intensity in parasites decreased with time in the absence of genetin sulphate (antibiotic G 418), thereby necessitating its regular addition.</td>
</tr>
</tbody>
</table>
Easier kinetic monitoring. Application for drug-drug screening is limited to promastigotes. Low cost and enhanced biosafety.

**β-galactosidase**

Colorimetric detection can be performed. Large size (the monomer is 116 kDa). Low sensibility. Endogenous expression of β-galactosidase by some mammalian cell types including macrophages.

**β-lactamase**


**Luciferase**

The method is rapid. Very sensitive. Highly reproducible. Does not require any very specialized instrument or training. Detection of only live, metabolically active cells by biphotonic imaging. Absence of background activity in the host cell. Compatible with HTS. Luminescent read out transient. Mixing of the samples and reagents needs to be timed with entering samples into the luminometer.

HTS, high throughput screening

These *in vitro* screening models have a major drawback related to their lack of biological relevance - they involve traditional cell culture methods (static and two-dimensional culture systems). Static cell culture systems that use the micro well plates are widely used. However, cells in human and animal tissues are sensitive to their microenvironment and face different mechanical stimulants due to interstitial flow and nutrient diffusion. Static cell culture systems are unable to provide these mechanical and physical factors arguably significantly limiting the cellular response *in vitro* (98, 99). Dynamic culture systems have the potential to overcome these limitations and better mimic the *in vivo* situation for drug discovery process (100).
1.6.2. *In vivo* assays

Different animal models (summarized in Table 1.4) are used to evaluate the effectiveness of drugs against leishmaniasis. These models imitate some of the pathological features and immunological responses shown in humans when exposed to *Leishmania* infections. *In vivo* assays allow the determination of drug activity in association with drug administration, excretion, and distribution. They can identify adverse events (toxic side effects) resulting from a particular treatment (101, 102). Murine models are widely used to evaluate the effectiveness of new drugs against leishmaniasis and to study the pathogenesis of this disease. *L. major*-BALB/c is the most used, with high reproducibility, and relatively fast progress of skin lesions (within 3 weeks). In this model, only potent drugs show anti-leishmanial efficacy as self-healing of CL is rare due to the immunological incapability of BALB/c mice (97, 101, 102).

The anti-leishmanial activity of compounds in the animal model is typically determined by a reduction of lesion size compared to untreated controls. However, inflammation plays a key role in lesion size. Therefore, size alone does not accurately reflect the anti-leishmanial activity. An additional indicator of therapeutic effect, e.g. determination of parasite burden should be considered. This can be achieved by different assays such as quantitative polymerase chain reaction (qPCR) or *in vivo* imaging (semi-quantitative) of bioluminescent parasites (97, 101, 102). A Therapeutic Index (TI) is often used to express the window between the required effective dose and the toxic/lethal doses of the drug (ED$_{50}$/LD$_{50}$) (95, 97).

<table>
<thead>
<tr>
<th>Animal Species</th>
<th>Examples</th>
<th>Main strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mice</td>
<td>BALB/c</td>
<td>Immunology, Vaccines, Chemotherapy</td>
</tr>
<tr>
<td></td>
<td>C57BL/6</td>
<td>Negative model-Immunology, Vaccines, Chemotherapy</td>
</tr>
<tr>
<td></td>
<td>Transgenic mice</td>
<td>Immunology</td>
</tr>
<tr>
<td>Hamster</td>
<td>Syrian golden hamster</td>
<td>Pathology, Chemotherapy</td>
</tr>
</tbody>
</table>
### 1.7. Drug development for CL: from pipeline to patients

The currently available drugs for CL have some drawbacks such as, low cure rates, toxicity, and high costs. These limitations clearly highlight the need for short, safe, efficacious, affordable and field-adapted treatments against *Leishmania* parasites (84). The process of developing and discovering new drugs is long, slow, expensive and challenging (Fig 1.12). For example, screening 100,000 compounds can lead to just one compound from a research and development (R&D) pipeline to a marketed drug and may take more than 10 years with an estimated expenditure around 2.6 billion USD$ (87). NTDs are generally considered commercially unattractive for Pharma research and development (R&D) (103) (81). In the last 40 years, only few new drugs have been developed for NTDs despite the great knowledge in the field of NTDs (88). NTDs usually involve populations with low purchasing power in low income countries - not a monetary incentive for the private sector to develop new therapies. Publication is typically the end of the line for NTD R&D or at other stages in the drug development pipeline (Fig 1.13) (88). Nonetheless, there has been a considerable moving forward in VL treatment and in the developing of new drugs of this disease. In contrast, no great attentions have been paid for CL drugs development (more details later) (90).

One of the strategies to overcome the high cost and long time lines of developing a new drug or chemical entity is “drug repurposing” (90). In this strategy, a known drug for a specific target is tested against different diseases. The drug has already been tested for toxicity, and other pharmacokinetic and pharmacodynamic studies have already been carried out, all in relation to its original indication. This can save time and money. As a result, getting a drug
to market can take less time. For instance, three drugs have been repurposed and used clinically for leishmaniasis: AmB was originally developed for fungal infections, paromomycin was primarily used for amoeba infections as an oral drug, and miltefosine was developed as an anti-cancer treatment (104, 105). Many researchers, worldwide, have identified a large number of compounds that show anti-leishmanial activity, either via re-purposing (tamoxifen, nelfinavir, imipramine, delamanid, fexinidazole) or isolating new chemical entities (NCEs) from natural sources (quinones, pyrimidines) (90). Another strategy is optimising the drug by reformulation of the active ingredient in the current drug or by using drug delivery systems for currently active drugs (more details later) (104, 105, 106).

Figure 1.12. The process of drug discovery and drug development. a) drug discovery stages b) drug development (107)
Figure 1.13. The Drug Development Pipeline - potential drugs for NTDs are frequently stuck in the early stage of development as a result of pipeline gaps (108)

1.8. New CL drugs

Despite limited resources, there have been recent developments in the NTD drug development arena. Some non-profit organisations such as TDR, have been involved in the development of 12 new drugs for NTDs. Another collaborative, patients' needs-driven, non-profit drug R&D organisation developing new treatments for neglected diseases, is the Drugs for Neglected Diseases initiative (DNDi). The partners and collaborators, which include academic institutions, Pharma and other non-governmental organizations (NGOs) work together, using their knowledge of NTDs, clinical trials and the capability of manufacturing drugs. DNDi facilitate these complex partnerships to enable rapid development and deployment to patients (80, 108, 109).

Drug discovery for CL is especially complex as CL is not a single disease with a single etiological agent, by contrast it is caused by more than 15 different Leishmania species with known variability in susceptibility to drugs. Identifying

---

3 the Special Programme for Research and Training in Tropical Diseases, supported by the WHO, the United Nations Children’s Fund (UNICEF) and the United Nations Development Programme (UNDP).
a drug that shows activity against all forms and species of CL is a tall order (109, 110).

Some potential new treatment candidates in the DNDi pipeline for NTDs are shown in Fig 1.14. Those for CL are:

- CPG-D35 oligonucleotides - synthetic DNA molecules working as an immunomodulator (by activating skin immune cells) for use as a monotherapy or in combination (111).
- DNDi-6148 and DNDi-0690 from oxaborole and nitroimidazole classes respectively, are undergoing Phase I clinical studies after completing the pre-clinical development as drugs for VL and CL.
- A combination of miltefosine (orally for three weeks) with thermotherapy (50°C for 30 seconds once during the treatment course) is in Phase II clinical trial (111).

**Figure 1.14.** New treatment candidates for leishmaniasis (111)
1.9. Drug delivery systems for leishmaniasis

Great attention has been paid in the field of drug development to drug delivery systems (DDs). These systems are used to increase efficacy and decrease toxicity of already active drugs by controlling their pharmacokinetic properties, such as absorption, distribution, metabolism, and excretion and also by enabling drug targeting to infected tissues/cells (112, 113). The accomplishment of CL treatment depends on the physical accessibility of the drug delivery systems to the infected macrophages in the dermis. The DDs should be able to penetrate the infected macrophages and by the time, the drug reaching the infection site of CL; the drug must cross the infected macrophage membrane, then permeate through the membrane of the PV and at the end crossing the plasma membrane of the *Leishmania* parasite, releasing the drug inside the PVs, leading to a local high concentration of the drug (Fig 1.15) (67, 113).

**Figure 1.15.** Route a drug must take to access intracellular *Leishmania* amastigotes within macrophages (A) (114) and DDs to intracellular *Leishmania* amastigotes(B) (115). A drug-loaded lipid or polymeric nanoparticle (yellow) is reaching the infected macrophage (1). This DDs is successfully phagocytosed by this infected macrophage (2). The DDs-including endolysosome (or phagolysosome) fuses with the amastigote-including parasitophorous vacuole (3). Drug is released from phagocytized DDs to kill *Leishmania* amastigotes (4).
Another promising approach for leishmaniasis treatment is related to the use of anti-leishmanial drugs with nanocarriers (DDs). There are different classes of nanocarriers e.g. particles, liposomes, emulsions etc, and many drug delivery systems have been evaluated in CL treatment (summarised in Table 1.5), some with promising results. Liposomal nanocarriers are the most studied over the past 30 years (67, 112, 113).

Using these nanocarriers DDs for CL therapies may facilitate drug solubility, reduce the toxicity, improve efficacy, modulate drug pharmacokinetics, permit sustainable drug release at the site of infection and protect the drug from degradation (113). An additional potential benefit is reducing the number of doses and the total dose, which would be significant for a drug like amphotericin B. The physicochemical properties (size, charge, morphology) and the rate of drug release from these DDs will significantly affect drug release into surrounding tissues, both before and after reaching cells at the uptake site (114). Generally, the efficacy of these DDs against CL depends on the administration route (Fig 1.16), for example in 1997, the intravenous administration once a day on six alternate days of AmBisome® (liposomal AmB) in a BALB/c L. major model of CL produced a dose-response effect, while the treatment was ineffective by the subcutaneous route (51). Liposomal SbV by the intravenous route is effective (116).

<table>
<thead>
<tr>
<th>Routes</th>
<th>Drug</th>
<th>Nanosystem</th>
<th>Parasite</th>
<th>Efficacy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parenteral</td>
<td>Amphotericin B</td>
<td>Chitosan and chondroitin sulphate nanoparticles</td>
<td>L. amazonensis</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Amphotericin B</td>
<td>Poloxamer 407-micelles</td>
<td>L. amazonensis</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Amphotericin B</td>
<td>PLGA-DMSA nanoparticles</td>
<td>L. amazonensis</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Amphotericin B</td>
<td>Liposome</td>
<td>L. tropica</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Amphotericin B</td>
<td>Liposome (AmBisome®)</td>
<td>L. major</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Amphotericin B</td>
<td>DSHemsPC-liposome</td>
<td>L. major</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Table 1.5. Experimental studies using nanosystems for CL treatment copied from (115).
<table>
<thead>
<tr>
<th>Compound</th>
<th>Delivery System</th>
<th>Species</th>
<th>Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amphotericin B</td>
<td>Nanodisks</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>PADRE-derivatized dendrimer complexed with liposome</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Chalcone DMC</td>
<td>PLA Nanoparticles</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Nanoselenium</td>
<td>Inorganic nanoparticle</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Paromomycin</td>
<td>Solid lipid nanoparticle</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Paromomycin</td>
<td>Solid lipid nanoparticle</td>
<td><em>L. tropica</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Pentamidine</td>
<td>Methacrylate nanoparticles</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Pentavalent antimonial</td>
<td>Nanohybrid hydrosols</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Sodium stibogluconate</td>
<td>Liposome</td>
<td><em>L. mexicana / L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Quercetin</td>
<td>Lipid-core nanocapsules</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Meglumine antimoniate</td>
<td>Beta-cyclodextrin</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Meglumine antimoniate</td>
<td>Polarity-sensitive nanocarrier</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Liposome</td>
<td><em>L. mexicana</em></td>
<td>NO</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Gamma-cyclodextrin</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Chalcone CH8</td>
<td>Liposome</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Paromomycin</td>
<td>Liposome</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Paromomycin</td>
<td>Liposome</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Meglumine antimoniate</td>
<td>Liposome</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Nano silver</td>
<td>Inorganic nanoparticles</td>
<td><em>L. major</em></td>
<td>No</td>
</tr>
<tr>
<td>Nano silver</td>
<td>Inorganic nanoparticles</td>
<td><em>L. major</em></td>
<td>No</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td>Liposome</td>
<td><em>L. major</em></td>
<td>No</td>
</tr>
<tr>
<td>Chalcone CH8</td>
<td>PLGA microparticles</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Nano silver</td>
<td>Inorganic nanoparticles</td>
<td><em>L. amazonensis</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Meglumine antimoniate</td>
<td>Liposome</td>
<td><em>L. major</em></td>
<td>No</td>
</tr>
<tr>
<td>Miltefosine</td>
<td>Liposome</td>
<td><em>L. major</em></td>
<td>Yes</td>
</tr>
<tr>
<td>Paromomycin</td>
<td>Liposome</td>
<td><em>L. major</em></td>
<td>No</td>
</tr>
<tr>
<td>Paromomycin</td>
<td>Solid lipid nanoparticle</td>
<td><em>L. tropica</em></td>
<td>Yes</td>
</tr>
</tbody>
</table>
Sodium stibogluconate | Liposome | L. mexicana/ L. major | Yes
--- | --- | --- | ---
Note: Chalcone DMC – 2',6'-dihydroxy-4'-methoxychalcone; Chalcone CH8 – 3-nitro-2'-hydro-4',6'-dimethoxychalcone; DMSA – dimercaptosuccinic acid; DSHemsPC – 1,2-distigmasterylhemi-succinoyl-sn-glycero-3-phosphocholine; PADRE – pan DR-binding epitope; PLA – poly(D,L-lactide); PLGA – poly(lactic-co-glycolic acid); UVB – ultraviolet B radiation.

Despite the promising effectivity against leishmaniasis and the safety profile of liposomal formulations, their high cost decreases their use in the leishmaniasis field. Subsequently, more attention has been paid recently to polymeric nanoparticles, Carvalho et al found that a nanoparticle delivery system (consisting of free deoxycholate AmB encapsulated in polylactic-co-glycolic acid (PLGA)) was more active in the treatment of experimental cutaneous leishmaniasis (L. amazonensis) in C57BL/6 mice than free drug (117). Kumar et al demonstrated that PLGA-PEG (poly(D,L-lactide–co--

**Figure 1.16.** Administration routes of DDs and anatomical barriers. A. Intravenous route. B. Subcutaneous, intramuscular and intraperitoneal route. C. Oral route. D. Topical route. E. CL causes regional inflammation is associated with leaky vasculature. In this situation, particles in blood circulation can permeate barrier to become close to the infected cells. F. Particles in blood circulation (67).
glycolide)--block--poly(ethylene glycol)) encapsulated amphotericin B nanoparticles were significantly more effective than free amphotericin B against *L. donovani* strain MHOM/IN/83/AG83 in both *in vitro* and *in vivo* (Female hamsters) studies (117). Similarly, Ahmed *et al* found that a noncovalent complex of amphotericin B (AmB) and poly (α-glutamic acid) (PGA) with a size of ~100 nm, to be significantly less toxic against KB-cells in comparison with free amphotericin B and amphotericin B deoxycholate (Fungizone™) whilst keeping the same anti-leishmanial activity against *L. major* (MHOM/SA/85/JISH118) or *L. donovani* (MHOM/ET/67/HU3) intracellular amastigotes (118). Unfortunately, most of these delivery systems required organic solvents or heat for preparation - using these solvents or temperatures is not ideal as they can influence the integrity of the polyenic substances used and besides increase the toxicity of the DDs (119). In contrast, ionotropic gelation is a widely used method for preparing polymer nanoparticles and this method does not require the use of organic solvents or heat (120). In this method, nanoparticles are prepared by the interaction between two oppositely charged groups (120). Some benefits of this method are the ease of preparation, aqueous environment, low toxicity and protection of the chemical structure of the encapsulated drug (120).

Recently, a nanoparticle delivery system for AmB has been developed using the ionotropic gelation method with chitosan as a positive molecule and chondroitin sulphate (glycosaminoglycans in the extracellular matrix of cartilage) as a negative one. These loaded nanoparticles were 10 times less toxic than unincorporated AmB against murine macrophages and showed *in vitro* anti-leishmanial activity against *L. infantum* and *L. amazonensis* promastigotes and amastigotes. The efficacy of these AmB-loaded nanoparticles against *L. amazonensis*-infected BALB/c mice have been evaluated and showed a significant reduction in parasite load at 1 mg/kg/day/intravenously for 10 days. These nanoparticles induced significantly higher levels of IFN-γ and IL-12 in the mice (121, 122).

Chitosan is a widely used compound in drug delivery systems because of its interesting structure - chitosan has a cationic feature, is soluble in acidic media and has mucoadhesive properties (123). Chitosan is reported to have
immune-stimulatory effects which include inducing NO and ROS production (124, 125, 126) and antimicrobial and anti-leishmanial activity (127, 128, 129). Chitosan can be used in various formulations in the drug delivery systems and these forms are summarised in Table 1.6 with some example of associated loaded drugs (130, 131, 132) and molecules (133, 134). Chitosan nanoparticles are biocompatible and biodegradable, important properties for drug safety and controlled release, and are increasingly being considered for a variety of biomedical applications, e.g. wound healing (130, 135). Chitosan nanoparticles can be prepared in different sizes and different charges, and are suitable for different routes of administrations (123) (more details later).

Table 1.6. Chitosan-based drug delivery systems.

<table>
<thead>
<tr>
<th>DDs</th>
<th>Method of preparation</th>
<th>Drug</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tablets</td>
<td>Matrix</td>
<td>theophylline, mesalamine, glipizide and diclofenac sodium</td>
<td>(120, 136, 137, 138, 139)</td>
</tr>
<tr>
<td>Capsules</td>
<td>Capsule shell</td>
<td>insulin</td>
<td>(140)</td>
</tr>
<tr>
<td>Microspheres/</td>
<td>Emulsion cross-linking, Coacervation/precipitation, Spray-drying</td>
<td>clarithromycin, propranolol HCl, gentamicin sulphate, famotidine and cimetidine</td>
<td>(141, 142, 143, 144, 145)</td>
</tr>
<tr>
<td>Microparticles</td>
<td>Emulsion-droplet coalescence, Ionotropic gelation, Reverse micellar method, Coacervation/precipitation</td>
<td>doxorubicin, cyclosporin A, gadopentetic acid, levofoxacin, amphotericin B and miltefosine</td>
<td>(120, 146, 147, 148, 149, 150)</td>
</tr>
<tr>
<td>Nanoparticles</td>
<td>Coacervation/ precipitation</td>
<td>insulin</td>
<td>(151)</td>
</tr>
<tr>
<td>Beads</td>
<td>Solution casting</td>
<td>ofloxacin and paclitaxel</td>
<td>(152, 153)</td>
</tr>
<tr>
<td>Films</td>
<td>Cross-linking</td>
<td>5-Fluorouracil</td>
<td>(154)</td>
</tr>
</tbody>
</table>

1.10. Nanoparticles and their interaction with skin lesions

The ease of administration and reduced systemic side effects of topical formulations prioritise them over systemic therapy for uncomplicated CL (30). Topical formulations for CL encounter different barriers in the skin and some
are shown in Fig 1.17. Nanoparticle carriers have been widely used in topical formulations to treat skin disease such as fungal infections, psoriasis and, for cosmetic purposes (155). The penetration of nanoparticles through the skin can occur by one of these three routes: intercellularly in between corneocytes, intracellularly through corneocytes or via dermal structures like the hair follicles (Fig 1.18) (155).

**Figure 1.17.** Factors to be considered in topical delivery (30).
Figure 1.18. Pathways of skin nanoparticles penetration. 1) via hair follicles, 2) intracellularly through corneocytes and 3) intercellularly around corneocytes (155).

The biological effects (toxicity, immune interactions), depth and mechanism of skin penetration of the nanoparticles are based on their structure and properties such as size, zeta potential, aggregation, solubility in the skin, skin lipid composition and drug release from these nanoparticles. The condition of the skin, healthy or otherwise, influences nanoparticle permeation. Current dogma avers that biodegradable polymeric nanoparticles accumulate in the hair follicle and on the surface (stratum corneum) of healthy skin (156). In CL, drug permeation may be influenced by the morphology of ulcers, such as necrotic centres and high borders to the lesion. CL causes an inflammation response involving higher permeability and vasodilatation of blood vessels of the dermis at the infection site, and moreover several types of immune cells, including macrophages, are infiltrated to the infection site and this could promote the permeation of the topical drug through the damaged epidermis (157), see Fig 1.19.

Despite this ease of drug permeation caused by the local inflammation in CL, the location of Leishmania parasites in the dermis, instead of the superficial portions of the epidermis where most fungi typically reside, forms a major impediment to the permeation of topical drugs (157).
The most favourable drug formulation for topical treatment of CL is the one in which the drug has a high anti-leishmanial activity and can permeate through the skin to reach *Leishmania* parasites located in the dermis, in high enough concentrations to act (115).

Chitosan nanoparticles have been reported to (i) improve the drug permeation into skin in comparison to other vehicles and, (ii) provide a sustain drug release from these nanoparticles. Moreover, chitosan has wound healing effects, mucosal adhesion properties and antimicrobial activity (158). Many clinical studies demonstrated the positive effects of using chitosan as wound dressing in accelerating the rapid wound re-epithelialisation and the regeneration of the granular layer, haemostasis in patients undergoing plastic surgery (159), skin grafting (160, 161) and endoscopic sinus surgery (162). Chitosan nanoparticles have been repeatedly administered for topical skin delivery; retinol encapsulated in chitosan-TPP nanoparticles showed less toxicity than unloaded retinol and potential activity for acne and anti-wrinkle treatment (163). Acyclovir (an antiviral medication) loaded chitosan-TPP nanoparticles caused an increase in the drug stability and stimulated drug penetration through porcine skin (164). Chitosan–dextran sulphate nanoparticles also showed mucoadhesive properties and potent activity in the treatment of ocular surface infections (165). Therefore, these encouraging properties make chitosan a suitable candidate for further studies in terms of topical treatment of CL.
Figure 1.19. Interaction of nanoparticles with lesions of CL (156). A high parasitic load, low lymphocytes infiltrate and small tissue necrosis is observed in nodular lesions. On the contrast, the parasite load is low with higher lymphocytes infiltrate and tissue injury. While in the necrotic tissue the neutrophils are gathered, infected macrophages and lymphocytes are situated in the border of the ulcers. In nodular lesions, nanoparticles are applied to the epidermis. Based on their physiochemical properties they either (i) stay on the surface, (ii) penetrate the epidermis (small, deformable NPs), and/or(iii) fuse with the epidermis. After which a drug release will occur and diffuse to the dermis to meet the infected macrophage and then this drug being eliminated by lymphatic and blood clearance. The time of retention in the dermis is crucial for the treatment efficacy. However, these drug carriers encounter fibrotic and necrotic dermal tissue with infiltration of neutrophils in the centre of the ulcer. The chance to reach infected macrophages is higher in ulcerative lesions. Nanoparticles can promote stress and proinflammatory signalling that enable the elimination of parasites and accelerate the wound healing and according to the nanoparticles physicochemical properties this can be happened by either direct influence on macrophages or indirectly by their effects in keratinocytes and neutrophils. The design of these nanoparticles should take into consideration the maximal eradication of parasites and lowering the tissue injury.

1.10.1. Mathematical models of skin permeability

Small uncharged drug molecules mainly permeate through skin by passive diffusion in which, move from an area of higher concentration to an area of lower concentration (Fig 1.20 ) (166).

Infinite dose permeation experiment is usually used to examine the permeation behaviour of a compound or to determine the influences of penetration enhancer on percutaneous permeation. Infinite dose is applied to
keep a steady rate of absorption of the compound through the skin, that is called the steady state flux and will produce a cumulative permeation amount of compound permeated through a unit area of membrane over time (166, 167).

**Figure 1.20.** The passive diffusion of drug through a membrane (168)

Fick’s first law can give the main equation (Equation 1) to identify the skin permeation after exposing the skin membrane to the diffusing molecules on one side of the skin when steady state conditions have been reached. This equation states the amount of molecule (Q) permeating the skin membrane of area (A) over a period of time (T) with the steady concentration gradient across the skin membrane, ΔCs (in mol/cm$^3$) and it relates the diffusion coefficient in the skin membrane, D (diffusion coefficient) (in cm$^2$/s), and the path length, h(in cm) (166, 169).

Equation (1) $Q = \frac{DAT\Delta Cs}{h}$, $\Delta Cs = C_0 - C_i$ (C$^0$ represents the concentration of compound applied to the skin surface and Ci stands for the concentration of compound inside the skin)

This equation assumes that the skin barrier (stratum corneum (SC)) is acting as a pseudo homogenous membrane and no changes happen in SC properties with time and position.

Fick’s first law, applied when steady state conditions have been reached, states that the rate of transfer of the diffusing molecules per unit area is proportional to the concentration gradient measured across the membrane (Equation 2). Therefore, equation 2 is indicated as the flux of the permeant per unit area (in mol/(cm$^2\cdot$s)) (166, 169),
Equation (2) \( J = \frac{Q}{AT} = \frac{K_D \Delta C_s}{h} \), \( J \) is the flux of the permeant per unit area (in mol/(cm\(^2\)·s))

As in most practical situations \( C_0 \gg \gg C_i \) therefore equation 2 becomes:

Equation (3) \( J = \frac{Q}{AT} = \frac{K_D C_0}{h} \)

Additionally, the permeability coefficient (\( k_p \)) is described as the flux of the permeant per unit area normalised by the concentration gradient and by characterising the skin as a single pseudo-homogenous membrane therefore \( k_p \) (in cm/s) is identified as

Equation (4) \( k_p = \frac{K_D}{h} \), \( K \) is the stratum corneum-formulation partition coefficient

Accordingly, from both equations 3 and 4, the flux of the permeant per unit area (in mol/(cm\(^2\)·s)) is:

Equation (5) \( J = K_p C_0 \) (166, 169)

Several assumptions should be taken into consideration before applying these equations to skin permeation into the experimental design including:

1. The stratum corneum forms the rate-limiting barrier
2. The stratum corneum is considered isotropic and its natures is not altered by the application of the vehicle of the drug formulation
3. The drug diffusion is not based on time, concentration or distance
4. The diffusing compound dissolves in the stratum corneum

However, in the clinical situations, patients mostly apply finite dose of the formulation. The amount of permeated compound through stratum corneum will accomplish a peak and stay constant (equation 6) and the diffusion is determined as bellow:

Equation (6) \( \frac{\partial c}{\partial t} = D \frac{\partial^2 c}{\partial x^2} \), \( c \) is the concentration of the permeating molecule at time \( t \) at depth \( x \) within the skin. This equation is applied just by presuming a unidirectional diffusion through an isotropic membrane (166, 167, 169).

1.11. Pharmacokinetics of ant-leishmanial drugs

Pharmacodynamic (PD) refers to the link between drug concentration and the influences on the biological system and illustrates how the drug influences the
parasite and can be determined in regards with effectiveness ($\text{EC}_{50}$, $\text{EC}_{90}$), potency (maximal effect) and the kill rate (time-dependence of the effect) (70, 71). While pharmacokinetic (PK) refers to the study of time course of the drug absorption, distribution, metabolism and excretion (ADME). The basic PK parameters are summarised in Table 1.7. Some concepts that affect importantly PK of CL drugs encompass: i) the target site of CL drug as *Leishmania* parasites survive and multiply in the macrophages of the dermis of the skin lesion and ii) the route of drug administration, for example, a topical drug should have the ability to penetrate through the stratum corneum of the epidermis and to retain in the dermis of the lesion. Moreover, iii) the metabolism of the parasite or the host (in macrophages, skin and liver) can activate or inactivate the drug. Drugs are divided into three groups according to PK/PD profile: 1- concentration-dependent antimicrobial effect -, 2- time-dependent antimicrobial effect or 3- dependent on both time and concentration (70, 71).

**Table 1.7. Basic PK parameters copied from (71)**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Description</th>
<th>Unit (example)</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose</td>
<td>D</td>
<td>the dose of drug administered</td>
<td>Mg</td>
<td>Design parameter</td>
</tr>
<tr>
<td>Dose interval</td>
<td>T</td>
<td>once per day (QD) twice per day (BD) trice per day (TID)</td>
<td>Per hour, per day</td>
<td>Design parameter</td>
</tr>
<tr>
<td>$C_{\text{max}}$</td>
<td>$C_{\text{max}}$</td>
<td>the maximal concentration in a specific matrix (usually in plasma, but can be in any part of the body) after drug administration</td>
<td>$\mu$g/ml</td>
<td>Direct measurement</td>
</tr>
<tr>
<td>$t_{\text{max}}$</td>
<td>$t_{\text{max}}$</td>
<td>the time corresponding to $C_{\text{max}}$</td>
<td>Hours</td>
<td>Direct measurement</td>
</tr>
<tr>
<td>Volume of distribution</td>
<td>$V_d$</td>
<td>the apparent volume in which a drug is distributed. Relates drug concentration to the amount of drug in the body and can give information about tissue distribution</td>
<td>Litre</td>
<td>$=D/C_0$</td>
</tr>
<tr>
<td>Elimination rate constant</td>
<td>$K_e$</td>
<td>the rate at which a drug is removed from the system</td>
<td>Per hour</td>
<td>$=\frac{\text{Cl}}{V_d}$ = $\ln (2)/T_{1/2}$</td>
</tr>
</tbody>
</table>
Regarding the pharmacokinetic of leishmaniasis drugs, pentavalent antimony has a long terminal half-life because of the intracellular conversion of SbV to SbIII which forms with the quick renal excretion the main characterisations of this drug pharmacokinetics. Miltefosine pharmacokinetics are characterised mainly by the accumulation in peripheral blood mononuclear cells (PBMCs) and long terminal half-life (70). However, paromomycin is characterised by the fastest excretion by the kidneys from the body in comparison with other leishmaniasis drugs. On the other hand, AmB pharmacokinetics have not been evaluated widely in leishmaniasis. It has been reported that the renal and faecal excretion of liposomal AmB (AmBisome®) is much slower than AmB deoxycholate (Fungizone) excretion which leads to higher exposure (70, 71). Wijnant et al reported that liposomal AmB (AmBisome®) caused a higher plasma peak and systemic exposure compared with AmB deoxycholate (Fungizone, after a single dose of 1 mg/Kg/ i.v. in L. major-infected mice) (170) and Table 1.8 summarizes the PK of leishmaniasis drugs including pentavalent antimonial, paromomycin, miltefosine, Fungizone and AmBisome® in clinical and mouse model studies. The application of PK and PD comprehension and understanding the relation between PK and PD produce a fundamental base for detecting the optimal dosage and effective therapeutic management of drugs for CL treatment and will be helpful in antileishmanial drugs combination to increase in an attempt to improve drug efficacy and decrease the duration of treatment (70, 170)
Table 1.8. Pharmacokinetic profile of leishmaniasis drugs (70, 170)

<table>
<thead>
<tr>
<th>Patients</th>
<th>Weight (kg)</th>
<th>Daily dose</th>
<th>Sampling day</th>
<th>C&lt;sub&gt;max&lt;/sub&gt; (µg/ml)</th>
<th>C&lt;sub&gt;trough&lt;/sub&gt; (µg/ml)</th>
<th>t&lt;sub&gt;max&lt;/sub&gt; (h)</th>
<th>k&lt;sub&gt;a&lt;/sub&gt; (day&lt;sup&gt;-1&lt;/sup&gt;)</th>
<th>V&lt;sub&gt;d&lt;/sub&gt;/F (L)</th>
<th>CL&lt;sub&gt;r&lt;/sub&gt;/F (L/h)</th>
<th>AUC (mg.h/L)</th>
<th>t&lt;sub&gt;1/2&lt;/sub&gt; (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SbV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Humans</td>
<td>Adults:</td>
<td>62 (56–120)</td>
<td>20 mg/kg/days, 20 days (IM)</td>
<td>Day 19</td>
<td>38.8 ± 2.1</td>
<td>0.198 ± 0.023</td>
<td>1.0 (1.0–2.0)</td>
<td>NA</td>
<td>0.30 ± 0.01&lt;sup&gt;b,c&lt;/sup&gt;</td>
<td>0.106 ± 0.006&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AUC&lt;sub&gt;24&lt;/sub&gt;: 190 ± 10</td>
</tr>
<tr>
<td></td>
<td>Children:</td>
<td>15 (13–18)</td>
<td>20 mg/kg/day, 20 days (IM)</td>
<td>Day 19</td>
<td>32.7 ± 0.9</td>
<td>0.113 ± 0.015</td>
<td>0.875 (0.5–1.5)</td>
<td>NA</td>
<td>0.39 ± 0.03&lt;sup&gt;b,c&lt;/sup&gt;</td>
<td>0.185 ± 0.013&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AUC&lt;sub&gt;24&lt;/sub&gt;: 111 ± 7</td>
</tr>
<tr>
<td>Paromomycin</td>
<td></td>
<td>35.5±11 .8&lt;sup&gt;a&lt;/sup&gt;</td>
<td>15 mg/kg (11 mg/kg base), 21 days (IM)</td>
<td>Day 1</td>
<td>20.5 ± 7.01</td>
<td>4.53 ± 6.71</td>
<td>NA</td>
<td>2.11 (7.68%)&lt;sup&gt;e&lt;/sup&gt;</td>
<td>15.3 (2.27%)&lt;sup&gt;e&lt;/sup&gt;</td>
<td>4.06 (3.05%)&lt;sup&gt;e&lt;/sup&gt;</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Data given as either mean ± standard deviation or median (range), unless indicated otherwise. AUC area under the concentration–time curve, AUC<sub>24</sub> AUC from time zero to 24 h, CL<sub>r</sub> clearance, C<sub>max</sub> peak plasma concentration, C<sub>trough</sub> trough plasma concentration 24 h after dose, F bioavailability, k<sub>a</sub> absorption rate constant, NA not available, t<sub>1/2</sub> plasma elimination half-life, t<sub>1/2,β</sub> elimination half-life, t<sub>1/2,24–48 h</sub> apparent half-life between 24 and 48 h (an approximation of the c-elimination half-life), t<sub>max</sub> time to C<sub>max</sub>, V<sub>d</sub> volume of distribution. <sup>b</sup> Per kg, <sup>c</sup> V<sub>β</sub> apparent volume of distribution during the b-elimination phase and <sup>e</sup> Mean (% standard error)
<table>
<thead>
<tr>
<th>Miltefosine</th>
<th>(CL) Humans Adults:</th>
<th>70.84 ± 11.73</th>
<th>2.11 ± 0.16 mg/kg/day, 28 days (Orally)</th>
<th>31.9 (17.2–42.4)</th>
<th>NA</th>
<th>NA</th>
<th>NA</th>
<th>NA</th>
<th>NA</th>
<th>NA</th>
<th>628 (213–861)</th>
<th>34.4 (9.5–46.15)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(CL) Humans Children:</td>
<td>26.22 ± 7.62</td>
<td>2.27 ± 0.16 mg/kg/day, 28 days (Orally)</td>
<td>22.7 (17.0–29.3)</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>448 (304–583)</td>
<td>37.1 (7.4–47.0)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Patients</th>
<th>Weight (g)</th>
<th>Daily dose</th>
<th>C&lt;sub&gt;max&lt;/sub&gt; (µg/ml)</th>
<th>AUC (h · µg/ml)</th>
<th>Clearance (ml/h/kg)</th>
<th>t&lt;sub&gt;1/2&lt;/sub&gt; (h)</th>
<th>V (ml/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fungizone</td>
<td>L. major-infected mice</td>
<td>20</td>
<td>a single i.v. 1-mg/kg dose</td>
<td>1</td>
<td>30.2</td>
<td>18.9</td>
<td>39.7</td>
</tr>
<tr>
<td>AmBisome</td>
<td>L. major-infected mice</td>
<td>20</td>
<td>a single i.v. 1-mg/kg dose</td>
<td>8.2</td>
<td>71</td>
<td>13.5</td>
<td>8.5</td>
</tr>
</tbody>
</table>

AUC area under the concentration–time curve, CLr clearance, C<sub>ss</sub> steady-state concentration, F bioavailability, k<sub>a</sub> absorption rate constant, NA not available, Q intercompartmental clearance, t<sub>max</sub> time to C<sub>max</sub> within one dosing interval, V volume of distribution, t<sub>p</sub> plasma elimination half-life, V central volume of distribution of the central compartment, V<sub>peripheral</sub> volume of distribution of the peripheral compartment.  

* Miltefosine accumulates during treatment and reaches C<sub>ss</sub> during the last week of treatment
* AUCD28 (AUC from start to end of treatment) unless indicated otherwise
* Unclear whether this is the mean C<sub>ss</sub> or the maximum C<sub>ss</sub>
* AUC from start of treatment to infinity (AUC∞)
1.12. Treatment challenges

As described in this chapter, currents treatments for CL have drawbacks, for instance high toxicity (nephrotoxicity, cardiotoxicity, hepatotoxicity etc), the high cost (such liposomal amphotericin B), instability, or sometimes low cure rates etc and this draws the attention to the need for new safe, effective, economically feasible new treatments for CL. Drug discovering and developing is a long, slow and very expensive process (71).

Drug delivery system is considered one of the effective strategies to overcome the cost and long process of developing new drugs in which DDs of already know active drugs and can be used to increase the activity of loaded drugs and to reduce their toxicity.

Chitosan has shown promising features in therapeutic delivery systems because of its biocompatibility, biodegradability, cationic structure, mucoadhesive properties, wound healing effects and the antimicrobial activity (130, 135). Therefore, chitosan has been chosen in this study as a carrier for AmB and the potential to treat CL, AmB is a high active drug against CL but its use has been decreased because of the toxicity and we aimed to improve the therapeutic window of AmB by using chitosan as a nanocarrier (171).

There are different methods for chitosan nanoparticles preparation and the ionotropic gelation method was chosen in this study as this method is a simple and quick method and can be used to synthesize spherical nanoparticles with different sizes and charges. Moreover, this method has been reported to produce very stable chitosan nanoparticles with sustainable drug release (171).

In literatures, chitosan nanoparticles showed encouraging properties as DDs for the treatment of leishmaniasis. However, there is just a study used the ionotrop gelation method and used chitosan nanoparticles with positive surface charge and with size of size= 136±11 nm and these studies need more detailed and controlled studies (83, 122).

Topical treatments have many advantages over systemic treatment for instance, (i) increasing the compliance with patients, (ii) affording a high local concentration of the drug at the lesion site and (iii) reduce the toxic effects of systemic drugs (158). Therefore, the possibility of use AmB loaded chitosan nanoparticles in this route could be of interest to benefit from the small size of the nanoparticles, mucoadhesive and wound healing effects of chitosan.
1.13. Aims and objectives

The overall aim of this project was to optimise an effective, safe and economically feasible nanoparticle delivery system of amphotericin B with the potential to treat cutaneous leishmaniasis.

- Hence, the aim of the first experimental chapter (chapter 2) was to:
  I. Determine the *in vitro* anti-leishmanial activity of chitosan and its derivatives against *L. major* and *L. mexicana* promastigotes and intracellular amastigotes at two different pH values of the culture medium (the medium pH 7.5 and at lower pH 6.5)
  II. To evaluate the *in vitro* role of chitosan in the activation of macrophage and M1 proinflammatory phenotype, via the measurement of NO, ROS and TNF-α production by host cells and by measuring parasite survival
  III. Investigate chitosan uptake by macrophages to explain activity against intracellular amastigotes.

- The purpose of the second experimental chapter (chapter 3) was to:
  I. Prepare two types of chitosan nanoparticles by using the inotropic gelation method; one with a positive surface charge using tripolyphosphate sodium (TPP) and the other with a negative surface charge, using dextran sulphate.
  II. Evaluate the characterisations of blank and amphotericin B loaded chitosan TPP or dextran sulphate nanoparticles by studying their physicochemical properties (size, morphology, zeta-potential and stability). The optimal conditions for nanoparticle preparation were chosen with regard to the smallest sizes and different charges.
  III. Determine amphotericin B loading and drug release from the amphotericin B loaded chitosan TPP or dextran sulphate nanoparticle

- The third experimental chapter aimed to (chapter 4):
  I. Evaluate the *in vitro* effectiveness of blank and amphotericin B loaded chitosan TPP or dextran sulphate nanoparticles against *L. major* and *L. mexicana* promastigotes and amastigotes after evaluating their haemolytic activity and cytotoxicity against KB-cells.
II. Evaluate the intravenous activities of blank and amphotericin B loaded chitosan TPP or dextran sulphate nanoparticle in vivo in BALB/c mice infected with *L. major*.

III. Measure the permeation of blank and amphotericin B loaded chitosan TPP or dextran sulphate nanoparticle through uninfected and *L. major* infected mouse BALB/c skin by *In vitro* Franz diffusion cell permeation studies.

- The fourth experimental chapter (chapter 5) aimed to:

I. Study the effects of media perfusion on the *in vitro* host cell phagocytosis and macropinocytosis.

II. Study the effects of the flow on the *in vitro* anti-leishmanial activity of chitosan solution and blank and amphotericin B loaded chitosan TPP or dextran sulphate nanoparticles
2. Activity of chitosan and its derivatives against *Leishmania major* and *mexicana* in vitro.

2.1. What is chitosan?

Chitosan is produced by the deacetylation of chitin (Fig 2.1). Chitin is the second most abundant natural polysaccharide and originates from the shells of crustaceans and the cell walls of fungi (172). Chitosan is a biodegradable, biocompatible and positively charged nontoxic mucoadhesive biopolymer (172, 173).

![Figure 2.1. Structure of chitin and chitosan and method of preparation chitosan from raw materials (172, 174).](image)

2.2. Chitosan solubility

Chitosan is insoluble at alkaline pH but is soluble in dilute acidic solvents like glacial acetic acid and acid solvents to form a cationic polymer (–NH\textsubscript{3}+ groups) (Fig 2.2). Chitosan in acidic media has a positive charge and the ability to form gels at low pH values because it is hydrophilic and can retain water in its structure (175). Chitosan pKa is approximately 6.3 and therefore, the approximate ionisation degree of chitosan is a 61% and 6% at pH 6.5 and 7.5 respectively.
2.3. Chitosan toxicity

Chitosan is widely considered as a non-toxic, biological polymer and has been approved by the FDA for use in wound dressings. Chitosan has been recognized by FDA as GRAS (Generally Recognized As Safe, GRAS Notice No. GRN 000073, EU 2011) (176) and approved for use in dietary applications in Italy and France (173). The lethal dose, 50% (LD$_{50}$) of chitosan for mice and rats are orally 16000 and 1500 mg/kg respectively (177).

2.4. Chitosan in wound healing

Chitosan has a similar chemical structure to hyaluronic acid and additionally enhances the functions of inflammatory cells such as polymorphonuclear leukocytes (PMN) (phagocytosis, production of osteopontin and leukotriene B4), macrophages (phagocytosis, production of interleukin (IL-1), transforming growth factor and platelet-derived growth factor) and fibroblasts (production of IL-8) (135, 175). Because of these properties, chitosan promotes skin wounds granulation with improving collagen production, angiogenesis and re-epithelialization of skin tissue. As a result, chitosan induces wound healing and produces less scarring. Also using chitosan hydrogel will provide a painless, antimicrobial and ideal dressing for wounds (135, 175).

2.5. Chitosan derivatives

The poor solubility of chitosan and the loss of the cationic nature charge at neutral and alkaline pH are two of the major obstacles to the usefulness consideration of chitosan as a useful antimicrobial material. Recently, the
chemical modification of chitosan to produce various derivatives to improve its solubility and widen its application, has gained a great attention. The presence of certain functional –NH₂ and –OH groups on chitosan structure provides the basis for many methods of structural modification. The most used method is graft copolymerization (an attractive technique to conjugate bioactive molecules on the surface). Derivation of chitosan can be used to improve its antimicrobial activity, solubility and the mucoadhesive properties (178, 179).

2.6. Antimicrobial activity of chitosan

Many reports mention that chitosan has a potential activity against microbes (detailed in Table 2.1) but the actual mechanism has not yet been fully elucidated [35]. Three mechanisms have been suggested to explain this activity. The first one is the interaction between the protonated NH₃⁺ groups of chitosan and the negative cell membrane of microbes. This interaction leads to change the permeability of the microbes' membrane wall, causes osmotic imbalances, and as a result prevents the growth of the microbes (174, 180). Another mechanism is that chitosan binds to microbial DNA and inhibits DNA transcription and mRNA synthesis (180, 181). The third mechanism is the chelation of metals and binding the basic nutrients for microbes. These three mechanisms lead to killing of the microbes (124). A fourth indirect mechanism of action may be related to the known pro-inflammatory activity effect of chitosan on macrophages. This involves stimulation of tumour necrosis factor (TNF-α), interleukin 6 (IL-6), nitric oxide (NO), reactive oxygen species (ROS) and interferon gamma (IFN-γ) which play critical roles in the proinflammatory response against intracellular microbes (by enhancing the production of microbicidal reactive nitrogen species) (125, 126, 182, 183, 184).
Table 2.1. The antimicrobial activities of chitosan and its derivatives (173, 185, 186, 187)

<table>
<thead>
<tr>
<th>Microbe</th>
<th>MIC of chitosan or derivative</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Escherichia coli</em></td>
<td>Chitosan 8 μg/ml; chitosan nanoparticles 0.0625 μg/ml; Cu loaded chitosan nanoparticles 0.0313 μg/ml</td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>Chitosan 0.0125%(w/v); chitosan-Zn complex 0.00625%(w/v); N, N-diethyl-N-methyl-chitosan 32 μg/ml</td>
</tr>
<tr>
<td><em>Proteus mirabilis</em></td>
<td>Chitosan 0.025%(w/v); chitosan-Zn complex 0.00625%(w/v)</td>
</tr>
<tr>
<td><em>Salmonella enteritidis</em></td>
<td>Chitosan 0.05%(w/v); chitosan-Zn complex 0.00625%(w/v)</td>
</tr>
<tr>
<td><em>Enterococcus faecalis</em></td>
<td>Chitosan 8 μg/ml; chitosan-Zn complex 0.125; N, N-diethyl-N-methyl-chitosan 16 μg/ml</td>
</tr>
<tr>
<td><em>Staphylococcus aureus</em></td>
<td>Chitosan 0.025%(w/v); chitosan-Zn complex 0.0125%(w/v); N-methyl-chitosan 16 μg/ml</td>
</tr>
<tr>
<td><em>Candida albicans</em></td>
<td>Chitosan 5 μg/ml; chitosan-Zn complex 0.1%(w/v)</td>
</tr>
</tbody>
</table>

2.7. Anti-leishmanial activity of chitosan

A few researchers have evaluated the activity of chitosan against *Leishmania* parasites under different conditions and the results are summarised in Table 2.2. It was observed that chitosan presented an anti-leishmanial activity with EC$_{50}$ (50% effective concentration) values ranging from around 50 to 240 μg/ml against different strains of *Leishmania* promastigotes and amastigotes. In these studies, there are inconsistent values of the activity of chitosan against *Leishmania* parasites. Accordingly, a lot of clarification and detailed controlled studies are needed to determine whether chitosan is a suitable candidate to find new chemotherapeutic alternatives for the treatment of leishmaniasis. The aim of this chapter was to: (i) determine the *in vitro* anti-leishmanial activity of chitosan and its derivatives against *L. major* and *L. mexicana* promastigotes and intracellular amastigotes at two different pH values (the culture medium pH of 7.5 and a lower pH of 6.5, which are both suitable for macrophage and parasite growth (188, 189, 190), (ii) to evaluate
the \textit{in vitro} role of chitosan in the activation of macrophage and M1 proinflammatory phenotype, via the measurement of NO, ROS and TNF-\textalpha production by host cells and by measuring parasite survival and (iii) investigate chitosan uptake by macrophages to explain its activity against intracellular amastigotes.
<table>
<thead>
<tr>
<th>Strain</th>
<th>Drug</th>
<th>Type of study</th>
<th>Results</th>
</tr>
</thead>
</table>
| L. infantum| Chitosan solution in acetic acid    | *In vitro*, against promastigotes and amastigotes | - EC<sub>50</sub> = 112.64 μg/ml promastigotes  
- EC<sub>90</sub> = 100.81 μg/ml amastigotes (127) |
| L. amazonensis | Chitosan solution in acetic acid | *In vitro*, against promastigotes and amastigotes | - EC<sub>50</sub> = 73.00 μg/ml promastigotes.  
-100 μg/ml: Percentage of infected macrophages after treatment 66% (122) |
| L. amazonensis | Chitosan nanoparticles          | *In vitro*, against promastigotes and amastigotes | - EC<sub>50</sub> = 52 μg/ml promastigotes.  
-100 μg/ml: Percentage of infected macrophages after treatment 39% (122) |
| L. chagasi  | Chitosan solution in acetic acid    | *In vitro*, against promastigotes       | EC<sub>50</sub> = 67 μg/ml promastigotes (122). |
| L. chagasi  | Chitosan nanoparticles            | *In vitro*, against promastigotes       | EC<sub>50</sub> = 46 μg/ml promastigotes (122). |
| L. infantum| Chitosan solution                  | *In vitro*, against promastigotes       | EC<sub>50</sub> = 240 μg/ml (128) |
| L. major    | nanochitosan film                  | *In vivo* study, female BALB/c mice were treated with nano-chitosan film four times/day | There was no significant difference between nanochitosan and Glucantime in reduction of lesion size of *L. major* infected mice (191) |
| L. major    | Chitosan microparticles            | *In vivo*, (100 μg/100 μl) were subcutaneously injected in the infected BALB/c mice) with two-day intervals until two weeks | Lesions of *L. major* infected mice were significantly smaller in chitosan treated groups (1.2 ± 0.8 mm) than in the control group (6.2 ± 1.7 mm) (129) |
| L. major    | Chitosan solution in acetic acid   | *In vivo* (BALB/c mice), chitosan 200 and 400 μg/ml were applied topically for 28 continuous days | Lesion size *L. major* infected mice was 8.47 mm for untreated group and 2.07 and 1.05 mm in groups treated with the 200 and 400 μg/ml of chitosan, respectively (192). |
| A pilot clinical study for 10 patients with CL (patients did not respond to traditional treatment) | poly (vinyl alcohol)/chitosan/clay nanocomposite film | Chitosan films were applied on the lesion for 7 days. This dressing was repeated every week until the complete healing. | Lesions were completely cured after 16 weeks with no side effects or recurrences (193). |
2.8. Materials and methods

2.8.1. Drugs and chemicals

Stocks of amphotericin B deoxycholate (5.2 mM [aq]) (Fungizone; Gibco, UK) were prepared, aliquoted, and kept at -20°C until use. Chitosan with three different molecular weights and its derivatives were used and are summarised in Table 1. Solutions of chitosan and derivatives were prepared by dissolving 1 g in 100 ml of 1% (v/v) acetic acid solution at room temperature with continuous stirring for 24 hours until a clear solution was obtained. The pH of the solution was adjusted to approximately 6 by adding sodium hydroxide 2N (NaOH, Sigma, UK) solution with a pH meter (Orion Model 420A). The chitosan solutions were autoclaved (121 °C; 15 mins). Phosphorylcholine substituted chitosan was kindly provided by Prof F Winnik (Montreal University, Canada) generated through reductive amination of PC-glyceraldehyde with primary amines of deacetylated chitosan (57 KDa). Percentage of substitution was controlled and determined by NMR (194). In our study, two pH values have been used: 7.5 is the medium culture pH and a lower pH 6.5. pH 6.5 is a suitable and safe pH for both macrophages and parasites, while pH<6.5 affects the growth of both macrophages and intracellular amastigotes.

Chitosan pKa is approximately 6.3 and therefore, the approximate ionisation degree of chitosan is a 61% and 6% at pH 6.5 and 7.5 respectively.

Table 2.3. Details of chitosan and its derivatives used in the study

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Properties</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>HMW (source: crustacean shells)</td>
<td>MW=310-375 KDa</td>
<td>Sigma, UK</td>
</tr>
<tr>
<td>MMW (source: crustacean shells)</td>
<td>MW=190-310 KDa</td>
<td>Sigma, UK</td>
</tr>
<tr>
<td>LMW (source: crustacean shells)</td>
<td>MW=50-190 KDa</td>
<td>Sigma, UK</td>
</tr>
<tr>
<td>Fungal chitosan (white mushroom)</td>
<td>MW=110-150 KDa</td>
<td>Dr Somavarapu</td>
</tr>
<tr>
<td>Chitosan oligosaccharide(synthetic)</td>
<td>MW=≤ 5KDa</td>
<td>Dr Somavarapu</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>-----------</td>
<td>---------------</td>
</tr>
<tr>
<td>Chitosan oligosaccharide lactate (synthetic)</td>
<td>MW=average Mn 5, oligosaccharide 60%</td>
<td>Dr Somavarapu</td>
</tr>
<tr>
<td>Chitosan- HCl (synthetic)</td>
<td>MW= 47 - 65 KDa</td>
<td>Dr Somavarapu</td>
</tr>
<tr>
<td>Carboxymethyl chitosan (synthetic)</td>
<td>MW=543.519 Da, level of substitution is 95%</td>
<td>Dr Somavarapu</td>
</tr>
<tr>
<td>PC1-CH (Phosphorylcholine substituted chitosan) (synthetic)</td>
<td>MW=33 KDa, PC(mol%)=30</td>
<td>Prof Winnik (194)</td>
</tr>
<tr>
<td>PC2-CH (synthetic)</td>
<td>MW=108 KDa, PC(mol%)=20</td>
<td>Prof Winnik (194)</td>
</tr>
<tr>
<td>PC3-CH (synthetic)</td>
<td>MW=109 KDa, PC(mol%)=30</td>
<td>Prof Winnik (194)</td>
</tr>
</tbody>
</table>
2.8.2. Ethics statement.

All animal work is carried out under a UK Home Office project licence according to the Animal (Scientific Procedures) Act 1986 and the new European Directive 2010/63/EU. The Project Licence (70/8427) has been reviewed by LSHTM Animal Welfare & Ethical Review Board prior to submission and consequent approval by the UK Home Office.

2.8.3. Cell lines

Preparation of macrophages
- Peritoneal mouse macrophages (PEMs) were obtained from 8-12-week-old female CD1-mice (Charles River Ltd, UK). Two ml of a 2% (w/v) starch solution in phosphate-buffered saline (PBS, Sigma, UK) was injected intraperitoneally (i.p.). After 24 h, the animal was sacrificed and the PEMs were harvested by peritoneal lavage with cold RPMI 1640 medium (Sigma, UK) containing 200 units penicillin and 0.2 mg streptomycin/ml (PenStrep; Sigma, UK). Subsequently, PEMs were centrifuged at 450 g at 4°C for 15 min and then the pellet was resuspended in RPMI 1640 with 10% (v/v) heat-inactivated fetal calf serum (HiFCS; Gibco, UK).

- Bone marrow-derived macrophages (BMMs) were obtained from femurs of 8-12-week-old female BALB/c mice (Charles River Ltd). Briefly, the bone marrow cells were carefully flushed from the bone with Dulbecco’s Modified Eagle’s Medium (DMEM; Thermofisher, UK) with 10% (v/v) HiFCS, 100 U/ml penicillin and 100 mg/ml streptomycin (Sigma, UK). Cells were pelleted by centrifugation (450 g, 10 min) and re-suspended in 10 ml DMEM with 10% (v/v) HiFCS and human macrophage colony stimulating factor 50ng/ml (HM-CSF; Thermofisher, UK). After plating out in T175 flasks (Greiner Bio-One, Stonehouse, UK), BMMs were kept at 37°C, 5% CO_2 for 7-10 days after which they were harvested, counted and used.

- THP-1 cell is a human leukemic monocyte-like derived cell line. THP-1 cells were cultured in RPMI 1640 medium supplemented with L-glutamine and 10% HiFCS. THP-1 cells were incubated in RPMI 1640 plus 10% (v/v) HiFCS and 20 ng/ml phorbol 12-myristate 13-acetate (PMA; Sigma, UK) at 37°C and 5% CO_2 for 72 h to induce maturation transformation of these monocytes into adherent macrophages (198)

**Human squamous carcinoma (KB) cells** are adherent cells derived from epidermal carcinoma from the mouth. KB cells were cultured in RPMI 1640 medium 10% HiFCS (199).

The number of cells and macrophages was estimated by counting with a Neubauer haemocytometer by light microscopy (x 400 total magnification).
2.8.4. Parasites

Four *Leishmania* species; two GFP labelled species (*L. major* (MHOM/SU/73/5ASKH) and *L. mexicana* (MNYC/BZ/62/M379), kindly donated by Dr G Getti (University of Greenwich, UK) were used for the fluorescence microscope study. They were cultured in Schneider's insect medium (Sigma, UK) with 23% (v/v) HiFCS, 1× penicillin-streptomycin-glutamine (Gibco-Invitrogen) and supplemented with 700 μg/ml G418 (an aminoglycoside antibiotic, Sigma, UK). *L. major* (MHOM/SA/85/JISH118) and *L. mexicana* (MNYC/BZ/62/M379) were used for other experiments as described, minus the G418. Promastigotes were incubated at 26°C, maximum passage number used = 7.

2.8.5. *In vitro* cytotoxicity assays

Re-suspended KB cells (4 x 10⁴/100 µl) were allowed to adhere to the bottom of 96-well plate overnight and then exposed to specific concentrations of the compounds for 72 h at 37°C and 5% CO₂ incubator. Podophyllotoxin (Sigma, UK) was included as a positive control at a starting concentration of 0.05 μM. Cytotoxicity was evaluated by a cell viability assay using the resazurin sodium salt solution (AlamarBlue, Sigma, UK) which was prepared according to the manufacturer’s instructions. 20μL of the resazurin solution was added to each well of the plates and fluorescence (cell viability) was measured over a period of 1 to 24 h using a Spectramax M3 plate reader (EX/EM 530 / 580 nm and 550 nm cut off). Results were expressed as percentage inhibition = (100 – x)% viability (means ± standard deviation σ). Cytotoxicity was evaluated in RPMI 1640 at two pH values (at normal pH of RPMI 7.5 and at a lower pH 6.5). The pH of RPMI 1640 was reduced from 7.5 to 6.5 by adding 0.05M acidic buffer, 2-N-morpholino ethanesulfonic acid (MES, Sigma, UK). RPMI 1640 plus MES (0.05M) at pH=6.5 did not show any toxicity to KB-cells.

2.8.6. *In vitro* 72-hour activity of chitosan and its derivatives against extracellular *L. major* and *L. mexicana* promastigotes

Promastigotes in RPMI 1640 medium were tested while in the exponential growth phase. The promastigotes were diluted to a density of 5x10⁶
promastigotes/ml and then exposed to different concentrations of (HMW, MMW, and LMW) chitosan, chitosan derivatives and Fungizone (positive control) in 96 well plates for 72 h at 26°C. The activity of the compounds against promastigotes was evaluated using the Alamar Blue™ assay as previously described. pH plays a critical role in the solubility and protonation of chitosan, so the activity against promastigotes was evaluated at two different pH values (pH=7.5 and a lower pH of 6.5 by adding MES). In addition to the colorimetric method of measuring parasite viability, promastigotes were manually counted microscopically in a Neubauer haemocytometer. Results were expressed as percentage inhibition= 100% - x% viability (means ± SD).

2.8.7. In vitro 72-hour activity of chitosan and its derivatives against intracellular amastigotes of *L. major* and *L. mexicana*

One hundred microliters of PEMs culture at 4 x 10⁵ cells/ml, dispensed into each well of a16-well LabTek tissue culture slide (Thermo Fisher, UK) at pH 7.5 or pH 6.5 and incubated for 24 h at 37 °C in 5 % CO₂. After 24 h, the wells were washed with fresh culture medium to remove non-adherent cells. Stationary phase, low-passage-number *Leishmania* promastigotes were then added at a ratio of 5 :1 PEM. This infection ratio was previously found to give sufficiently high and reproducible infection levels. Slides were incubated for another 24h at 34 °C to mimic dermal temperatures in 5 % CO₂ (201). Any free, extracellular parasites were removed by washing the wells with cold culture medium. One slide was fixed with 100 % methanol for >30sec and stained with 10 % Giemsa for 5 minutes. The number of PEMs infected with *Leishmania* amastigotes per 100 macrophages was microscopically counted. All the experiments were conducted at macrophages infection levels above 80% prior to addition of chitosan. Chitosan, its derivatives and Fungizone© (a micellar suspension) at a range of concentrations (in quadruplicate) were added to the wells (100µl) and the slides were incubated for 72 h at 34 °C in 5 % CO₂. After 72 hours, the slides were fixed with 100% methanol for >30sec and stained with 10% Giemsa for 5 min. The slides were examined and the % of macrophages which were infected was counted. The anti-leishmanial activity of compounds was expressed as percentage reduction in infected
macrophages compared to untreated control wells (202). RPMI 1640 plus MES (0.05M) with pH=6.5 had no activity against *Leishmania* amastigotes.

### 2.8.8. Influence of the origin of the host cell on the *in vitro* activity of HMW chitosan against *L. major* amastigotes

A further two host cell types, THP-1 and BMMs were infected with *L. major* and the activity of HMW chitosan was assessed. THP-1 cells (were cultured in RPMI 1640 with 10% HiFCS) and BMMs (were cultured in DMEM with 10% HiFCS) were used to assess the host cell dependence of the anti-leishmanial activity of HMW chitosan (198). The experiment was conducted as described in section (vii) at pH 6.5.

### 2.8.9. Influence of incubation duration on chitosan activity against *L. major* amastigotes

The experiment was conducted using *L. major* amastigotes in BMMs host cell at pH 6.5 as described in section 2.8.7 after 4, 24, 48 and 72h of incubation with HMW chitosan and Fungizone as a positive control.

### 2.8.10. The role of HMW chitosan on BMMs activation

One hundred microliters of BMMs, PEMs and THP-1 macrophages (4 x 10\(^5\)/ml) in DMEM (BMMs) or RPMI (PEMs and THP-1) at pH=6.5 were plated in each well of 96 well plates (standard clear plates for nitric oxide assay and black wall/clear bottom plates for ROS and TNF-α assay) and incubated for 24 hours at 37 °C in 5 % CO2. Plates were washed with DMEM (BMMs) or RPMI (PEMs and THP-1) to remove non-adherent macrophages. *L. major* at 1:5 ratio (5 parasites per host cell) was then added to the wells and the plates were incubated for 24h at 34 °C in 5 % CO2 to allow infection of the adherent macrophages. After 24h incubation with macrophages infection rate more than 80%, the immune stimulatory effects of HMW chitosan was determined by quantifying the release of TNF-α, ROS and NO by the macrophages, as described below at pH 6.5. Then We have chosen BMMs to evaluate if the immunostimulatory effects of HMW chitosan have any important role in its anti-amastigotes activity as these macrophages are more homogenous than...
PEMs and THP-1 cells (203). Both PEMs and BMMs have been reported to have a similar acidic pH ≈ 5.5 of parasitophorous vacuoles of L. amazonensis infected PEMs and BMMs (204, 205, 206).

2.8.10.1. Measurement of TNF-α

HMW chitosan at a range of concentrations (1.64, 4.9,14.8, 44.4, 133.3 and 400 µg/ml) was added to infected and uninfected macrophages and the plates were incubated for 4 and 24 h at 34°C in 5% CO2. Lipopolysaccharides from Escherichia coli O26:B6 (LPS, 100ng/ml; Sigma, UK) was used as a positive control and inducer. TNF-α release by the macrophages was measured using a mouse TNF-α ELISA kit (ab208348, abcam, UK) according to the manufacturer's instructions using a Spectramax M3 microplate reader (wavelength 450 nm) to determine if HMW chitosan stimulates T helper 1 or T helper 2 cells.

2.8.10.2. Measurement of ROS

ROS was measured using a 2′,7′-dichlorofluorescein diacetate (DCFDA, cellular reactive oxygen species detection assay kit, abcam, UK). Uninfected and infected macrophages were treated with 25 µM DCFDA in in phosphate-buffered saline (PBS) buffer for 45 min at 37°C and then washed once in buffer. The cells were cultured at 34°C in 5% CO2 for 0.5, 1, 2, 4, 8 and 24 h, with a range of concentrations (1.64, 4.9,14.8, 44.4, 133.3 and 400 µg/ml) of HMW chitosan or in the presence of H2O2 (25mM) (Thermofisher, UK) as a positive control in DMEM (BMMs) or RPMI (PEMs and THP-1) + 10% HiFCS (pH=6.5) in quadruplicate wells. In some experiments, cells were pre-treated with a selective inhibitor of ROS, N-acetyl-L-cysteine (NAC, 5mM; Sigma, UK), for 2 hours before the addition of the inducer or chitosan. At 0.5, 1, 2, 4, 8 and 24 h the plates were read, using a Spectramax M3 microplate reader (Ex=485nm, Em=535nm).

2.8.10.3. Measurement of NO

NO was measured by the Griess reagent (Thermofisher, UK). HMW chitosan at a range of concentrations (1.64, 4.9,14.8, 44.4, 133.3 and 400 µg/ml) was
added to infected and uninfected macrophages and the plates were incubated at 4 and 24 h at 34°C in 5% CO2. LPS (100ng/ml) was used as a positive control. In some experiments, cells were pre-treated with a selective inhibitor of nitric oxide with NG-methyl-L-arginine acetate salt (0.4 mM, L-NMMA; Sigma, UK) for 2 hours before the addition of LPS. NO was quantified according to the kit protocol, Briefly, 150 μl of the cell culture supernatants (particulates were removed by centrifugation) was mixed gently with 150μl of the Griess reagent in a 96 well plates and the mixture was incubated for 30 mins at room temperature. The absorbance was measured using a Spectramax M3 plate reader (wavelength 548 nm). Sodium nitrite (Sigma, UK) at different concentrations was used to create a standard curve.

2.8.11. Uptake of chitosan by macrophages

The uptake of HMW chitosan was evaluated using two methods. The first method used two endocytosis inhibitors: cytochalasin D (1μg/ml , Sigma, UK) which is a phagocytosis inhibitor and dynasore (30 μg/ml, Sigma, UK) which inhibits pinocytosis (clathrin-mediated endocytosis (CME) by blocking GTPase activity of dynamin) (208, 209, 210). The second method used dynasore and rhodamine-labelled chitosan (MW 200 kDa, Creative PEGWorks, USA) to track cellular uptake of chitosan over time by fluorescence microscopy. We chose BMMs to evaluate the uptake of chitosan by macrophages as these macrophages are more homogenous than PEMs (203).

2.8.11.1. Activity of chitosan after inhibition of the endocytic pathway of BMMs

One hundred microliters of BMMs culture (4 x 10⁵/ml) in DMEM at pH 6.5 or pH=7.5 were dispensed into each well of 16-well LabTek™ culture slides and were infected with stationary phase L. major promastigotes. Some of the infected BMMs were pretreated with dynasore (30 μg/ml) or cytochalasin D (1μg/ml) for two hours. Subsequently, HMW chitosan was added to each well at concentrations of 1.64, 4.9,14.8, 44.4, 133.3 or 400 μg/ml and macrophages were incubated for 4 or 24 h at 34 °C in 5% CO₂. After each point, the slides were examined as described in section (vii). The inhibition activity of the uptake
(phagocytosis or pinocytosis) of the two inhibitor was evaluated by using a fluorescence plate reader, by using fluorescent latex beads and pHrodo™ Red dextran (211). We showed that cytochalasin caused 94% and 84% phagocytosis inhibition of fluorescent latex beads (Sigma-Aldrich, UK) after 4 h and 24 h respectively and dynasore caused 95% and 90% pinocytosis inhibition of pHrodo™ Red dextran (Mw= 10,000 MW, Thermo Fisher, UK) after 4 h and 24h respectively.

2.8.11.2. Microscopic imaging of the cellular uptake of rhodamine-labelled chitosan

The qualitative characterisation of chitosan uptake of cells was carried out by wide-field microscopy (Nikon Ti-E inverted microscope). Briefly, after deriving BMMs, 500μl of the BMMs (in DMEM plus 10% HiFCS at pH 6.5, 4 x 10^4 macrophages per ml) was seeded on each well of a 4-well LabTek tissue culture slide (Thermo Fisher, UK) and incubated for 24h at 37°C in 5% CO₂. Subsequently, 5 μg/ml of Hoechst 33342 stain (Ex/Em = 350/461 nm, Thermofisher, UK) as a nuclear dye was added and the slides were incubated for 30 min at 37°C in 5% CO₂. The macrophages were washed with PBS, *L. major*-GFP of *L. mexicana*-GFP was then added, at a ratio of 10:1 and further incubated for 24h at 34°C in 5% CO₂ (We used 10:1 ratio not 5:1 as previously as at this experiment different species of *L. major*-GFP and *L. mexicana*-GFP were used and the ratio 10:1 was sufficient to obtain a high infection rate). Macrophages were then washed with PBS and 500 µl of LysoTracker® far Red (50 nM, Ex/Em;647/668nm; Thermo Fisher, UK) was added to each well. The labelled, infected macrophages were then exposed to 30 μg/ml rhodamine-labelled chitosan (MW 200kDa, Creative PEGWorks, USA) in 500 µl of fresh DMEM plus 10% HiFCS pH 6.5 and incubated for 4 h and 24h at 37°C with live imaging at each time point. In some experiments, infected BMMs were pre-incubated with dynasore 30 μg/ml for 2 h before adding rhodamine-labelled chitosan. All the images were collected using a Nikon Ti-E inverted microscope equipped with (63x objective) using Nikon Elements software. Three images for each experiment were then analysed using ImageJ software (v 1.52, National Institutes of Health, USA). The degree of correlation
between pixels in the red and green channels was assessed by the Colocalization Colormap plugin in the ImageJ software. This plugin enables quantitative visualisation of colocalization by calculating the normalized mean deviation product (nMDP) in a colour nMDP scale (from -1 to 1): negative refers (cold colours) to no co-localization while indexes more than 0 display co-localization and the higher number refers to more colocalization (212, 213).


Dose-response curves and EC$_{50}$ values were calculated by using GraphPad Prism version 7.02 software and the corresponding sigmoidal dose-response curves were established by using a nonlinear fit with variable slope models. Results represent means ± SD. EC$_{50}$ values were compared by using extra-sum-of-squares F tests. ANOVA and t-test were used to compare differences between two groups means or more. p values of 0.05 were considered statistically significant.

2.9. Results

2.9.1. Cytotoxicity of chitosan and its derivatives against KB cells in RPMI (pH 7.5 and pH 6.5)

The cytotoxicity of chitosan and its derivatives against KB cells was clearly observed in a dose-dependent manner at two pH values (6.5 and 7.5) as shown in Fig 2.4. Chitosan and its derivatives had a low toxicity toward KB cells at both pH values and there was no significant difference in the cytotoxicity at these two pH values ($p<0.05$ by an extra sum-of-squares F test) (Table 2.4). No significant difference in the cytotoxicity was observed between
the chitosans and the derivatives with LD$_{50} \approx 800$ μg/ml, except carboxymethyl chitosan with LD$_{50} \approx 1100$ μg/ml was significantly lower toxic ($p<0.05$ by an extra sum-of-squares F test).

**Table 2.4. In vitro cytotoxicity of chitosan and its derivatives against KB cells at two pH values after 72h of incubation**

<table>
<thead>
<tr>
<th>Compound</th>
<th>pH=7.5*,**</th>
<th>pH=6.5*,**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LD$_{50}$ μg/ml</td>
<td>LD$_{90}$ μg/ml</td>
</tr>
<tr>
<td>Podophyllotoxin</td>
<td>0.8 ± 0.03</td>
<td>2 ± 0.3</td>
</tr>
<tr>
<td>Fungizone</td>
<td>61 ± 8</td>
<td>228 ± 9</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>751 ± 88</td>
<td>3146 ± 377</td>
</tr>
<tr>
<td>MMW chitosan</td>
<td>752 ± 87</td>
<td>3033 ± 410</td>
</tr>
<tr>
<td>LMW chitosan</td>
<td>811 ± 93</td>
<td>3095 ± 425</td>
</tr>
<tr>
<td>Fungal chitosan</td>
<td>734 ± 95</td>
<td>3046 ± 377</td>
</tr>
<tr>
<td>Chitosan Oligosaccharide</td>
<td>727 ± 97</td>
<td>3115 ± 402</td>
</tr>
<tr>
<td>Chitosan Oligosaccharide-lactate</td>
<td>777 ± 98</td>
<td>3134 ± 388</td>
</tr>
<tr>
<td>Chitosan HCL</td>
<td>748 ± 90</td>
<td>3340 ± 409</td>
</tr>
<tr>
<td>PC1-CH</td>
<td>757 ± 91</td>
<td>3398 ± 388</td>
</tr>
<tr>
<td>PC2-CH</td>
<td>794 ± 90</td>
<td>3613 ± 400</td>
</tr>
<tr>
<td>PC3-CH</td>
<td>777 ± 90</td>
<td>3484 ± 357</td>
</tr>
<tr>
<td>Carboxymethyl chitosan</td>
<td>1183 ± 89</td>
<td>3800 ± 488</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown). Chitosan and its derivatives had a low toxicity at both pH values (6.5 and 7.5) toward KB-cells and there was no significant difference in the cytotoxicity at these two pH values ($p<0.05$ by t-test). ** No statistically significant difference was found in LD$_{50}$ (50% lethal dose) values between three types of chitosan and other derivatives against KB-cells (except carboxymethyl chitosan which is the least toxic) ($p>0.05$ by an extra sum-of-squares F test).
**Figure 2.4.** Dose-response curves of the cytotoxicity of chitosan and its derivatives against KB cells at pH=7.5 (A) and 6.5 (B). KB cells were cultured in the presence of different concentrations of chitosan and its derivatives. The toxicity of drugs was measured after 72 hours by measuring the inhibition of metabolic activity. Values are expressed as % inhibition of KB cells relative to untreated controls. No statistically significant difference was observed in LD50 values of chitosan and its derivatives against KB cells between pH=6.5 and pH=7.5 (p>0.05 by an extra sum-of-squares F test).
2.9.2. Analysis of anti-promastigotes activity

Anti-leishmanial activity (against promastigotes) of high, medium and low molecular weight (HMW, MMW and LMW respectively) chitosan and its derivatives (a total of 11) was tested. Dose-dependent activity (Fig 2.5) against *Leishmania* promastigotes was observed for chitosan and its' derivatives except for carboxymethyl chitosan which showed no activity against parasites within the experimental parameters tested (pH 7.5 or 6.5 and concentrations up to 400 µg/ml). Chitosan and its derivatives showed a higher anti-leishmanial activity (with around 7-20 times) at low pH compared with higher pH. Furthermore, (HMW, MMW and LMW) chitosan from crustacean source and fungal chitosan at pH= 6.5 showed a remarkable activity against *L. major* and *L. mexicana* promastigotes and were more active than other derivatives (*p*<0.05 by an extra sum-of-squares F test) as shown in Table 2.5.
Table 2.5. *In vitro* activity of chitosan and its derivatives against promastigotes at two pH values after 72h of incubation

<table>
<thead>
<tr>
<th>Compound</th>
<th>pH=7.5 * L. major</th>
<th>pH=7.5* L. <em>mexicana</em></th>
<th>pH=6.5*,** L. major</th>
<th>pH=6.5*,** L. <em>mexicana</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EC50 µg/ml</td>
<td>EC90 µg/ml</td>
<td>EC50 µg/ml</td>
<td>EC90 µg/ml</td>
</tr>
<tr>
<td>Fungizone</td>
<td>0.05 ± 0.01</td>
<td>0.2 ± 0.02</td>
<td>0.14 ± 0.01</td>
<td>0.3 ± 0.03</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>105 ± 12</td>
<td>1549 ± 525</td>
<td>140 ± 12</td>
<td>2187 ± 928</td>
</tr>
<tr>
<td>MMW chitosan</td>
<td>113 ± 9</td>
<td>1277 ± 580</td>
<td>150 ± 12</td>
<td>2223 ± 681</td>
</tr>
<tr>
<td>LMW chitosan</td>
<td>118 ± 11</td>
<td>1238 ± 582</td>
<td>157 ± 13</td>
<td>2225 ± 723</td>
</tr>
<tr>
<td>Fungal chitosan</td>
<td>118 ± 11</td>
<td>1228 ± 560</td>
<td>150 ± 13</td>
<td>1991 ± 580</td>
</tr>
<tr>
<td>Chitosan Oligosaccharide</td>
<td>153 ± 15</td>
<td>1680 ± 506</td>
<td>190 ± 20</td>
<td>2366 ± 461</td>
</tr>
<tr>
<td>Chitosan Oligosaccharide-lactate</td>
<td>98 ± 9</td>
<td>1226 ± 130</td>
<td>125 ± 14</td>
<td>765 ± 83</td>
</tr>
<tr>
<td>Chitosan HCL</td>
<td>96 ± 7</td>
<td>1189 ± 211</td>
<td>110 ± 24</td>
<td>746 ± 169</td>
</tr>
<tr>
<td>PC1-CH</td>
<td>111 ± 20</td>
<td>1875 ± 230</td>
<td>176 ± 14</td>
<td>2832 ± 412</td>
</tr>
<tr>
<td>PC2-CH</td>
<td>104 ± 6</td>
<td>1485 ± 259</td>
<td>170 ± 8</td>
<td>2744 ± 377</td>
</tr>
<tr>
<td>PC3-CH</td>
<td>119 ± 19</td>
<td>1860 ± 365</td>
<td>187 ± 16</td>
<td>3175 ± 580</td>
</tr>
<tr>
<td>Carboxymethyl chitosan</td>
<td>No activity up to 400 µg/ml</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown). *Statistically significant differences were found for the EC50 values of chitosan and its derivatives at pH=6.5 and pH=7.5 (p<0.05 by using t-test). ** *L. major* promastigotes were significantly more susceptible to chitosan and derivatives than *L. mexicana* (p<0.05 by an extra sum-of-squares F test). Amphotericin B deoxycholate (Fungizone) was used as a positive control. Both pH of 6.5 and chitosan solvent did not show any activity against promastigotes. Both RPMI alone pH 6.5 and chitosan solvent did not show any activity against promastigotes.
Figure 2.5. Dose-response curves of the activity of chitosan and its derivatives against *Leishmania* promastigotes at two pH values. A: *L. major* at pH=7.5; B: *L. mexicana* at pH = 7.5; C: *L. major* at pH = 6.5; D: *L. mexicana* at pH = 6.5. Promastigotes were cultured in the presence of different concentrations of chitosan and its derivatives. The activity of drugs was measured after 72h by measuring the inhibition of metabolic activity. Values are expressed as % inhibition of promastigotes relative to untreated controls. Statistically significant difference was observed in EC$_{50}$ values of chitosan and its derivatives against *L. mexicana* and *L. major* promastigotes between pH=6.5 and pH=7.5 (p<0.05 by t-test). Both RPMI alone pH 6.5 and chitosan solvent did not show any activity against promastigotes.

### 2.9.3. Analysis of anti-amastigotes activity in PEMs

Anti-leishmanial activity (against amastigotes) of high, medium and low molecular weight (HMW, MMW and LMW respectively) chitosan and its derivatives (a total of 11) was tested. Dose-dependent activity (Fig 2.6) against *Leishmania* amastigotes was observed for chitosan and its’ derivatives except for carboxymethyl chitosan which showed no activity against amastigotes.
within the experimental parameters tested (pH 7.5 or 6.5 and concentrations up to 400 µg/ml). In the 72 hour assays, chitosan and its derivatives were significantly more active (with around 7-20 times) against intracellular *L. major* and *L. mexicana* amastigotes at pH 6.5 than pH 7.5 (*p*<0.05 by a paired t-test) as shown in Fig 2.6. (HMW, MMW and LMW) chitosan from crustacean source and fungal chitosan exhibited a significantly higher activity against *L. major* and *L. mexicana* intracellular amastigotes (EC$_{50}$ ≈ 12 µg/ml against *L. major* and 16 µg/ml against *L. mexicana*) than other derivatives at pH= 6.5 (*p*<0.05 by an extra sum-of-squares *F* test) as shown in Table 2.6.
Table 2.6. *In vitro* activity of chitosan and its derivatives against amastigotes infecting PEMs after 72h of incubation

<table>
<thead>
<tr>
<th>Compound</th>
<th>pH 7.5*</th>
<th></th>
<th>pH 6.5*</th>
<th></th>
<th>pH 6.5*</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L. major</td>
<td>L. mexicana</td>
<td>L. major</td>
<td>L. mexicana</td>
<td>L. major</td>
<td>L. mexicana</td>
</tr>
<tr>
<td></td>
<td>EC₅₀ µg/ml</td>
<td>EC₉₀ µg/ml</td>
<td>EC₅₀ µg/ml</td>
<td>EC₉₀ µg/ml</td>
<td>EC₅₀ µg/ml</td>
<td>EC₉₀ µg/ml</td>
</tr>
<tr>
<td>Fungizone</td>
<td>0.07 ± 0.01</td>
<td>0.13 ± 0.05</td>
<td>0.19 ± 0.05</td>
<td>1.5 ± 0.2</td>
<td>0.06 ± 0.01</td>
<td>0.11 ± 0.06</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>98 ± 6</td>
<td>1635 ± 245</td>
<td>119 ± 9</td>
<td>1804 ± 304</td>
<td>11.4 ± 1</td>
<td>69 ± 18</td>
</tr>
<tr>
<td>MMW chitosan</td>
<td>103 ± 8</td>
<td>1652 ± 287</td>
<td>125 ± 10</td>
<td>1793 ± 323</td>
<td>12.9 ± 1</td>
<td>81 ± 18</td>
</tr>
<tr>
<td>LMW chitosan</td>
<td>102 ± 7</td>
<td>1651 ± 282</td>
<td>125 ± 10</td>
<td>1795 ± 320</td>
<td>12.1 ± 1</td>
<td>74 ± 14</td>
</tr>
<tr>
<td>Fungal chitosan</td>
<td>102 ± 7</td>
<td>1650 ± 276</td>
<td>124 ± 9</td>
<td>1796 ± 316</td>
<td>12.6 ± 3</td>
<td>92 ± 27</td>
</tr>
<tr>
<td>Chitosan Oligosaccharide</td>
<td>145 ± 12</td>
<td>2473 ± 500</td>
<td>175 ± 14</td>
<td>2543 ± 505</td>
<td>73 ± 4</td>
<td>260 ± 32</td>
</tr>
<tr>
<td>Chitosan Oligosaccharide- lactate</td>
<td>93 ± 7</td>
<td>1957 ± 174</td>
<td>120 ± 9</td>
<td>2365 ± 239</td>
<td>39 ± 1</td>
<td>201 ± 16</td>
</tr>
<tr>
<td>Chitosan HCl</td>
<td>97 ± 11</td>
<td>2080 ± 516</td>
<td>121 ± 15</td>
<td>2402 ± 667</td>
<td>40 ± 2</td>
<td>210 ± 23</td>
</tr>
<tr>
<td>PC1-CH</td>
<td>144 ± 10</td>
<td>1292 ± 217</td>
<td>169 ± 12</td>
<td>1365 ± 212</td>
<td>68 ± 3</td>
<td>246 ± 26</td>
</tr>
<tr>
<td>PC2-CH</td>
<td>133 ± 6</td>
<td>1005 ± 194</td>
<td>159 ± 6</td>
<td>1705 ± 170</td>
<td>60 ± 3</td>
<td>202 ± 22</td>
</tr>
<tr>
<td>PC3-CH</td>
<td>163 ± 11</td>
<td>1052 ± 144</td>
<td>187 ± 10</td>
<td>1107 ± 142</td>
<td>71 ± 4</td>
<td>251 ± 30</td>
</tr>
<tr>
<td>Carboxymethyl chitosan</td>
<td>No activity up to 400 µg/ml</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown). *Statistically significant differences were found between the EC₅₀ values of chitosan and its derivatives at pH=6.5 and pH=7.5 (*p*<0.05 by using t-test). Both pH of 6.5 and chitosan solvent did not show any activity against amastigotes. Both RPMI alone pH 6.5 and chitosan solvent did not show any activity against amastigotes.
Figure 2.6. Dose-response curves of the activity of chitosan and its derivatives against *Leishmania* amastigotes at two pH values. A: *L. major* at pH=7.5; B: *L. mexicana* at pH = 7.5; C: *L. major* at pH = 6.5; D: *L. mexicana* at pH = 6.5. PEMs were infected with stationary-phase promastigotes and exposed to various concentrations of chitosan and its derivatives, followed by microscopic counting of the number of infected macrophages*. Values are expressed as % inhibition of infection relative to untreated controls. Chitosan and its derivatives are significantly more active in pH 6.5 than in pH 7.5 (p<0.05 by t-test). * Macrophage infection rate was >80% after 24h. Both RPMI alone pH 6.5 and chitosan solvent did not show any activity against amastigotes.

To allow like-for-like comparison, EC\textsubscript{50} values were recalculated in terms of molarity using estimated molecular weights (HMW: MW= 342.5 KDa, MMW: MW=250 KDa, LMW: MW= 120 KDa and fungal chitosan MW=130 KDa) at pH = 6.5. Based on molarity (Table 2.7 and 2.8 ), HMW chitosan was significantly more
active against *L. major* and *L. mexicana* promastigotes and amastigotes and hence used in all subsequent studies. Fig 2.7 observes the morphology of infected macrophages before and after treatment with HMW chitosan is taken by a microscope provided with a digital camera.

**Table 2.7.** *In vitro* activity of chitosans against promastigotes based on molarity

<table>
<thead>
<tr>
<th>Compound</th>
<th>pH=6.5*,**</th>
<th>pH=6.5*,**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><em>L. major</em></td>
<td><em>L. mexicana</em></td>
</tr>
<tr>
<td></td>
<td>EC₅₀ µM</td>
<td>EC₉₀ µg/ml</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>0.017 ± 0.001</td>
<td>0.10 ± 0.02</td>
</tr>
<tr>
<td>MMW chitosan</td>
<td>0.024 ± 0.001</td>
<td>0.172 ± 0.03</td>
</tr>
<tr>
<td>LMW chitosan</td>
<td>0.05 ± 0.001</td>
<td>0.33 ± 0.06</td>
</tr>
<tr>
<td>Fungal chitosan</td>
<td>0.05 ± 0.003</td>
<td>0.31 ± 0.005</td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD HMW chitosan is significantly more active against *Leishmania* promastigotes than other types (p <0.05 by one-way ANOVA)

**Table 2.8.** *In vitro* activity of chitosans against amastigotes based on molarity

<table>
<thead>
<tr>
<th>Compound</th>
<th>pH 6.5*</th>
<th>pH 6.5*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><em>L. major</em></td>
<td><em>L. mexicana</em></td>
</tr>
<tr>
<td></td>
<td>EC₅₀ µM</td>
<td>EC₉₀ µM</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>0.03 ± 0.01</td>
<td>0.2 ± 0.05</td>
</tr>
<tr>
<td>MMW chitosan</td>
<td>0.05 ± 0.04</td>
<td>0.32 ± 0.07</td>
</tr>
<tr>
<td>LMW chitosan</td>
<td>0.1 ± 0.002</td>
<td>0.6 ± 0.1</td>
</tr>
<tr>
<td>Fungal chitosan</td>
<td>0.09 ± 0.008</td>
<td>0.7 ± 0.2</td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD HMW chitosan is significantly more active against *Leishmania* promastigotes than other types (p <0.05 by one-way ANOVA)

**Figure 2.7.** Morphology of infected (PEMs) with *L. major* and *L. mexicana* after treatment with HMW chitosan. Slides were fixed with 100% methanol for 5 minutes and stained with 10 % Giemsa for 5 minutes. These figures have been taken by a microscope attached to a digital camera. A: *L. major* infected macrophages before treatment (*L. major* amastigotes with tight vacuoles). B: *L. mexicana* infected macrophages before treatment (*L. mexicana* amastigotes with large vacuoles). C: *L. major* infected macrophages after treatment with HMW chitosan.
2.9.4. Host cell dependence of antileishmanial activity of HMW chitosan and time to kill assay on amastigotes at pH 6.5

We aimed to assess the host cell dependence of anti-leishmanial activity of HMW chitosan and Fungizone by evaluating the *in vitro* activity against *L. major* amastigotes in three different host cells (PEMs, BMMs and THP-1). EC$_{50}$ and EC$_{90}$ values of HMW chitosan and Fungizone against amastigotes infecting three different macrophage populations are summarized in Table 2.9. As can be seen, there was a significant difference in the activity of HMW chitosan and Fungizone depending on the type of the host cells ($p < 0.05$ by one-way ANOVA) and both HMW chitosan and Fungizone displayed higher activity in PEMs and BMMs than in differentiated THP-1 cells. The results in Fig 2.8 clearly show that both HMW chitosan and Fungizone had time-dependent effects against intracellular amastigotes in RPMI with pH=6.5.

**Table 2.9.** HMW chitosan activity against *L. major* amastigotes in three different macrophage cultures after 72 h

<table>
<thead>
<tr>
<th>Host cell/infection rate % at 24h</th>
<th>HMW chitosan</th>
<th>Fungizone</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EC$_{50}$ µg/ml</td>
<td>EC$_{90}$ µg/ml</td>
</tr>
<tr>
<td>PEMs / &gt; 80%</td>
<td>10.31 ± 1.22*</td>
<td>89.07 ± 20.46</td>
</tr>
<tr>
<td>BMMs / &gt; 80%</td>
<td>14.60 ± 1.79*</td>
<td>145.7 ± 36.2</td>
</tr>
<tr>
<td>THP-1 / &gt; 80%</td>
<td>24.28 ± 2.87*</td>
<td>200.1 ± 48.8</td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown). *,** statistically significant difference in EC$_{50}$ values between the three types of cells (Fungizone and were significantly more active in PEMs and BMMs compared with THP-1 cells) ($p<0.05$ by an extra sum-of-squares F test) taking into consideration that infection levels were higher in PEMs and THP-1 than BMMs. % infection rate gives the percentage of infected macrophages. . Both pH of 6.5 and chitosan solvent did not show any activity against amastigotes.
Figure 2.8. Influence of incubation duration on the chitosan and Fungizone activity against *L. major* intracellular amastigotes in BMMs. BMMs were infected with stationary-phase promastigotes and exposed to fixed concentrations of chitosan HMW and Fungizone for 4, 24, 48 and 72 h, followed by microscopic counting of the number of infected macrophages. (A) *In vitro* time-to-kill for Fungizone (B) *In vitro* time-to-kill for chitosan HMW. Results shown are the means ± SD of quadruplicates and represent one experiment of three performed.

2.9.5. Effects of HMW chitosan on the production of TNF-α by uninfected or *L. major* infected macrophages at pH = 6.5

The activation of M1 macrophages by Th1 lymphocyte plays an important role in the control of CL (14, 15, 16) therefore, we measured TNF-α production by macrophages stimulated by HMW chitosan. Following the stimulation by HMW chitosan, the TNF-α production by macrophages (BMMs, PEMs and THP-1) was found to be in a dose-dependent manner in both infected and uninfected cells as shown in Fig 2.9. After 24 h, the levels of TNF-α in the culture fluid of macrophages (both infected and uninfected BMMs, PEMs and THP-1) containing concentrations of HMW chitosan (14.8, 44.4 and 133.3 μg/ml) was significantly higher than untreated macrophages, with TNF-α being highest at 44.4 μg/ml chitosan. While at other concentrations (1.64, 4.9 and 400 μg/ml), HMW chitosan did not stimulate macrophages to produce TNF-α (p < 0.05 by t-test). HMW chitosan at concentrations 14.8, 44.4 and 133.3 μg/ml stimulated uninfected BMMs to produce TNF-α with 87± 4.5 - 712± 9 - 48±3 pg/ml, uninfected PEMs with 67± 5 - 570± 8 - 33±3 pg/ml and uninfected THP-1 with 47± 3.5 - 412± 10 -
22±3 pg/ml respectively and 56± 3.5 - 464± 10 - 32±4 pg/ml, 46± 5 - 400± 7 - 22±4 pg/ml and 36± 2 - 310± 10 - 15±4 pg/ml in L. major infected BMMs, PEMs and THP-1 respectively. In other words, HMW chitosan stimulated less amount of TNF-α in L. major infected than uninfected macrophages (p < 0.05 by t-test) and BMMs produced higher levels of TNF-α after the stimulation in comparison with PEMs and THP-1(p < 0.05 by one-way-ANOVA). Less TNF-α was generated when the chitosan concentration was increased to 133.3 µg/ml and above. Lipopolysaccharides from Escherichia coli O26:B6 (LPS; positive control) stimulated TNF-α production in both uninfected and infected BMMs, PEMs and THP-1 after a 24 h incubation period and at a significantly higher level than chitosan (p < 0.05 by t-test). Our results indicated that HMW chitosan activated M1 macrophages.
Figure 2.9. TNF-α production in uninfected and *L. major* infected BMMs, PEMs and THP-1 macrophages* after 24 h of exposure to 1.64, 4.9, 14.8, 44.4, 133.3 and 400 µg/ml of chitosan at pH = 6.5. The dose-response in both uninfected and *L. major* infected macrophages was bell-shaped. TNF-α production was significantly decreased (p < 0.05 by t-test) by infecting the cells with *L. major*. TNF-α stimulation was higher with the rank BMMs, PEMs and THP-1. Experiments were conducted in quadruplicate, data are expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown). Positive control= macrophages treated with LPS 10 µg/ml. Negative control = macrophages not exposed to chitosan. *Initial macrophage infection rate was >80% after 24 h. Chitosan solvent did not cause any TNF-α production.

### 2.9.6. Effects of HMW chitosan on the production of ROS by macrophages at pH = 6.5

ROS plays an important role in the killing of intracellular amastigotes (14, 15, 16) therefore, we measured ROS production by macrophages stimulated by HMW chitosan. HMW chitosan (at concentrations 14.8, 44.4 and 133.3 µg/ml) increased the production of ROS (indicated by H2DCFDA fluorescence) after 4 h of incubation but did not stimulate ROS after 8 h of incubation (Table 2.10). Other concentrations of HMW chitosan (1.64, 4.9 and 400 µg/ml) did not stimulate BMMs, PEMs or THP-1 to produce ROS after 4 h or 8 h of incubation.

The ROS response in both uninfected and infected BMMs, PEMs and THP-1 was in bell shaped – similar to that seen with TNF-α. Increasing chitosan concentration (more than 14.8 µg/ml) increased ROS production until concentration 44.4 µg/ml (the maximum production of ROS), after which increasing concentration reduced
ROS production. In addition, we showed that ROS production by macrophages was significantly decreased (p < 0.05 by t-test) by infecting the cells with *L. major* as shown in Fig 2.10. BMMs produced higher levels of ROS after the stimulation in comparison with PEMs and THP-1 (p < 0.05 by one-way-ANOVA).

**Table 2.10.** ROS production in uninfected and *L. major* infected BMMs after 8 h of exposure to different concentrations of HMW chitosan at pH=6.5

<table>
<thead>
<tr>
<th>Chitosan µg/ml</th>
<th>Uninfected BMMs</th>
<th>Infected BMMs</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.64</td>
<td>4000 ± 100</td>
<td>2650 ± 100</td>
</tr>
<tr>
<td>4.9</td>
<td>3999 ± 200</td>
<td>2550 ± 150</td>
</tr>
<tr>
<td>14.81</td>
<td>4020 ± 150</td>
<td>2650 ± 100</td>
</tr>
<tr>
<td>44.4</td>
<td>4050 ± 100</td>
<td>2750 ± 200</td>
</tr>
<tr>
<td>133.3</td>
<td>4000 ± 200</td>
<td>2564 ± 150</td>
</tr>
<tr>
<td>400</td>
<td>3959 ± 100</td>
<td>2400 ± 100</td>
</tr>
<tr>
<td><strong>Negative control = BMMs not exposed to chitosan</strong></td>
<td>4750 ± 100</td>
<td>2850 ± 100</td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate, data is expressed as mean +/- SD (experiment was reproduced a further two times with confirmed similar data (not shown). Chitosan solvent alone did not cause any ROS production.
Figure 2.10. ROS production in uninfected and L. major infected BMMs, PEMs and THP-1 macrophages * after 4 h of exposure to 1.64, 4.9, 14.8, 44.4, 133.3 and 400 µg/ml of HMW chitosan at pH=6.5. High levels of ROS were induced by both uninfected and L. major infected macrophages exposed to HMW chitosan compared to those that were not (P <0.05 by t-test). Maximum production of ROS occurred at 44.4 µg/ml of chitosan. ROS production by L. major infected macrophages was significantly lower compared to uninfected cells (p < 0.05 by t-test). ROS stimulation was higher with the rank BMMs, PEMs and THP-1. Experiments were conducted in quadruplicate, data is expressed as mean +/- SD (experiment was reproduced a further two times with confirmed similar data (not shown). Positive control = macrophages treated with H$_2$O$_2$ 25 mM (a known ROS inducer). Negative control = macrophages not exposed to chitosan. *Initial macrophage infection rate was >80% after 24 h. Chitosan solvent did not cause any ROS production.

We found that HMW chitosan had an in vitro stimulatory effect on BMMs ROS production after 4h of incubation. We therefore investigated whether this ROS plays any role in the activity of HMW chitosan against intracellular amastigotes. For these experiments, the 4 h post treatment time point was taken because ROS
peaked at this point in BMMs in response to chitosan treatment at a time when chitosan does not induce NO in BMMs \( \textit{ibid} \). Scavenging of ROS by the ROS scavenger, 5mM N-acetyl-L-cysteine (NAC), had no significant impact on the activity of chitosan against intracellular amastigotes \( (p > 0.05 \text{ by } t\text{-test}) \) – see Fig 2.11. ROS scavenger (N-acetyl-L-cysteine (NAC), 5mM) caused a complete scavenging of ROS after 4 h (Table 2.11). and had no cytotoxicity against KB cells or leishmanicidal against \textit{L. major} amastigotes. Even though chitosan stimulated ROS production but this did not play a role in the anti-leishmanial activity of chitosan.

![Graph](image)

\textbf{Figure 2.11.} Activity of HMW chitosan against \textit{L. major} amastigotes in BMMs* after 4 h, with and without ROS scavenger at pH = 6.5. Infected macrophages were pre-incubated with 5 mM NAC for 2 h, after which HMW chitosan at concentrations 1.64, 4.9,14.8, 44.4, 133.3 and 400 µg/ml was added and the cells were incubated for a further 4 h. Chitosan activity against intracellular amastigotes was evaluated as described in section (vii). Values are expressed as % inhibition of infection relative to untreated controls. After 4h, there was no significant difference in the anti-leishmanial activity of chitosan after scavenging of ROS \( (p >0.05 \text{ by } t\text{-test}) \). Experiments were conducted in quadruplicate, data is expressed as mean +/- SD. Experiment was reproduced further two times with confirmed similar data (not shown). *Initial macrophage infection rate was >80% after 24 h.
Table 2.11. ROS production in uninfected and L. major infected BMMs after exposure to chitosan in the presence of ROS scavenger

<table>
<thead>
<tr>
<th>Chitosan µg/ml</th>
<th>ROS (Relative Fluorescence Intensity) after 4 h in:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Uninfected BMMs pre-treated with ROS scavenger</td>
</tr>
<tr>
<td>1.64</td>
<td>4700 ± 200</td>
</tr>
<tr>
<td>4.9</td>
<td>4800 ± 250</td>
</tr>
<tr>
<td>14.81</td>
<td>4750 ± 100</td>
</tr>
<tr>
<td>44.4</td>
<td>4800 ± 100</td>
</tr>
<tr>
<td>133.3</td>
<td>4900 ± 150</td>
</tr>
<tr>
<td>400</td>
<td>4950 ± 100</td>
</tr>
<tr>
<td><strong>Positive control (ROS)</strong> = BMMs treated with 25 mM H₂O₂</td>
<td>4800 ± 250</td>
</tr>
<tr>
<td><strong>Negative control</strong> = BMMs not exposed to chitosan or to H₂O₂</td>
<td>4800 ± 100</td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate, data is expressed as mean +/- SD (experiment was reproduced a further two times with confirmed similar data (not shown). ROS was measured after 4 h of exposure to HMW chitosan.

2.9.7. Effects of HMW chitosan on the production of NO by macrophages at pH = 6.5

NO plays an important role in the killing of intracellular amastigotes (14, 15, 16) therefore, we measured NO production by macrophages stimulated by HMW chitosan. We showed that chitosan did not have a stimulatory effect on BMMs, PEMs and THP-1 NO production after 4 h of incubation (Table 2.12). However, after a 24 h incubation, HMW chitosan at pH=6.5 had a stimulatory effect on BMMs, PEMs and THP-1 NO production in a clear bell-shaped dose-dependent manner. HMW chitosan at concentrations 14.8, 44.4 and 133.3 µg/ml induced uninfected BMMs to produce NO with 14.9± 0.3 - 34±1.2 - 11±1 µM, uninfected PEMs with 10.9± 0.4 - 26±1.2 - 8.5±1 µM and uninfected THP-1 with 8.9± 0.2 - 20±1 - 6.1±0.5 µM respectively and 11±1 - 26 ± 2.5 - 8 ± 1.2 µM, 8 ±1 - 20 ± 2 - 6 ± 1.2 µM and 6± 0.1 - 14±1 - 4.1±0.5 µM in L. major infected BMMs, PEMs and THP-1 respectively, NO being highest at 44.4 µg/ml. While other concentrations of HMW chitosan (1.64, 4.9 and 400 µg/ml) did not stimulate macrophages to produce NO after 24 h of incubation. In other words, HMW chitosan stimulated a
lower quantity of NO in infected than uninfected macrophages \((p < 0.05 \text{ by t-test})\) and BMMs produced higher levels of NO after the stimulation in comparison with PEMs and THP-1\((p < 0.05 \text{ by one-way-ANOVA})\) (Fig 2.12).

LPS caused significantly higher NO production compared to HMW chitosan \((p < 0.05 \text{ by t-test})\) in both uninfected and infected BMMs, PEMs and THP-1. The levels of NO produced by \(L. \text{major}\) infected BMMs exposed to LPS (positive control) or HMW chitosan were significantly lower than levels produced by uninfected macrophages \((p < 0.05 \text{ by t-test})\) (Fig 2.12).

**Table 2.12.** NO production in uninfected and \(L. \text{major}\)-infected BMMs after 4h of exposure to different concentrations of HMW chitosan at pH=6.5

<table>
<thead>
<tr>
<th>Chitosan (\mu g/ml)</th>
<th>Uninfected BMMs</th>
<th>Infected BMMs</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.64</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>4.9</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>14.81</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>44.4</td>
<td>0.05 ± 0.01</td>
<td>0</td>
</tr>
<tr>
<td>133.3</td>
<td>0.06 ± 0.01</td>
<td>0.05 ± 0.01</td>
</tr>
<tr>
<td>400</td>
<td>0.05 ± 0.01</td>
<td>0.04 ± 0.01</td>
</tr>
</tbody>
</table>

Negative control = BMMs not exposed to chitosan

<table>
<thead>
<tr>
<th></th>
<th>Uninfected BMMs</th>
<th>Infected BMMs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative control = BMMs not exposed to chitosan</td>
<td>0.07 ± 0.01</td>
<td>0.05 ± 0.01</td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate, data is expressed as mean +/- SD (experiment was reproduced a further two times with confirmed similar data (not shown). Chitosan solvent alone did not cause any NO production.
Figure 2.12. NO production in uninfected and *L. major* infected BMMs, PEMs and THP-1 macrophages * after 24 h of exposure to 1.64, 4.9, 14.8, 44.4, 133.3 and 400 μg/ml of chitosan at pH = 6.5. The response in both uninfected and infected macrophages was bell-shaped in relation to chitosan concentration. Maximal production of NO was stimulated by 44.4 μg/ml of chitosan. NO production was significantly decreased (p < 0.05 by t-test) when the cells had been infected with *L. major*. NO stimulation was higher with the rank BMMs, PEMs and THP-1. Experiment was conducted in quadruplicate cultures, data expressed as mean +/- SD (experiment was reproduced a further two times with confirmed similar data and data not shown). Positive control = macrophages treated with LPS 10 μg/ml. Negative control = macrophages not exposed to chitosan.

*Initial macrophage infection rate was >80% after 24 h. Chitosan solvent did not cause any NO production.*

As HMW chitosan had an *in vitro* stimulatory effect on BMMs NO production after 24h of incubation we investigated further whether NO has any role in the activity of HMW chitosan against intracellular amastigotes. Inhibition of NO production by the NO inhibitor NG-methyl-L-arginine acetate salt (L-NMMA) at 0.4mM, had no
significant influence on the activity of chitosan against intracellular amastigotes (p > 0.05 by t-test) (Fig 2.13). The NO inhibitor (L-NMMA, 0.4 mM) caused 90% reduction in NO production (Table 2.13) after 24 h and had no cytotoxicity effects against KB cells and no leishmanicidal against intracellular *L. major* amastigotes. Even though chitosan stimulated NO production but this did not play a role in the anti-leishmanial activity of chitosan.

**Figure 2.13.** Activity of HMW chitosan against *L. major*-infected BMMs* after 24 h in the presence or absence of an NO inhibitor at pH = 6.5. Infected macrophages were pre-incubated with the NO inhibitor L-NMMA (0.4 mM) for 2 h, following which HMW chitosan at concentrations 1.64, 4.9, 14.8, 44.4, 133.3 and 400 µg/ml was added and the cells were incubated for a further 24 h. Chitosan activity against intracellular amastigotes was evaluated as described in section (vii). Values are expressed as % inhibition of infection relative to untreated controls. After 24 h, there was no significant difference in the activity of chitosan after inhibition of NO (p >0.05 by t-test). Experiment was conducted in quadruplicate cultures, data expressed as mean +/- SD. Experiment was reproduced a further two times and confirmed the results (data not shown). *Initial macrophage infection rate was >80% after 24 h.

**Table 2.13.** NO production in uninfected and *L. major* infected BMMs after exposure to chitosan in the presence of NO inhibitor at pH=6.5

<table>
<thead>
<tr>
<th>Chitosan µg/ml</th>
<th>NO µM after 24 h in:</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Uninfected BMMs pre-treated with NO inhibitor</td>
<td>Infected BMMs pre-treated with NO inhibitor</td>
<td></td>
</tr>
<tr>
<td>1.64</td>
<td>1.4 ± 0.4</td>
<td>0.15 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>4.9</td>
<td>1.5 ± 0.3</td>
<td>0.16 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>14.81</td>
<td>1.9 ± 0.5</td>
<td>0.17 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>44.4</td>
<td>1.6 ± 0.2</td>
<td>0.15 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>133.3</td>
<td>1.2 ± 0.4</td>
<td>0.14 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>400</td>
<td>1.0 ± 0.6</td>
<td>0.15 ± 0.1</td>
<td></td>
</tr>
</tbody>
</table>
Positive control (NO) = BMMs treated with 10 µg/ml LPS

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.4 ± 0.3</td>
<td>0.16 ± 0.1</td>
</tr>
</tbody>
</table>

Negative control = BMMs not exposed to chitosan or to LPS

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.7 ± 0.3</td>
<td>0.13 ± 0.1</td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate, data is expressed as mean +/- SD (experiment was reproduced a further two times with confirmed similar data (not shown). NO was measured after 24 h of exposure to HMW chitosan.

2.9.8. Cellular uptake of HMW chitosan and inhibition of endocytosis

We found that the activation of M1 macrophages by HMW chitosan did not play a role in its activity against intracellular amastigotes therefore, we investigated whether the anti-leishmanial effects of HMW chitosan against intracellular amastigotes after 4 h and 24 h exposure were dependent on the direct activity of chitosan following its entry into the macrophages at pH=6.5. No significant difference was observed in the activity of chitosan against intracellular amastigotes when it was added after prior phagocytosis inhibition with cytochalasin D (p > 0.05 by t-test). In contrast, dynasore (an inhibitor of pinocytosis, a clathrin-mediated endocytosis (CME) inhibitor) did significantly affect chitosan mediated parasite killing at pH = 6.5 as shown in Fig. 33(p< 0.05 by t-test). The same activity was seen at pH 7.5. – see Fig 2.14, panel C. We found that cytochalasin caused 94 and 84% phagocytosis inhibition of fluorescent latex beads (Sigma-Aldrich, UK) after 4 h and 24 h respectively and dynasore caused 95 and 90% pinocytosis inhibition of pHrodo™ Red dextran (Mw= 10,000 MW, Thermo Fisher, UK) after 4h and 24h respectively (Table 2.14). The two inhibitors had no activity against intracellular L. major amastigotes at the concentrations used. Pinocytosis (CME) played a critical role in the efficacy of HMW chitosan against intracellular amastigotes.
Figure 2.14. Activity of HMW chitosan against *L. major* infected BMMs* after 4 h, pH=6.5 (A), 24 h, pH=6.5 (B) and at 24 h, pH=7.5 with or without phagocytosis inhibitor or pinocytosis (CME) inhibitor. We found that chitosan requires pinocytosis (CME) not phagocytosis by BMMs for killing of *L. major* amastigotes at pH = 6.5 and 7.5. BMMs were infected with stationary-phase promastigotes. Some of the infected macrophages were pre-incubated with cytochalasin D (phagocytosis inhibitor) or dynasore (pinocytosis (CME) inhibitor) and exposed to various concentrations (1.64, 4.9, 14.8, 44.4, 133.3 and 400 µg/ml) of chitosan for 4 h and 24 h, followed by microscopic counting of the number of infected macrophages. There was no significant difference in the activity of HMW chitosan after inhibition of phagocytosis (p >0.05 by t-test). While a significant inhibition of chitosan mediated parasite killing occurred in the presence of dynasore at two pH values (p >0.05 by t-test). Values are expressed as % inhibition of infection relative to untreated controls. Experiment was conducted in quadruplicate cultures, data expressed as mean +/- SD>. Experiment was reproduced a further two times and confirmed the results (data not shown). *Initial macrophage infection rate was >80% after 24 h.
Table 2.14. Phagocytosis and pinocytosis by *L. major* infected BMMs in the presence of the uptake inhibitors

<table>
<thead>
<tr>
<th>Time/Hour</th>
<th>Number of latex beads ± SD *10^5/mg protein</th>
<th>Concentration of dextran ± SD µg/mg protein</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Without cytochalasin D</td>
<td>With cytochalasin D</td>
</tr>
<tr>
<td>4</td>
<td>108 ± 8</td>
<td>6 ± 1</td>
</tr>
<tr>
<td>24</td>
<td>456 ± 30</td>
<td>73 ± 8</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate, data is expressed as mean +/- SD (experiment was reproduced a further two times with confirmed similar data (not shown).

2.9.9. Fluorescence microscopy of the uptake of chitosan by macrophages

Rhodamine-labelled chitosan was used to track the delivery of chitosan to the parasitophorous vacuole (PV) of *Leishmania* infected macrophages. Fig 2.15 illustrates the cellular uptake of chitosan by *L. major*-GFP- or *L. mexicana*-GFP-infected BMMs after 4 h and 24 h rhodamine-labelled chitosan exposure. There was co-localization of chitosan and intracellular amastigotes after 4 h and 24 h with nMDP colour index 0.7 and 1 respectively (see nMDP material and methods). The uptake of chitosan increased in a time-dependent manner. Fig 2.15 (Panels D and E) shows this uptake after 4 h and 24 h respectively, and the accumulation of chitosan in PVs (shown as yellow that indicates co-localization of rhodamine and GFP). Fig 2.15 (Panel F) also shows that the inhibition of pinocytosis (CME) with dynasore prevented the uptake of chitosan with a negative nMDP colour index that represents no co-localization of chitosan and amastigotes. This is also supporting evidence for the uptake by pinocytosis as seen in Fig 2.14.
Figure 2.15. Fluorescence microscopy images of the cellular uptake of rhodamine-labelled chitosan over 4 and 24 h at pH=6.5 by BMMs infected with *L. major* (XA) or with *L. mexicana* (XB). Green represents intracellular amastigotes, red represents labelled chitosan and yellow represents merged red chitosan and green *Leishmania*. Panels A-F represent the following: Infected BMMs unexposed to chitosan after 4 h (panel A) or 24 h (panel B); Infected BMMs exposed to chitosan after 4 h (panel D) or 24 h (panel E); Infected BMMs unexposed to chitosan after 24 h (panel C) and Infected BMMs exposed to chitosan and pinocytosis inhibitor (dynasore) after 24 h (panel F)
2.10. Discussion

The literature on the anti-leishmanial activity of chitosan and its derivatives is limited, especially pertaining to its mechanism of action (124, 214, 215). In this study, we assessed the anti-leishmanial activity of various forms of chitosan, including low, medium and high molecular weight chitosan, and chitosan derivatives. Chitosan derivatives are generally produced by chemical modification of the amino or hydroxyl groups of chitosan for the optimization of the physicochemical properties. We found that chitosan and its derivatives had minimal cytotoxicity against KB-cells with LD$_{50}$ values $\geq$700 $\mu$g/ml and other macrophages (PEMs, BMMs and THP-1) at pH 7.5 or 6.5. This data supports previous reports of chitosan’s low cytotoxicity against CCRF-CEM (human lymphoblastic leukaemia) and L132 (human embryonic lung) cells with similar LD$_{50}$ values (173).

We determined that a lower pH 6.5, compared to 7.5, enhanced by 7-20 times the anti-leishmanial activity of chitosan and its derivatives against $L$. major and $L$. mexicana promastigotes and amastigotes. This higher activity of chitosan at the lower pH 6.5 could be due to its greater ionisation (protonation of the amino groups; PK$_a$ of chitosan=6.3). The greater positive charge could increase the chitosan antimicrobial activity by interacting with the negatively charged microbial membrane – in accordance with the first postulated mechanism of antimicrobial activity (124, 174). A higher chitosan activity at lower pH (pH $\approx$ 5) has previously been reported against Escherichia coli and Salmonella typhimurium (216, 217). Our study is the first to show the pH dependence of the anti-leishmanial activity of chitosan and its derivatives and could explain why literature reports of the anti-leishmanial activity of chitosan have shown such variability, with EC$_{50}$ values ranging from 70 to 240 $\mu$g/ml against $L$. infantum, $L$. amazonensis and $L$. chagasi promastigotes and amastigotes (122, 127, 129, 191, 218, 219). For example, in one study, the EC$_{50}$ of chitosan against $L$. infantum amastigotes (in PEMs) in RPMI 1640 medium was 100.81 $\mu$g/ml, but the pH at which the experiment was conducted was not mentioned (127). Moreover, Malli et al (2019) reported that chitosan solution (LMW) showed no activity until 100 $\mu$g/ml against $L$. major
promastigotes or amastigotes without mentioning the pH of the experiment (220).

Influence of pH was also seen when the anti-leishmanial activity of chitosan (of the different molecular weights) and chitosan derivatives were compared. While the different chitosans and derivatives showed minor differences in their anti-leishmanial activity at pH 7.5, the derivatives were 3 to 5 times less active than the HMW, MMW, LMW and fungal chitosan at lower pH 6.5. This reduced activity could be due to the lower number of amino groups on the chitosan derivatives (see Fig 2.3). These derivatives are more soluble at a higher pH and have similar activity to chitosan, but at a lower pH the higher protonation of the chitosan improves the anti-leishmanial activity significantly (221, 222). Carboxymethyl chitosan had no anti-leishmanial activity - most of the amino groups on this derivative have been substituted by carboxymethyl moieties making the molecule negatively charged (223).

HMW, MMW, LMW and fungal chitosan have a wide range of molecular weights. To allow like-for-like comparison, EC50 values were recalculated in terms of molarity using estimated molecular weights (HMW: MW= 342.5 KDa, MMW: MW=250 KDa, LMW: MW= 120 KDa and fungal chitosan MW=130 KDa) at pH = 6.5. Based on molarity (Table 2.7 and 2.8), HMW chitosan was significantly more active against *L. major* and *L. mexicana* promastigotes and amastigotes and further studies were conducted using HMW chitosan. The higher anti-leishmanial activity of HMW chitosan compared to MMW and LMW chitosan mirrors its greater antibacterial activity in another study against *Escherichia coli*, *Pseudomonas aeruginosa* and *Staphylococcus aureus* (224). HMW has a long chain, and therefore more glucosamine units, and possesses more amino groups (Fig 2.3) resulting in more protonated groups (-NH3+) than MMW and LMW (224) which could explain its greater potency.

We also showed that the anti-leishmanial activity of chitosan is significantly greater against *L. major* infected PEMs or BMMs compared to differentiated THP-1 cells in the order PEMs>BMMs>THP-1 cells underlining the need to take the host cell into consideration when conducting similar experiments.
In order to understand the potential anti-amastigote mechanism(s) of chitosan, we investigated whether the activity of HMW chitosan against the intracellular amastigotes was via direct uptake into the host cell and localisation in the parasitophorous vacuole or indirectly via the activation of M1 macrophages, given that the cellular immune responses in cutaneous leishmaniasis play a critical role in self-cure (225, 226).

The activation of M1 macrophages by Th1 lymphocyte subpopulation, which produces different cytokines, primarily IFN-γ and TNF-α is crucial for the elimination of the intracellular Leishmania via the triggering of an oxidative burst and therefore, the host cells increase the production of ROS and NO which are responsible for killing of the parasite (38, 39). We found that HMW chitosan stimulated TNF-α production by macrophages and this would be expected to be an indicator of an M1 macrophage that would have greater leishmanicidal activity. Our results show that chitosan stimulated BMMs, PEMs and THP-1 ROS production with a peak after 4 h and led to a significant increase in the TNF-α and NO production after 24 h in a bell-shaped response. Similar findings have been reported showing that HMW chitosan had in vitro stimulatory effect on PEMs (from male rats) NO production (126) and LMW chitosan stimulated RAW264.7 macrophage TNF-α production (184). Another study demonstrated that LMW chitosan induced ROS production in an epithelial, human breast cancer cell line (227). The bell-shaped responses are consistent with a study that showed that chitosan stimulated NO and TNF-α production in peritoneal macrophages in a dose-dependent manner and their levels tended to decrease at higher concentrations of chitosan (320μg/ml) (228). This type of response has also been reported previously for tucaresol for both its immunomodulatory and activity against experimental *L. donovani* infections, albeit at lower doses (229). We found that BMMs had high levels expression of TNF-α, NO and ROS and this could be explained as BMMs are more homogenous than PEMs, and they are characterised with their high yield, homogeneity and long lifespan (230).

BMMs were chosen to evaluate if the anti-leishmanial activity of HMW chitosan is through indirect way (through the immunostimulatory effects) or direct way (by the uptake of chitosan by macrophages) or both of them, as these
macrophages are more homogenous than PEMs cells (203). Despite the observed chitosan-induced ROS and NO production there was no evidence that this contributed to the anti-leishmanial activity in our study – the inhibitors we used to mitigate their production had no effect on the ability of chitosan to kill intracellular Leishmania amastigotes (Figs 2.11 and 2.13). This led us to investigate the cellular uptake of HMW chitosan and its relationship to the anti-leishmanial activity.

The uptake of the large charged molecule HMW chitosan has not been systematically studied before and there is no clear evidence of its penetrating cell membranes or of its uptake mechanism. Macrophages are known to take up extracellular materials and plasma by endocytosis. Endocytosis mainly occurs via two different cellular uptake mechanisms: pinocytosis or phagocytosis, where pinocytosis is fluid-phase endocytosis and phagocytosis is the process of engulfing large particles (231). Inhibition of pinocytosis (CME) significantly reduced the anti-leishmanial activity of HMW chitosan. Therefore, in our study pinocytosis (CME) was considered to be the main mechanism for the uptake of HMW chitosan by BMMs, indicating a direct anti-leishmanial effect of this molecule against amastigotes. Other researches have also reported pinocytosis as the pathway for the uptake of chitosan of different molecular weights by HEK293 epithelial cells (232). The fluorescence imaging in our study showed that in BMMs HMW chitosan is taken up into the parasitophorous vacuole (PV) where the Leishmania amastigotes reside, with the labelled chitosan being internalized within 4 h and increasing up to 24 h later. This scenario is consistent with another study where rhodamine isothiocyanate- chitosan (RITC-chitosan 98-10 K) was found to be directly delivered to the U937 macrophage lysosome after 24 h (233). The accumulation of chitosan in the PV might be due to chitosan's relatively high pKa 6.3, making it more soluble and protonated in the acidic contents of the vacuole. This is consistent with a study using bafilomycin to inhibit acidification and prevent chitosan accumulation within macrophages (233).

In summary, our studies indicate that chitosan and its water-soluble derivatives showed anti-leishmanial activity against both L. major and L. mexicana promastigotes and amastigotes in a pH-dependent manner. At pH 6.5 HMW
chitosan is more active than MMW and LMW chitosan and chitosan derivatives, in particular those where the amino groups are substituted. In addition, HMW chitosan activated M1 macrophages, stimulating them to produce NO and ROS. However, the anti-leishmanial activity of chitosan was not due to such immune activation, as an NO inhibitor and a ROS scavenger failed to reduce the anti-leishmanial activity. Instead, the anti-leishmanial activity was related to direct uptake of chitosan into the parasitophorous vacuole by pinocytosis (CME). HMW chitosan demonstrated effective in vitro anti-leishmanial activity with minimal cytotoxicity and future work will focus on in vivo studies, formulations and routes for drug administration.
3. Preparation and characterisation of amphotericin B loaded chitosan nanoparticles

3.1. Introduction

As mentioned in Chapter 1, the polyene antibiotic AmB (a standard treatment for systemic fungal infections) was classified as a second line treatment for VL and MCL, particularly for pentavalent antimonial resistant *Leishmania*. However, the toxic side effects of AmB restrict its use. Great efforts have been spent to develop drug delivery systems (DDs) of AmB, to reduce its toxicity and improve the efficacy of the drug, such as AmBisome®, a liposomal formulation of AmB, which is significantly less toxic than the free drug and is effective against VL and CL and then has been promoted as first line for VL in the Indian subcontinent (ISC). However, the drawbacks are (i) high cost, where donated free of charge by WHO for VL, not for CL and (ii) need for cold chain due to stability guaranteed only up to 25°C (54, 55, 56, 58, 234). Polymeric nanoparticles technology has also gained a great interest in the DDs field, giving opportunities for controlled drug release, drug protection of enzymatic degradation and retention period of drug. We mentioned in Chapter 1 that chitosan nanoparticles are gaining a lot of attention in DDs in the medical field as they are both biodegradable and biocompatible (119). There are different methods for the preparation of chitosan nanoparticles and they are summarised in Fig 3.1.
Figure 3.1. Preparation methods of chitosan nanoparticles. (A) Emulsion cross-linking in which chitosan is stabilized by a surfactant and then is emulsified in an oil phase (water-in-oil emulsion) such as chitosan aqueous solution in toluene, using Span 80® as emulsifier and is then reacted with an appropriate cross linking agent (e.g. formaldehyde, glutaraldehyde, genipin, glyoxal etc.) followed by washing and drying of the nanoparticles, (B) Ionotropic gelation which represents the method we used and will be described in details, (C) Emulsion-droplet coalescence in which chitosan solution is dispersed in liquid paraffin oil to prepare an emulsion and then sodium hydroxide solution is added to the first emulsion under high speed mixing which produces nanoparticles which are centrifuged and dried, (D) Precipitation in which a compressed air nozzle is used to inject chitosan solution into basic organic solvent (sodium hydroxide, NaOH methanol or ethanediamine), (E) Reverse micelles in which a surfactant (e.g. sodium 10 bis (ethyl hexyl) sulfosuccinate or cetyl trimethylammonium bromide) is dissolved into an organic solvent (e.g. n-hexane) to which aqueous chitosan solution is added under continuous stirring. Subsequently, a cross-linking agent (e.g. glutaraldehyde) is added and maintained under stirring overnight, and the organic solvent is removed by evaporation(F) spray drying in which an aqueous acetic acid solution of chitosan is prepared then, drugs are suspended or dissolved in the chitosan solution and then a cross-linking agent (glutaraldehyde or sodium tripolyphosphate. Small droplets are formed upon the atomization and the formation of flowing particles with evaporation of solvent. These techniques except ionotropic gelation frequently require the use of organic solvents or heat, which are undesirable steps and may affect encapsulated drug and may increase cytotoxicity effects (171, 235, 236)

The ionotropic gelation method is described as an easy and simple technique in which, nanoparticles are formed by an electrostatic interaction between the cationic amino groups of chitosan and negatively charged anions of other compounds (such as tripolyphosphate sodium (TPP), dextran sulphate, chondroitin sulphate, etc) with mechanical stirring at room temperature leading to spherical nanoparticles. The use of different pH values of media and ratios
of chitosan and polyanions can result in the synthesis of particles at different sizes and surface charges. This technique has many advantages such as the usage of aqueous condition, low toxicity and not changing the chemistry of the encapsulated drug (120, 237). Moreover, these nanoparticles can be prepared in small and different sizes and charges, they can be used for different routes of administration and offer a sustained drug release (112).

All nanoparticles used in our study were prepared using the inotropic gelation method. Chitosan nanoparticles were prepared via the interaction between the oppositely charged groups of chitosan (positive amino groups) and TPP (polyanions) or dextran sulphate (negative groups) (Fig 3.2, give structure of TPP and dextran sulphate). Dextran sulphate is a biodegradable and biocompatible polysaccharide with a negative charge and is soluble in water. These properties enable dextran sulphate to produce nanoparticles when interacts with positively charged molecules to give positively or negatively charged nanoparticles according to the mass ratios used (238). Because of the biodegradability, biocompatibility and the possibility of dextran to interact with chitosan to produce negative charged nanoparticles, we chose dextran sulphate as a cross-linker.

Tripolyphosphate sodium (TPP) is a popular and commonly used polyanion to prepare chitosan nanoparticles because of its safety (TPP is approved as safe by the FDA; Sec. 182.1810 sodium tripolyphosphate (239)) and gelation properties and furthermore, TPP has a role in the stability of nanoparticles (122, 240). Because of the safety profile and the ability of TPP to interact with chitosan to produce positive charged nanoparticles, we chose TPP as another cross-linker.

There are several possible mechanisms for drug (AmB) release from chitosan nanoparticles as shown in Fig 3.3 and chitosan nanoparticles show a pH-dependent drug release because of its solubility. Therefore, the aims of this chapter were (i) to produce two types of chitosan nanoparticles containing AmB, one by using TPP to obtain positively charged nanoparticles and the other with dextran sulphate to obtain negatively charged nanoparticles, both with smallest possible sizes. After optimizing the preparation parameters, the
aims were (ii) to characterise the produced blank and AmB loaded nanoparticles in terms, of size, charge, morphology and stability and (iii) to evaluate the amphotericin B loading and drug release from the amphotericin B loaded chitosan TPP or dextran sulphate nanoparticle.

Figure 3.2. Chemical structure of TPP and dextran sulphate (241)

Figure 3.3. Mechanisms of drug release from chitosan nanoparticles. a) In diffusion release, a permeation of the drug is happening through the interior of the matrix of polymer to the near medium, b) in the swelling release, an absorption of water into the polymer is occurred until the dissolving of polymer, c): erosion release which can be homogenous (at the same rate throughout the matrix) and heterogeneous (erosion of the polymer from the surface towards the inner core). Polymer degradation may be due to the surrounding media or the presence of enzymes. (120)
3.2. Material and methods

3.2.1. Preparation of blank chitosan nanoparticles

Nanoparticles were prepared by inotropic gelation by mixing positively charged chitosan with negatively charged TPP or dextran sulphate as shown schematically in Fig 3.4.

1- HMW chitosan (MW=310-375 KDa, Sigma, UK) was dissolved at various concentrations (0.33, 1, 3 and 9 mg/ml) in 1% aqueous acetic solution (Sigma, UK). The pH of the resulting chitosan solution was adjusted to pH of 3, 4, 5, 6 and 7 by adding NaOH solution (Sigma, UK) and this enabled investigation into the influence of pH on particles formation.

2- The sodium tripolyphosphate (TPP, Mw= 367.85 g/mol, Fisher scientific, US) and dextran sulphate (Mw= 40 kDa, Sigma, UK) solutions were prepared by dissolving TPP or dextran sulphate in double-distilled water at various concentrations

3- The nanoparticles were formed at chitosan: TPP or chitosan: dextran sulphate mass ratios of 3:1, 5:1, 10:1, 20:1, 1:1, 1:3, 1:5, 1:10 and 1:20). TPP or dextran sulphate aqueous solution (10 ml) was added dropwise using a 10 ml syringe into the chitosan solution (10ml) under magnetic stirring (Fig 3.4). Directly after adding the TPP or dextran sulphate solution, the nanoparticles suspension was sonicated to reduce the particles size by using a probe sonicator Soniprep 150 (Richmond Scientific Ltd, Lancashire, UK); the diameter of the microprobe was 3mm, operating at an output frequency of 23kHz with an amplitude of 14-16 nm for 15 mins (15 mins was found to be the optimal time after testing for 1, 5, 15 and 20 min) with 1 min rest after every 5 min of sonication to decrease possible overheating of the sample and resulting degradation of the AmB. Subsequently, the nanoparticle suspension
was filtered through a 0.2 µm size syringe filter (Millex, Merck Millipore, UK) to remove aggregates and larger particles. The nanoparticles were concentrated by centrifugation (8,000 x g) using high recovery centrifugal filters (Spin-X UF concentrators, 20 ml, 30 kDa, Corning, UK).

4- The nanoparticles suspension was analysed directly by using a Zetaziser (Malvern Instruments Ltd., UK) to determine the size, polydispersity index (PDI) and zeta potential of the nanoparticles. Nanoparticles were then lyophilised using a freeze dryer (Micro Modulyo, Richmond Scientific, UK). In this process, D-mannitol (Mw=182.17 g/mol, Sigma, UK) 5% or sucrose (Mw=342.3 g/mol, Sigma, UK) 5% v/v was used as a cryoprotectant to protect the nanoparticles from the freezing and desiccation stresses (the stress of freezing and dehydration) (242). After 48 hours, lyophilized nanoparticles were collected, weighed and stored at 4°C for further analysis. The lyophilized blank nanoparticles were white cotton-like substance.

3.2.2. Preparation of AmB loaded chitosan nanoparticles

The optimal parameters determined for producing blank nanoparticles which gave the smallest sizes and PDI (which refers to homogeneity of nanoparticle size (243)) were chosen to prepare the AmB loaded chitosan nanoparticles (Fig 3.4).

1- 10 mg of AmB (Purity ≥ 95%, Cambridge Bioscience, UK) was dissolved in 0.5 ml of DMSO (high-performance liquid chromatography grade; Fisher Chemical, United Kingdom) and sonicated in a Camlab TransSonic T460/H water bath for 15 min at room temperature.

- AmB is insoluble in water at pH 6 to 7. It is soluble in DMSO (30–40 mg/ml) and in dimethylformamide (2–4 mg/ml). Molecular weight of AmB is 924.08 g/mol and logP is -0.66.

2- To prepare AmB loaded chitosan-TPP nanoparticles, AmB solution (0.5 ml of 10 mg) was added to 10 ml of TPP solution (6 mg in 10 ml distilled water) and this solution was added dropwise to 10 ml of HMW
chitosan solution (30 mg in 10 ml AC 1%) of pH of 5 under magnetic stirring.

3- To prepare AmB loaded chitosan-dextran sulphate nanoparticles, AmB solution (0.5 ml of 10 mg) was added to 10 ml of dextran solution (30 mg in 10 ml double distilled water) and this solution was added dropwise to 10 ml of HMW chitosan solution (10 mg in 10 ml AC 1%) of pH of 5 under magnetic stirring.

4- Subsequently, nanoparticle suspension was sonicated directly after adding the gelation material, filtered, purified and freeze dried (using a cryoprotectant) as described for blank nanoparticles in section 3.3.1. The lyophilised AmB loaded nanoparticles were yellow cotton-like material. Each experiment was repeated three times.
3.2.3. Physicochemical properties of the nanoparticles (size, charge and morphology)

1- The size of the nanoparticles was measured by dynamic light scattering (DLS) using a Zetaziser (Malvern Instruments Ltd., UK) with the following parameters: dispersant: water, dispersant refractive indices (RI): 1.33, viscosity (cP): 0.8872, material RI: 1.33, temperature (°C): 25.0, measurement position (mm): 3 and attenuator: 9. The result is expressed as Z-Average (nm) and polydispersity index (PDI). Z-average reflects the intensity weighted mean hydrodynamic size of the particles measured by DLS. PDI represents the distribution of the nanoparticles sizes in the sample (243). DLS is identified as technique for measuring the size and size distribution of molecules and particles which are dispersed or dissolved in liquid and measures hydrodynamic diameter based on the light dispersion properties of samples Tyndall effect (light scattering) and Brownian motion (the random motion of particles suspended in a fluid because of the bombardment by the solvent molecules that surround them). DLS gives the PDI value which reflects the size distribution of the nanoparticles which is classified to monomodal (one population) or plurimodal (several populations) and monodisperse (narrow distribution) or polydisperse (broad distribution) assuming that lower PDI less than 0.4 refers to a homogenous population and 0.1 or less to higher homogeneity in the particle population (246, 247).

2- Zeta-potential (representing surface charge of nanoparticles) of the nanoparticles was measured by the Zetaziser with same parameters for the sizing except for measurement position being (mm): 2 mm and attenuator: 11. Zetaziser measures the zeta potential through the monitoring of the mobility of charged particles on the surface of the nanoparticles by application of an electrical potential (248). Data analysis was performed using the Malvern ZetaSizer software.
Measurements were repeated three times for sizes and 6 times for zeta-potential for each sample.

3- The morphology of the nanoparticles was examined using a scanning electron microscopy (SEM, UCL, School of Pharmacy) and a transmission electron microscopy (TEM, UCL, School of Pharmacy). For the SEM, a fragment of sample was attached to a self-adhesive carbon disc mounted on a 25 mm aluminium stub. The stub was coated with 25 nm of gold using a sputter coater. The stub was then placed into a FEI Quanta 200 FEG SEM for imaging at 5kV accelerating voltage using secondary electron detection (249).

Liquid samples for TEM were dropped with a Pasteur pipette onto a copper grid coated with a carbon/formvar support film. After 15 seconds, a filter paper was blotted off to remove the excess sample. Then a drop of negative stain (1% uranyl acetate) was added and blotted after 15 seconds. The grid was placed into a specimen holder and inserted into a Phillips/FEI CM 120 BioTwin TEM for imaging at 200kV (250).

3.2.4. Stability of nanoparticles regarding size and zeta potential

This stability was evaluated by keeping nanoparticles in distilled water, PBS or RPMI (pH 7.5 or pH 5) and in mouse (BALB/c) plasma (pooled female, BioIVT, UK) in rubber-capped glass vials at temperatures of 4, 34 or 37 °C for 30 days. Particle sizes and zeta-potential were measured after 0, 1, 7 and 30 days.

Stability of dried nanoparticles was identified by resuspending them in water after 0, 1, 7 and 30 days and measuring their size and charge and they were highly stable.

3.2.5. Determination of drug encapsulation efficiency and AmB loading and release

Nanoparticles were prepared as described in section 4-2-2. Following sonication of the suspensions in the probe sonicator and filtration, the AmB
loaded nanoparticle suspension was centrifuged (8,000 x g) to remove the free AmB by using High recovery centrifugal filters (Spin-X UF concentrators, 30 kDa, Corning, UK). Filtrates and supernatants were collected and analysed for AmB concentrations by HPLC as described in section 3.3.6.1. Because of the molecular weight cut-off of the filtration tubes, only the free drug could pass through the membrane. The encapsulation efficiency (EE), drug loading (122) and yield (118) were calculated using the following equations:

\[
\text{EE}\% = \frac{\text{Mass of total AmB} - \text{mass of free AmB}}{\text{Mass of total AmB}} \times 100
\]

\[
\text{Drug loading}\% = \frac{\text{Mass of total AmB} - \text{mass of free AmB}}{\text{Mass of chitosan + dextran sulphate or TPP}} \times 100
\]

\[
\text{Yield}\% = \frac{\text{Mass of nanoparticles after freeze-drying}}{\text{Mass of AmB + chitosan+ dextran sulphate or TPP}} \times 100
\]

Additionally, the AmB loading was evaluated again after freeze drying by dissolving the yielded yellow powder in DMSO, in acidic pH 3 (by using 1% (v/v) acetic acid), and then by measuring the quantity of AmB by HPLC as described previously in Chapter 3, in section 3.3.6.1. There was no significant difference in the loading value between these two methods, and the data in the thesis will be expressed according to the first method (using High recovery centrifugal filters).

3.2.6. *In vitro* release of AmB

The release of AmB from chitosan-TPP or chitosan-dextran sulphate nanoparticles was evaluated by the dialysis method. One ml of the nanoparticles suspension (1 mg/ml AmB equivalent prepared in double distilled water) was added to either one ml of PBS containing 5% DMSO or
one ml of mouse (BALB/c) plasma (pooled female, BioIVT, UK) containing DMSO (5%) (for the solubility of AmB) (118). Subsequently, this 2 ml was placed in a dialysis bag (molecular mass cut off =12−14 kDa, Sigma, UK) and dialyzed against 50 ml of PBS containing 5% DMSO at pH of 7.5, 6.5 or 5. After immersing the dialysis bag in the release medium, the dialysis set up was left under stirring at 4, 34 or 37 °C for 168 h. The temperatures 4, 34 and 37 °C were chosen to mimic the storage, skin and body temperatures respectively, while pH 5 was chosen to simulate the release in the endosomal compartment of macrophages, pH 7.4 to simulate physiological conditions (251) and pH 6.5 to mimic our in vitro study (anti-leishmanial activity) conditions.

After 6, 24, 48, 72, 96,120, 144 and 168 h the total dialysis medium was replaced with fresh medium to avoid saturation of AmB, (maintaining strict sink conditions throughout the experiment). Release media was processed to quantify the released AmB using HPLC as described in section 3.3.6.1. The results were expressed as a cumulative percentage release of the total amount of AmB (%w/w) versus time according to the equation.

\[
\text{Cumulative release (\%) = } \frac{\text{Mass of released AmB at time } t \text{ (mg)}}{\text{Mass of total AmB (1 mg)}} \times 100
\]

Mass of released AmB at time \( t \) is a cumulative amount. For instance, mass of released AmB after 48h is the total amount released at 6, 24 and 48 h.

3.2.6.1. Quantification of AmB by HPLC

AmB was analysed by using a 1260 Infinity Agilent HPLC system. The column and settings used in our study are summarized in Table 3.1 (252). A stock solution of AmB was prepared by dissolving 1 mg of AmB in DMSO. Standard solutions were achieved by diluting this stock solution in PBS containing 5% DMSO.
### Table 3.1. HPLC parameters for AmB quantification

<table>
<thead>
<tr>
<th>HPLC column</th>
<th>Injection volume (μL)</th>
<th>Flow rate (ml/min)</th>
<th>Mobile phase</th>
<th>Detector wavelength nm</th>
<th>Retention time (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenomenex; Synergi–Hydro RP (250x4.6 mm; 5 μm)</td>
<td>20</td>
<td>1</td>
<td>5mM EDTA•2Na in methanol</td>
<td>450</td>
<td>7.65</td>
</tr>
</tbody>
</table>

### 3.3. Results

#### 3.3.1. Effects of the initial concentration of chitosan and sonication time on the quality of the nanoparticles

#### 3.3.1.1. Conditions that resulted in poor quality nanoparticles

Precipitation and poor quality of both types of nanoparticles were shown at pH values of 7 and 3 of chitosan solution at all tested conditions. Chitosan solutions at concentrations (0.3 or 9 mg/ml), at all tested pH with different mass ratios and after sonication of the nanoparticles suspension for 1, 5, 15 or 20 mins, gave poor quality nanoparticles with (high PDI>0.8) and with different peaks as seen in Table 3.2. Similarly, chitosan solutions at concentrations (1 or 3 mg/ml) at all pH values with different mass ratios and after sonication of the nanoparticles suspension for 1 or 5 mins produced poor quality nanoparticles (Table 3.2). Finally, chitosan solutions at concentrations (1 or 3 mg/ml) at all pH values with a mass ratio between chitosan and TPP (20:1, 1:1, 1/3, 1:5, 1:10 or 1:20) or a mass ratio between chitosan and dextran sulphate (1:5, 1:10, 1:20, 10:1 or 20:1) and after sonication for 1, 5, 15 or 20 mins caused a precipitation of particles or poor quality nanoparticles with a high PDI of 1.
Table 3.2. Conditions which did not produce good quality nanoparticles

<table>
<thead>
<tr>
<th>Chitosan mg/ml</th>
<th>pH</th>
<th>Chitosan: TPP or chitosan: dextran sulphate mass ratio</th>
<th>Sonication time mins</th>
<th>Resulted nanoparticles</th>
<th>Related figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.3</td>
<td>3, 4, 5, 6 or 7</td>
<td>3:1, 5:1, 10:1, 20:1, 1:1, 1:3, 1:5, 1:10 and 20:1</td>
<td>1, 5, 15 or 20</td>
<td>Poor quality nanoparticles, had a high polydispersity regarding sizes. These samples were deemed not suitable for further study as they might contain large particles or aggregates</td>
<td>3.5, a)</td>
</tr>
<tr>
<td>1</td>
<td>3 or 7</td>
<td>3:1, 5:1, 10:1, 20:1, 1:1, 1:3, 1:5, 1:10 and 20:1</td>
<td>1, 5, 15 or 20</td>
<td>Poor quality nanoparticles, had a high polydispersity regarding sizes. These samples were deemed not suitable for further study as they might contain large particles or aggregates</td>
<td></td>
</tr>
<tr>
<td>4, 5 or 6</td>
<td>3:1, 5:1, 10:1, 20:1, 1:1, 1:3, 1:5, 1:10 and 20:1</td>
<td>1 or 5</td>
<td>Poor quality nanoparticles with high PDI and very large nanoparticles with size≈ 800nm</td>
<td>3.5, b)</td>
<td></td>
</tr>
<tr>
<td>4, 5 or 6</td>
<td>Chitosan: TPP (20:1, 1:1, 1:3, 1:5, 1:10 or 1:20) Chitosan: dextran sulphate (1:5, 1:10, 1:20, 1:10 or 1:20)</td>
<td>15 or 20</td>
<td>A precipitation of particles or poor quality nanoparticles with high PDI of 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>3 or 7</td>
<td>3:1, 5:1, 10:1, 20:1, 1:1, 1:3, 1:5, 1:10 and 20:1</td>
<td>1, 5, 15 or 20</td>
<td>Poor quality nanoparticles, had a high polydispersity regarding sizes. These samples were deemed not suitable for further study as they might contain large particles or aggregates</td>
<td></td>
</tr>
<tr>
<td>4, 5 or 6</td>
<td>3:1, 5:1, 10:1, 20:1, 1:1, 1:3, 1:5, 1:10 and 20:1</td>
<td>1 or 5</td>
<td>Poor quality nanoparticles with high PDI and very large nanoparticles with size≈ 800nm</td>
<td>3.5, b)</td>
<td></td>
</tr>
<tr>
<td>4, 5 or 6</td>
<td>Chitosan: TPP (20:1, 1:1, 1:3, 1:5, 1:10 or 1:20) Chitosan: dextran sulphate (1:5, 1:10, 1:20, 1:10 or 1:20)</td>
<td>15 or 20</td>
<td>A precipitation of particles or poor quality nanoparticles with high PDI of 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>3, 4, 5, 6 or 7</td>
<td>3:1, 5:1, 10:1, 20:1, 1:1, 1:3, 1:5, 1:10 and 20:1</td>
<td>1, 5, 15 or 20</td>
<td>Poor quality nanoparticles, had a high polydispersity regarding sizes. These samples were deemed not suitable for further study as they might contain large particles or aggregates</td>
<td>3.5, c)</td>
</tr>
</tbody>
</table>

Experiment was reproduced further two times with confirmed similar data
**Figure 3.5.** Poor quality nanoparticles at different conditions. a) at initial chitosan concentration 0.3 mg/ml and all other different parameters, b) at initial chitosan concentration 1 or 3 mg/ml and sonication for 1 or 5 mins and c) at initial chitosan concentration 9 mg/ml other different parameters. There are several populations of particles, some are small in size e.g. less than 100 nm and some are large about 1 µm. Each colour represents one measurement as each sample was measured 3 times.
3.3.1.2. Conditions that resulted in good quality nanoparticles

Chitosan solutions at concentrations (1 or 3 mg/ml) at pH (4, 5 or 6) with a mass ratio between chitosan and TPP (3:1, 5:1 or 10:1) or a mass ratio between chitosan and dextran sulphate (1:3, 1:1, 3:1 or 5:1) and after sonication of the nanoparticles suspension for 15 mins gave good quality nanoparticles with (low PDI<0.4) (lower PDI means more homogenous and stable nanoparticles (210)) with one peak; with different Z-Averages according to the conditions that would be discussed later (Fig 3.6 and Fig 3.7). Sonication of the nanoparticles suspension for more than 15 mins (for example 20 mins) produced no significant changes in the quality (PDI) or physicochemical properties (sizes and charges) of the nanoparticles.

![Size Distribution by Intensity](image)

**Figure 3.6.** Good quality chitosan-dextran sulphate nanoparticles with one peak (one population of nanoparticles at initial chitosan concentration 3 mg/ml and sonication for 15 mins. Chitosan-dextran sulphate nanoparticles (Size = 145.8 nm, PDI =0.2). Each colour represents one measurement as each sample was done in three measurements.
Our results indicated that the optimal parameters to obtain good quality nanoparticles (for both types of nanoparticles with TPP or with dextran sulphate) are: initial concentration of chitosan with 1 or 3 mg/ml and sonication time of the nanoparticles suspension for 15 mins as sonicating for more than 15 mins gave same results regarding quality (PDI), size and charge.

3.3.2. Effects of pH of chitosan solution and the mass ratio on the size and charge of good quality nanoparticles

Chitosan and TPP with parameters (chitosan 3 mg/ml at pH 5 and TPP 0.6 mg/ml) produced the smallest and most quality nanoparticles of chitosan-TPP nanoparticles (with lowest PDI, homogenous suspension) with size 48 ± 6 nm, PDI = 0.1 ± 0.03 and positive charge (zeta potential = 32.1 ± 1.2 mv) (Table 3.3). However, chitosan and dextran sulphate with parameters (chitosan 1 mg/ml at pH 5 and dextran sulphate 3 mg/ml) produced the smallest and most quality nanoparticles of chitosan-dextran sulphate nanoparticles with size 145 ± 6 nm, PDI = 0.1± 0.05 and negative charge (zeta potential = -15.5 ± 1mv) (Table 3.4).
Table 3.3. Effect of pH and concentration of chitosan and mass ratio of the reactants on the physicochemical properties of blank chitosan-TPP nanoparticles

<table>
<thead>
<tr>
<th>Chitosan mg/ml</th>
<th>TPP μM</th>
<th>Ch:TPP mass ratio</th>
<th>Ch:TPP Molarity ratio</th>
<th>pH</th>
<th>Particle sizes nm</th>
<th>PDI</th>
<th>Zeta potential mv</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.002</td>
<td>0.3 0.81 3:1</td>
<td>1:405</td>
<td>4</td>
<td>100 ± 9</td>
<td>0.3 ± 0.1</td>
<td>17 ± 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.2 0.53 5:1</td>
<td>1:265</td>
<td>5</td>
<td>120 ± 8</td>
<td>0.3 ± 0.1</td>
<td>18.9 ± 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.1 0.27 10:1</td>
<td>1:135</td>
<td>4</td>
<td>170 ± 9</td>
<td>0.3 ± 0.1</td>
<td>19.8 ± 0.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.3 0.81 3:1</td>
<td>1:405</td>
<td>5</td>
<td>95 ± 10</td>
<td>0.3 ± 0.1</td>
<td>13.5 ± 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.2 0.53 5:1</td>
<td>1:265</td>
<td>6</td>
<td>108 ± 11</td>
<td>0.2 ± 0.05</td>
<td>17.5±0.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.1 0.27 10:1</td>
<td>1:135</td>
<td>6</td>
<td>169 ± 9</td>
<td>0.3 ± 0.1</td>
<td>19.2±0.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.3 0.81 3:1</td>
<td>1:405</td>
<td>5</td>
<td>135 ± 11</td>
<td>0.2 ± 0.05</td>
<td>11.2±0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.2 0.53 5:1</td>
<td>1:265</td>
<td>6</td>
<td>149 ± 12</td>
<td>0.3 ± 0.1</td>
<td>15.5±2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.1 0.27 10:1</td>
<td>1:135</td>
<td>5</td>
<td>190 ± 9</td>
<td>0.2 ± 0.04</td>
<td>17.5±0.9</td>
</tr>
<tr>
<td>3</td>
<td>0.008</td>
<td>1 2.72 3:1</td>
<td>1:340</td>
<td>4</td>
<td>141 ± 10</td>
<td>0.2 ± 0.05</td>
<td>23.9±1.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.6 1.6 5:1</td>
<td>1:200</td>
<td>4</td>
<td>99 ± 6</td>
<td>0.2 ± 0.02</td>
<td>34.3±0.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.3 0.81 10:1</td>
<td>1:101</td>
<td>5</td>
<td>220 ± 15</td>
<td>0.2 ± 0.02</td>
<td>44.8±1.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 2.72 3:1</td>
<td>1:340</td>
<td>6</td>
<td>140 ± 9</td>
<td>0.2±0.03</td>
<td>22.9±1.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.6 1.6 5:1</td>
<td>1:200</td>
<td>5</td>
<td>48 ± 6</td>
<td>0.1 ± 0.03</td>
<td>32.1±1.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.3 0.81 10:1</td>
<td>1:101</td>
<td>5</td>
<td>178 ± 12</td>
<td>0.3±0.02</td>
<td>40.2±1.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 2.72 3:1</td>
<td>1:340</td>
<td>6</td>
<td>174 ± 9</td>
<td>0.1±0.05</td>
<td>16.2±1.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.6 1.6 5:1</td>
<td>1:200</td>
<td>6</td>
<td>155 ± 8</td>
<td>0.2 ± 0.02</td>
<td>18.3±1.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.3 0.81 10:1</td>
<td>1:101</td>
<td>5</td>
<td>340 ± 19</td>
<td>0.3 ± 0.1</td>
<td>18.7±1.1</td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). The smallest size and PDI of these nanoparticles were 48 ± 6 nm and 0.1 ± 0.03 with positive surface charge (zeta potential = +32.1 ± 1.2)
Table 3.4. Effect of pH and concentration of chitosan and mass ratio of the reactants on the physicochemical properties of blank chitosan-dextran sulphate nanoparticles

<table>
<thead>
<tr>
<th>Chitosan (Ch)</th>
<th>Dextran sulphate(Dx)</th>
<th>Ch:Dx mass ratio</th>
<th>Ch:DX Molarity ratio</th>
<th>pH</th>
<th>sizes nm</th>
<th>PDI</th>
<th>Zeta potential mv</th>
</tr>
</thead>
<tbody>
<tr>
<td>mg/ml µM</td>
<td>mg/ml µM</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 0.002</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 0.075</td>
<td>1:3</td>
<td>1:37.5</td>
<td>4</td>
<td>160 ± 6</td>
<td>0.2 ± 0.05</td>
<td>-17.5 ± 1</td>
<td></td>
</tr>
<tr>
<td>1 0.025</td>
<td>1:1</td>
<td>1:12.5</td>
<td>177 ± 9</td>
<td>0.3 ± 0.1</td>
<td>-8 ± 0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3 0.007</td>
<td>3:1</td>
<td>1:3</td>
<td>190 ± 9</td>
<td>0.3 ± 0.1</td>
<td>+6 ± 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2 0.005</td>
<td>5:1</td>
<td>1:2.5</td>
<td>185 ± 8</td>
<td>0.3 ± 0.1</td>
<td>+8 ± 0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 0.075</td>
<td>1:3</td>
<td>1:37.5</td>
<td>145 ± 6</td>
<td>0.1 ± 0.05</td>
<td>-15.5 ± 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 0.025</td>
<td>1:1</td>
<td>1:12.5</td>
<td>169 ± 9</td>
<td>0.3 ± 0.1</td>
<td>-7 ± 0.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3 0.007</td>
<td>3:1</td>
<td>1:3</td>
<td>170 ± 5</td>
<td>0.2 ± 0.04</td>
<td>+4 ± 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2 0.005</td>
<td>5:1</td>
<td>1:2.5</td>
<td>185 ± 8</td>
<td>0.3 ± 0.1</td>
<td>+5 ± 0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 0.075</td>
<td>1:3</td>
<td>1:37.5</td>
<td>230 ± 6</td>
<td>0.2 ± 0.04</td>
<td>-12 ± 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 0.025</td>
<td>1:1</td>
<td>1:12.5</td>
<td>200 ± 6</td>
<td>0.4 ± 0.1</td>
<td>-6 ± 0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3 0.007</td>
<td>3:1</td>
<td>1:3</td>
<td>210 ± 7</td>
<td>0.3 ± 0.1</td>
<td>+3 ± 0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2 0.005</td>
<td>5:1</td>
<td>1:2.5</td>
<td>220 ± 5</td>
<td>0.4 ± 0.1</td>
<td>+4 ± 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9 0.225</td>
<td>1:3</td>
<td>1:28</td>
<td>340 ± 12</td>
<td>0.3 ± 0.1</td>
<td>-33±7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 0.075</td>
<td>1:1</td>
<td>1:9.3</td>
<td>307 ± 12</td>
<td>0.3 ± 0.1</td>
<td>-10 ± 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 0.025</td>
<td>3:1</td>
<td>1:3</td>
<td>332 ± 9</td>
<td>0.2 ± 0.04</td>
<td>+8 ± 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.6 0.015</td>
<td>5:1</td>
<td>1:1.8</td>
<td>303 ± 6</td>
<td>0.4 ± 0.1</td>
<td>+10 ± 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9 0.225</td>
<td>1:3</td>
<td>1:28</td>
<td>270 ± 10</td>
<td>0.2 ± 0.05</td>
<td>-35±7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 0.075</td>
<td>1:1</td>
<td>1:9.3</td>
<td>279 ± 11</td>
<td>0.3 ± 0.1</td>
<td>-15 ± 4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 0.025</td>
<td>3:1</td>
<td>1:3</td>
<td>290 ± 11</td>
<td>0.3 ± 0.1</td>
<td>+6 ± 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.6 0.015</td>
<td>5:1</td>
<td>1:1.8</td>
<td>285 ± 10</td>
<td>0.2 ± 0.05</td>
<td>+7 ± 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9 0.225</td>
<td>1:3</td>
<td>1:28</td>
<td>380 ± 10</td>
<td>0.2 ± 0.05</td>
<td>-39±7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 0.075</td>
<td>1:1</td>
<td>1:9.3</td>
<td>400 ± 11</td>
<td>0.3 ± 0.1</td>
<td>-19 ± 4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 0.025</td>
<td>3:1</td>
<td>1:3</td>
<td>380 ± 11</td>
<td>0.2 ± 0.05</td>
<td>+3 ± 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.6 0.015</td>
<td>5:1</td>
<td>1:1.8</td>
<td>450 ± 10</td>
<td>0.2 ± 0.03</td>
<td>+4 ± 1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). The smallest size and PDI of these nanoparticles were 145 ± 6 nm and 0.1 ± 0.05 with negative surface charge (zeta potential = -15.5 ± 1).
3.3.3. Effects of AmB loading and freeze-drying with and without cryoprotectants on physicochemical properties and the morphology of the nanoparticles

1- AmB loading increased the size of blank chitosan-TPP and dextran sulphate nanoparticles by 18.75% and 13% respectively. However, such loading did not cause any significant change to the zeta potential and PDI of both types of nanoparticles (Table 3.5) (p > 0.05 by t-test). Freeze drying process without the use of a cryoprotectant (sucrose or D-mannitol) resulted in poor quality nanoparticles with various sizes for both blank and AmB loaded chitosan-TPP and chitosan-dextran sulphate nanoparticles. In contrast, the use of sucrose as a cryoprotectant produced good quality nanoparticles and caused 39.5%, 17%, 21% and 6% increase in size for blank chitosan-TPP nanoparticles, blank chitosan-dextran sulphate nanoparticles, loaded AmB chitosan-TPP nanoparticles and loaded AmB chitosan-dextran sulphate nanoparticles respectively and did not lead to a significant difference in the zeta potential or PDI (p > 0.05 by t test).

2- D-mannitol as a cryoprotectant produced good quality nanoparticles, but caused 108%, 38%, 73% and 15.8% increase in size for blank chitosan-TPP nanoparticles, blank chitosan-dextran sulphate nanoparticles, loaded AmB chitosan-TPP nanoparticles and loaded AmB chitosan-dextran sulphate nanoparticles respectively and did not lead to a significant difference in the zeta potential (p > 0.05 by t test). When the two cryoprotectants are compared, sucrose produced significantly smaller nanoparticles with lower PDI for both types of nanoparticles, p < 0.05 t-test) (Table 3.5, Fig 3.8).

3- The morphological characteristics of blank chitosan-TPP or chitosan-dextran sulphate nanoparticles and AmB loaded nanoparticles were examined using TEM and SEM which showed a spherical structure for both chitosan-TPP or chitosan-dextran sulphate nanoparticles. The
TEM and SEM measured size of the four formulations, which was comparable to values measured by DLS. The incorporation of AmB into the nanoparticles did not change the shape of these nanoparticles, just increased the sizes (Fig 3.9 and Fig 3.10).

Table 3.5. Effect of cryoprotectants used during freeze drying on the physicochemical properties of unloaded and AmB loaded chitosan nanoparticles

<table>
<thead>
<tr>
<th></th>
<th>Nanoparticles</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Chitosan -TPP</td>
<td>AmB loaded chitosan – TPP</td>
<td>Chitosan – dextran sulphate</td>
<td>AmB loaded chitosan – dextran sulphate</td>
</tr>
<tr>
<td>Size (nm)</td>
<td>Before lyophilizing</td>
<td>48 ± 6</td>
<td>57 ± 7</td>
<td>145 ± 6</td>
</tr>
<tr>
<td></td>
<td>After lyophilization</td>
<td>+ sucrose 5%</td>
<td>67 ± 7</td>
<td>69 ± 8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>+ D- mannitol 5%</td>
<td>100 ± 9</td>
<td>99 ± 9</td>
</tr>
<tr>
<td></td>
<td>After lyophilization</td>
<td>+ sucrose 5%</td>
<td>0.25 ± 0.05</td>
<td>0.2 ± 0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>+ D- mannitol 5%</td>
<td>0.39 ± 0.07</td>
<td>0.4 ± 0.01</td>
</tr>
<tr>
<td>Zeta potential (mv)</td>
<td>Before lyophilizing</td>
<td>32.1 ± 1.2</td>
<td>29 ± 2</td>
<td>-15.5 ± 1</td>
</tr>
<tr>
<td></td>
<td>After lyophilization</td>
<td>+ sucrose 5%</td>
<td>28.5 ±1.9</td>
<td>25.5 ± 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>+ D- mannitol 5%</td>
<td>27 ± 2</td>
<td>24 ± 1</td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD (experiments were repeated three times with confirmed similar data). Sucrose more effectively protected the nanoparticles in comparison with D-mannitol (p < 0.05 by t-test).
**Figure 3.8.** Effectiveness of sucrose 5% and D-mannitol 5% as a cryoprotectant for freeze drying of blank and AmB loaded chitosan nanoparticles suspensions. 1: Blank chitosan-TPP nanoparticles before lyophilizing, 2: Blank chitosan-TPP nanoparticles after lyophilizing + sucrose 5%, 3: Blank chitosan-TPP nanoparticles after lyophilizing + D-mannitol 5%, 4: AmB loaded chitosan-TPP nanoparticles before lyophilizing, 5: AmB loaded chitosan-TPP nanoparticles after lyophilizing + sucrose 5%, 6: AmB loaded chitosan-TPP nanoparticles after lyophilizing + D-mannitol 5%, 7: Blank chitosan-dextran nanoparticles before lyophilizing, 8: Blank chitosan-dextran nanoparticles after lyophilizing + sucrose 5%, 9: Blank chitosan-dextran after lyophilizing + D-mannitol 5%, 10: AmB loaded chitosan-dextran nanoparticles before lyophilizing, 11: AmB loaded chitosan-dextran after lyophilizing + sucrose 5%, 12: AmB loaded chitosan-dextran nanoparticles after lyophilizing + D-mannitol 5%. Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data).
Figure 3.9. TEM micrographs of unloaded and amphotericin B loaded chitosan nanoparticles. A: Unloaded chitosan–TPP nanoparticles, B: AmB loaded chitosan–TPP nanoparticles, C: Unloaded chitosan – dextran sulphate nanoparticles, D: AmB loaded chitosan–dextran sulphate nanoparticles. TEM images indicate the nanoparticles to be spherical. Magnification: 40000x
3.3.4. Stability of physicochemical properties of AmB loaded chitosan nanoparticles

Both AmB loaded chitosan–TPP and chitosan–dextran sulphate nanoparticles did not show any significant change in their size or zeta potential at temperatures of 4, 34 and 37 °C when incubated in different media (water, PBS or RPMI at pH 7.5 or 6.5) or in mouse (BALB/c) plasma at 4 °C for a period of 30 days which indicated a high stability of these nanoparticles (Tables 3.6 and 3.7, Fig 3.11). Moreover, no significant difference in PDI was identified after 30 days in these different conditions.

Figure 3.10. SEM micrographs of unloaded and amphotericin B loaded chitosan nanoparticles. A: Unloaded chitosan–TPP nanoparticles, B: AmB loaded chitosan–TPP nanoparticles, C: Unloaded chitosan – dextran sulphate nanoparticles, D: AmB loaded chitosan–dextran sulphate nanoparticles. SEM images indicate the nanoparticles to be spherical and with similar sizes with the zetasizer. Magnification 50000x
Figure 3.11. Size of AmB loaded chitosan-TPP nanoparticle (A) and AmB loaded chitosan-dextran sulphate nanoparticle (B) in different media over time. The nanoparticles were stable in size after 30 days of storage in different media and temperatures. Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data).
Table 3.6. Variations of physicochemical properties of AmB loaded chitosan-TPP nanoparticles in different media upon storage at different temperatures

<table>
<thead>
<tr>
<th></th>
<th>Day 0</th>
<th>Day 1</th>
<th>Days 7</th>
<th>Days 30</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Size nm</td>
<td>PDI</td>
<td>Zeta potential mv</td>
<td>Size nm</td>
</tr>
<tr>
<td>Water at 4, 34 or 37 °C</td>
<td>70 ± 6</td>
<td>0.1 ± 0.02</td>
<td>25.5 ± 1</td>
<td>74 ± 5</td>
</tr>
<tr>
<td>PBS at 4, 34 or 37 °C</td>
<td>73 ± 5</td>
<td>0. ± 0.01</td>
<td>23.3 ± 1</td>
<td>75 ± 4</td>
</tr>
<tr>
<td>RPMI (pH=7.5) at 4, 34 or 37 °C</td>
<td>75 ± 6</td>
<td>0.2 ± 0.1</td>
<td>24.1± 1</td>
<td>79 ± 7</td>
</tr>
<tr>
<td>RPMI (pH=6.5) at 4, 34 or 37 °C</td>
<td>68 ± 7</td>
<td>0.1 ± 0.01</td>
<td>32 ± 6</td>
<td>74 ± 5</td>
</tr>
<tr>
<td>Plasma at 4 °C</td>
<td>75 ± 7</td>
<td>0.1 ± 0.01</td>
<td>29 ± 6</td>
<td>77 ± 6</td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). No significant difference was shown in the size, PDI or zeta potential between two types of the nanoparticles after 30 days storage (p >0.05 by t – test).
Table 3.7. Variations of physicochemical properties of AmB loaded-chitosan dextran sulphate nanoparticles in different media upon storage at different temperatures

<table>
<thead>
<tr>
<th></th>
<th>Day 0</th>
<th>Day 1</th>
<th>Days 7</th>
<th>Days 30</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Size (nm)</td>
<td>PDI</td>
<td>Zeta potential (mv)</td>
<td>Size (nm)</td>
</tr>
<tr>
<td>Water at 4, 34 or 37 °C</td>
<td>180 ± 6</td>
<td>0.2± 0.1</td>
<td>-14 ± 5</td>
<td>187 ± 5</td>
</tr>
<tr>
<td>PBS at 4, 34 or 37 °C</td>
<td>177 ± 5</td>
<td>0.2 ± 0.1</td>
<td>-15 ± 5</td>
<td>178 ± 4</td>
</tr>
<tr>
<td>RPMI (pH=7.5) at 4, 34 or 37 °C</td>
<td>180 ± 6</td>
<td>0.2± 0.1</td>
<td>-20 ± 5</td>
<td>183 ± 7</td>
</tr>
<tr>
<td>RPMI (pH=6.5) at 4, 34 or 37 °C</td>
<td>175 ± 7</td>
<td>0.2± 0.1</td>
<td>-11 ± 5</td>
<td>178 ±5</td>
</tr>
<tr>
<td>Plasma at 4 °C</td>
<td>177 ± 7</td>
<td>0.2± 0.1</td>
<td>-15 ± 5</td>
<td>179 ±5</td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). No significant difference was shown in the size, PDI or zeta potential between two types of the nanoparticles after 30 days storage (p >0.05 by t – test).
3.3.5. Nanoparticles loading and encapsulation properties

Both types of nanoparticles had a high encapsulation efficiency of more than 90%. The fluffy yellow yield was more than 90% for both types (Table 3.8). There was no significant difference in AmB loading between chitosan TPP and dextran sulphate nanoparticles ($p>0.05$ by t-test) (Fig 3.12).

Table 3.8. Percentage of AmB loading, encapsulation and yield

<table>
<thead>
<tr>
<th>Type of nanoparticles</th>
<th>EE %</th>
<th>AmB loading %</th>
<th>Yield %</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td>94 ± 5</td>
<td>26 ± 1</td>
<td>93 ± 6</td>
</tr>
<tr>
<td>AmB loaded chitosan-dextran sulphate nanoparticles</td>
<td>92 ± 8</td>
<td>23 ± 2</td>
<td>92 ± 6</td>
</tr>
</tbody>
</table>

data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). No significant difference was shown between AmB loaded chitosan-TPP and dextran sulphate nanoparticles ($p>0.05$ by t-test). AmB loaded chitosan-TPP nanoparticles size= 69 ± 8 nm and AmB loaded chitosan-dextran sulphate nanoparticles size= 174 ± 8 nm

Figure 3.12. Comparison of AmB encapsulation, loading and yield of the two types of nanoparticles. No significant difference was shown between AmB loaded chitosan-TPP and dextran sulphate nanoparticles regarding the encapsulation, loading and yield ($p>0.05$ by t-test). Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). AmB loaded chitosan TPP nanoparticles size= 69 ± 8 nm and AmB loaded chitosan dextran sulphate nanoparticles size= 174 ± 8 nm
3.3.6. *In vitro* release of AmB from the nanoparticles

AmB release from the two types of nanoparticles is shown in Fig 3.13 and Table 3.9. The chitosan-TPP and chitosan-dextran sulphate nanoparticles showed a slow release within 7 days in PBS (at two pH values of 7.5 and 6.5) at three temperatures 4, 34 and 37 °C and in mouse (BALB/c) plasma at 37 °C. Chitosan-TPP nanoparticles released AmB significantly quicker than chitosan-dextran sulphate nanoparticles at the different conditions (nanoparticle suspended in plasma or PBS and at different pHs and temperatures) (*p*<0.05, one-way-ANOVA). Neither AmB loaded chitosan-TPP nanoparticles nor AmB loaded chitosan-dextran sulphate nanoparticles showed any significant difference in the drug release after storing at 34 °C or 37 °C (*p*>0.05 by t-test) (Fig 3.13, Table 3.9). However, the pH influenced the drug release significantly with both types of nanoparticles, showing higher cumulative releases at the lower pH of 5 than at higher pH of 6.5 or 7.5 (*p*<0.05 by t-test) (Table 3.9 and Fig 3.13).
Table 3.9. *In vitro* cumulative release of AmB from the two formulations at different conditions

<table>
<thead>
<tr>
<th>Type</th>
<th>6 h %</th>
<th>24 h %</th>
<th>48 h %</th>
<th>72 h %</th>
<th>96 h %</th>
<th>120 h %</th>
<th>144 h %</th>
<th>168 h %</th>
</tr>
</thead>
<tbody>
<tr>
<td>AmB loaded chitosan–dextran sulphate nanoparticles</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBS, pH 7.4</td>
<td>4 °C</td>
<td>0.1±0.05</td>
<td>1±0.05</td>
<td>2.2±0.4</td>
<td>5.2±1</td>
<td>7.5±2</td>
<td>9.5±2</td>
<td>11±2</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>0.3±0.1</td>
<td>2.5±0.2</td>
<td>5.2±1</td>
<td>8.5±2</td>
<td>10±3</td>
<td>13.5±2</td>
<td>16.4±3</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>0.1±0.02</td>
<td>2±0.1</td>
<td>4.4±1</td>
<td>6.9±1</td>
<td>9.1±2</td>
<td>12.5±3</td>
<td>15.5±3</td>
</tr>
<tr>
<td>PBS, pH 6.5</td>
<td>4 °C</td>
<td>0.2±0.02</td>
<td>2±0.2</td>
<td>3.1±1</td>
<td>4.9±1</td>
<td>6.9±1</td>
<td>8.9±1</td>
<td>11.5±2</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>0.4±0.1</td>
<td>4±0.5</td>
<td>7.3±2</td>
<td>9.2±3</td>
<td>13.1±3</td>
<td>15±2</td>
<td>17.2±4</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>0.1±0.05</td>
<td>2.9±0.4</td>
<td>5.4±1</td>
<td>7.9±2</td>
<td>10.1±2</td>
<td>12.2±2</td>
<td>16.5±3</td>
</tr>
<tr>
<td>PBS, pH 5</td>
<td>4 °C</td>
<td>0.2±0.05</td>
<td>3.5±1</td>
<td>9.5±2</td>
<td>16.1±4</td>
<td>17.2±3</td>
<td>20.2±3</td>
<td>21.1±4</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>0.5±0.1</td>
<td>7.5±2</td>
<td>14.5±3</td>
<td>20.9±5</td>
<td>23±4</td>
<td>24.9±3</td>
<td>27.5±4</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>0.3±0.1</td>
<td>6.5±1</td>
<td>13.5±3</td>
<td>20.1±4</td>
<td>21.2±5</td>
<td>24.2±3</td>
<td>26.1±3</td>
</tr>
<tr>
<td>Plasma</td>
<td>37 °C</td>
<td>0.2±0.05</td>
<td>4.1±1</td>
<td>8.1±1</td>
<td>9.2±2</td>
<td>10.1±2</td>
<td>12±2</td>
<td>14.9±2</td>
</tr>
<tr>
<td>AmB loaded chitosan – TPP nanoparticles</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBS, pH 7.4</td>
<td>4 °C</td>
<td>0.5±0.1</td>
<td>5.1±1</td>
<td>9.2±1</td>
<td>11.5±2</td>
<td>13.8±2</td>
<td>15.9±1</td>
<td>18.9±2</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>1.2±0.3</td>
<td>9.9±2</td>
<td>15.6±2</td>
<td>20.6±3</td>
<td>24.5±5</td>
<td>26±4</td>
<td>28.9±5</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>1±0.2</td>
<td>10±2</td>
<td>14.9±3</td>
<td>19.5±2</td>
<td>23.5±5</td>
<td>24.5±3</td>
<td>27.5±4</td>
</tr>
<tr>
<td>PBS, pH 6.5</td>
<td>4 °C</td>
<td>0.3±0.1</td>
<td>4.1±1</td>
<td>10.2±2</td>
<td>12.5±2</td>
<td>15.8±5</td>
<td>17.9±2</td>
<td>19.9±3</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>1.5±0.3</td>
<td>10.5±2</td>
<td>16.4±4</td>
<td>21.9±4</td>
<td>26.3±5</td>
<td>27.8±3</td>
<td>29.8±5</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>1.2±0.4</td>
<td>9.8±1</td>
<td>15.2±3</td>
<td>20.2±3</td>
<td>24.1±5</td>
<td>25.6±4</td>
<td>28±4</td>
</tr>
<tr>
<td>PBS, pH 5</td>
<td>4 °C</td>
<td>0.9±0.2</td>
<td>16.5±3</td>
<td>19.8±3</td>
<td>25.5±4</td>
<td>26.2±4</td>
<td>34.5±4</td>
<td>40.2±6</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>1.5±0.4</td>
<td>21.2±4</td>
<td>27.2±5</td>
<td>31.2±3</td>
<td>34.6±6</td>
<td>39.8±5</td>
<td>41.9±5</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>1.7±0.4</td>
<td>20.2±3</td>
<td>26.5±6</td>
<td>30.2±4</td>
<td>33.1±4</td>
<td>40.2±5</td>
<td>45.2±4</td>
</tr>
<tr>
<td>Plasma</td>
<td>37 °C</td>
<td>1.7±0.3</td>
<td>11.2±2</td>
<td>14.5±4</td>
<td>20.9±2</td>
<td>25.3±3</td>
<td>27.3±4</td>
<td>29.9±4</td>
</tr>
<tr>
<td>AmB solution</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBS, pH 7.4</td>
<td>4 °C</td>
<td>84±2</td>
<td>100±1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>85±2</td>
<td>100±2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>86±3</td>
<td>100±2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>PBS, pH 6.5</td>
<td>4 °C</td>
<td>83±1</td>
<td>100±1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>86±2</td>
<td>100±3</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>88±4</td>
<td>100±2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>4 °C</td>
<td>84±1</td>
<td>100±2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>PBS, pH 5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------</td>
<td>-----------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
</tr>
<tr>
<td></td>
<td>34 °C</td>
<td>85±1</td>
<td>100±2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>37 °C</td>
<td>87±2</td>
<td>100±2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Plasma</td>
<td>37 °C</td>
<td>85±2</td>
<td>100±2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). Both types of nanoparticles showed significantly more cumulative release in the low pH of 5 than in higher pH of 6.5 or 7.5 (p<0.05 by t-test). The AmB release from chitosan-TPP nanoparticles was faster than chitosan dextran sulphate nanoparticles (p < 0.05 by t - test). AmB loaded chitosan-TPP nanoparticles size= 69 ± 8 nm and AmB loaded chitosan-dextran sulphate nanoparticles size= 174 ± 8 nm.
Figure 3.13. *In vitro* release profile of AmB loaded chitosan nanoparticles at 37 °C. **A**: AmB loaded chitosan-dextran sulphate nanoparticles in PBS (pH of 5, 6.5 or 7.5) and mouse (BALB/c) plasma, **B**: AmB loaded chitosan-TPP nanoparticles in PBS (pH of 5, 6.5 or 7.5) and mouse plasma, **C**: Comparison.
(BALB/c) plasma and C: comparison of AmB release from AmB solution, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan dextran sulphate nanoparticles in PBS at pH 5 and 7.5. Data expressed as mean +/- SD (experiment was reproduced three times with confirmed similar data). AmB loaded chitosan-TPP nanoparticles size= 69 ± 8 nm and AmB loaded chitosan-dextran sulphate nanoparticles size= 174 ± 8 nm
3.4. Discussion

Blank and AmB loaded chitosan-TPP and chitosan-dextran sulphate nanoparticles were successfully produced via the inotropic gelation method. Homogeneously dispersed nanoparticles with different sizes from 50 nm to around 400 nm, with low PDI, and with positive or negative surface charge were synthesised. The effects of experimental conditions and parameters (initial concentration of chitosan, chitosan: TPP or chitosan: dextran sulphate mass ratios, pH of chitosan solution and sonication time) on the physicochemical properties of the nanoparticles (size, PDI and charge) were determined. The aim was to create positively and negatively charged nanoparticles with the smallest size and lowest PDI. The PDI value indicates dispersion homogeneity and the distribution of the nanoparticles sizes in the sample and high PDI means variable ranges of sizes in the sample while lower PDI reflects constantly sized nanoparticles (253). We aimed and succeeded in synthesising the smallest sizes for both types of nanoparticles, as smaller nanoparticles with size 100 nm showed a 2.5-fold higher uptake in Caco-2 cells than larger particles with size 1 µm and a 6-fold higher uptake than particles sized 10 µm (254). Additionally, nanoparticles with small size have exhibited maximum deposition of their content in the skin dermis (after topical application) and small nanoparticles can facilitate macrophage targeting residence in the skin (after intravenous injection) (32, 246, 255, 256). Subsequently, smaller nanoparticles in literatures offered higher uptake rates, more permeability through skin and higher targeting to skin and these properties are substantial in CL treatment.

A paper reported that negatively charged nanoparticles are taken up significantly more than positively charged nanoparticles by Caco-2 epithelial cells (257).

This encouraged us to prepare two types of chitosan nanoparticles (positive and negative charged nanoparticles with smallest possible size).

**Influence of reactant mass ratio on the nanoparticles**
We indicated that the optimal mass ratio to obtain good quality nanoparticles with smallest size and lowest PDI was 5:1 for chitosan: TPP (at chitosan 3 mg/ml) and this was consistent with literatures (258, 259). It was shown that at this ratio, TPP anions are adequately incorporated into chitosan and as a result, a further boost in the cross-linking and tightening of the chitosan chains within the particle result, which explains the decrease in the nanoparticles sizes, as previously speculated by Masarudin et al (2015) (253). Regarding dextran sulphate, a mass ratio 1:3 of chitosan: dextran sulphate (at chitosan 1 mg/ml) gave the smallest nanoparticles size and the lowest PDI and this is similar with another published report published by Tiyaboonchai et al (238). As a more concentration of dextran sulphate in comparison with chitosan might increase the level of complexation of the nanoparticles and the chitosan chains can become entangled to a great extent (238, 260).

**Influence of initial concentration of chitosan on the nanoparticles**

We showed that using a high initial concentration of chitosan (9 mg/ml) led to poor quality and aggregation of nanoparticles and this was in agreement with a previously published report (261). At this high concentration of chitosan, more molecules of chitosan tend to entangle with each other and crosslink with counter ion (TPP) or sulphate groups of dextran sulphate to form larger particles (261) and moreover, this aggregation could be attributed to the higher number of positive groups as these positive groups can make the chitosan chain to stretch because of the intra chitosan chain repulsion (133, 262, 263).

**Influence of pH of chitosan solution on the nanoparticles**

Additionally, the pH of chitosan solution played an important role in the quality of the nanoparticles. The synthesized nanoparticles were aggregated and of poor quality at pH 3 and pH 7. At pH 3, chitosan is highly protonated with high charge density. While at pH 7, chitosan has a low charge and is partially solubilized. The greater positive charge at pH 3 can make the chitosan chain stretch because of the intra molecular repulsion while at higher pH of 7, there is a large reduction in the protonation degree of the nanoparticles and that large leads to reduce the inter particles electrostatic repulsion among these
nanoparticles. Therefore, there is a higher possibility of the aggregation (133, 262).

**Zeta potential of the nanoparticles and the Influence of sonication duration on the nanoparticles**

Zeta potential of the nanoparticles increased with increasing concentration and with decreasing pH of the initial chitosan solution used. This increase in zeta potential values could be explained as; the higher concentration of chitosan leads to more total amino groups and consequently more protonated positive \(-\text{NH}_3^+\) on the surface of the nanoparticles and lower pH of chitosan solution results in more positive amino groups (133, 262). To assess the influence of sonication time on the physicochemical characteristics of nanoparticles, the prepared nanoparticle dispersions were subjected to sonication for 1, 5, 15 and 20 mins. The sonication duration had a critical role in the quality of nanoparticles as sonicating for 15 mins resulted in favoured nanoparticles and this was in accordance with other reports. Too little sonication duration is insufficient to break the aggregation of the nanoparticles and after 20 mins the aggregation cannot be further broken so size and PDI remain constant (264, 265, 266).

**The freeze drying process of the nanoparticles**

The freeze drying process causes many stresses related to freezing and dehydration and these stresses can destabilize the nanoparticles suspensions and lead to poor quality and aggregation of the nanoparticles (242). This is what we found in our study. Therefore, it is recommended to use protectants for the nanoparticles to protect them from the freeze and dry stress. The most common cryoprotectants used in literatures are sugars as they can form a glassy matrix that can protect the nanoparticles from the mechanical stress and avoid aggregation, so we used D-mannitol and sucrose in our study. We determined that sucrose had a greater protective (2-3 x) effect on both types of nanoparticles than D-mannitol. A similar finding has previously been reported that sucrose is more successful than D-mannitol in protecting the nanoparticles from the lyophilisation (267) possibly due to the fact that sucrose
does not crystallize during lyophilisation process, unlike D-mannitol, as previously reported (242).

**Stability of the nanoparticles and their AmB encapsulation and loading properties**

We found that blank and AmB loaded chitosan-TPP or chitosan-dextran sulphate nanoparticles were stable in terms of size and zeta potential for 30 days at different temperatures (4, 34 and 37 °C) and in different media (water, PBS, RPMI and mouse (BALB/c) plasma). Another published report showed that sizes of AmB loaded chitosan-TPP nanoparticles were stable for 6 months in water at 4° C and at room temperature and chitosan-dextran nanoparticles were stable in terms of sizes and zeta potential for 4 weeks (240, 268). The encapsulation efficacy of AmB in both types of nanoparticles was around 90% and similar data was reported for AmB encapsulation in chitosan nanoparticles with TPP (80%) or with chondroitin sulphate (90%) (122, 240). While the loading of AmB was 23% and 26% w/w for chitosan-TPP and chitosan-dextran sulphate nanoparticles respectively with two times more loading in comparison with chitosan chondroitin sulphate nanoparticles (122).

**AmB release from the nanoparticles**

The release profiles of AmB from AmB solution (as a control) through synthetic membrane was significantly higher than from AmB loaded chitosan-TPP or chitosan-dextran sulphate nanoparticles (p<0.05 by t test). The nature of the complexation agent (TPP or dextran sulphate) did not influence the slow AmB release from both types of nanoparticles in PBS and mouse plasma. The nanoparticles stability in plasma (size and charge) and slow release of AmB in plasma would ensure that AmB does not bind to low density plasma lipoproteins thereby avoiding any potential AmB toxicity. This is consistent with another study of stability of a noncovalent complex of amphotericin B (AmB) with poly(α-glutamic acid) (PGA) in mouse CD/1 serum (118). Both types of chitosan nanoparticles exhibited a pH-dependent AmB release, with a greater release at a low pH of 5 than at higher pH of 7.5. This is likely to be due to the higher solubility of chitosan in acidic media (269).
Drug release from chitosan-TPP nanoparticles was faster than from chitosan-dextran sulphate nanoparticles and this could be explained as chitosan-TPP nanoparticles are significantly smaller than chitosan-dextran sulphate nanoparticle. The resulting larger surface area to volume ratio of chitosan-TPP nanoparticles would allow greater AmB release from the surface of the nanoparticles as more of the drug is closer to the surface (254). The negatively charge of chitosan-dextran sulphate nanoparticles would also play a role in slower release which reported previously of insulin release (270).

3.4.1. Conclusion

In summary, we successfully prepared two different types of AmB loaded chitosan nanoparticles, one smaller size nanoparticle with positive surface charge and the other with larger size and negative charge. The synthesized nanoparticles were able to efficiently encapsulate AmB. Different parameters such as chitosan concentration, chitosan: TPP or chitosan: dextran sulphate mass ratio and chitosan solution pH significantly affected the physicochemical characterization of the nanoparticles. Both positive and negative nanoparticles showed a high stability in terms of size and at different temperatures. As expected, these nanoparticles exhibited a prolonged AmB release. Therefore, they appear to be good candidates for further investigation into their anti-leishmanial activity by different routes of administration.
4. *In vitro* and *in vivo* activity of chitosan formulations in experimental cutaneous leishmaniasis

4.1. Introduction

The intravenous AmB (as mentioned in the introduction) is one of the available second-line drugs for leishmaniasis which acts by forming pores in the cell membrane of *Leishmania* via complexation with ergosterol. However, the use of the conventional deoxycholate amphotericin B (Fungizone) is clinically limited because of the infusion-related side effects such as, fever, nausea, vomiting, rigours and two more serious effects: anaemia and nephrotoxicity (271, 272). A great interest of research to develop the drug delivery system of AmB in leishmaniasis treatment arises. Accordingly, liposomal formulation (AmBisome®, size= 70-80 nm (272, 273)) with a better tolerated profile and low toxicity issues was developed and approved by FDA for the treatment of VL. It has showed clinical effectivity in CL patients, in multiple doses (3 mg/kg daily for a total of 21 mg/kg) (272, 274). Although AmBisome® is on the WHO Essential Medicines List, this formulation has some limitations in terms of the high price (at least 200 USD$ per vial of 50 mg, is donated by Gilead via WHO for free for VL, not for CL ), the need for cold chain, shelf-life related issues, slow infusion and the difficult to access the drug in many countries (170, 275, 276). Moreover, AmBisome® has a complex production process and an increase in particles size and a change in the drug content upon storage of AmBisome® have been reported (during 72 h of storage) (276, 277).

Other drug delivery systems (DDs) used in the *Leishmania* field encounter some disadvantages summarised in Table 4.1, in addition to accumulation of lipid in liver and spleen caused by the lipidic formulations that may cause pathological conditions (278).
Table 4.1. Disadvantages of different DDs (67, 112, 279, 280, 281, 282, 283, 284)

<table>
<thead>
<tr>
<th>DDs</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liposomes</td>
<td>High cost, low stability and using an organic solvent in the preparation</td>
</tr>
<tr>
<td>Niosomes</td>
<td>Instability, leaking of entrapped drug and Hydrolysis of encapsulated drugs which limiting the shelf life of the dispersion</td>
</tr>
<tr>
<td>Nanodiscs</td>
<td>Lack of size control, using an organic solvent and other drawback is the precipitation under low pH&lt;6</td>
</tr>
<tr>
<td>Emulsions</td>
<td>The need to use a high concentration of surfactants and cosurfactant, stability highly influenced by pH and temperatures and desorption of surfactants</td>
</tr>
<tr>
<td>Solid lipid nanoparticle</td>
<td>Organic solvent, low drug loading efficiency, fast drug burst release and the possibility of drug expulsion during storage because of the crystalline structure</td>
</tr>
</tbody>
</table>

Polymeric nanoparticles, prepared by inotropic gelation method (formed by interactions between two oppositely charged molecules), have gained a great interest in the DDs, with advantages over other DDs as their preparation is usually at lower costs, simple, quick, does not require the use of organic solvents (generally) and the long shelf life of these nanoparticles at room temperature (114).

In Chapter 3, we successfully synthesised two types of AmB loaded nanoparticles (one was positively charged by using TPP and the other was negatively charged by using dextran sulphate) by using ionotropic gelation methods without using any organic solvents. This process was fast, simple and with low cost of 55 USD$ approx. for 1 g of AmB nanoparticles. These nanoparticles, in contrast to liposomal formulations, showed a high stability in different media (water, PBS, RPMI and mouse (BALB/c) plasma) at different temperatures for a period of 30 days, and they showed a slow drug release in these media. All these characteristics of AmB loaded chitosan nanoparticles (the high stability for a long time in different conditions in terms of size and charge, slow drug release, easy preparation method and low cost etc), made them a suitable candidate for further investigations for CL treatment.

In the literature, chitosan nanoparticles have shown a good activity against a wide range of microbes and are sometimes more active than chitosan solution (MMW and HMW) (Table 4.1). Some studies showed that both chitosan
solution and chitosan nanoparticles have the same antimicrobial mechanism i.e. by interacting with microbial cell membrane or binding with microbial DNA (285). AmB encapsulated in different types of chitosan nanoparticles has been evaluated against leishmaniasis with promising results in vitro and in vivo and the studies are summarised in Table 4.2. Most of these studies used positively charged nanoparticles with a size greater than 100 nm. In contrast, we decided to investigate smaller nanoparticles (for possible skin permeation, and as smaller size of nanoparticles facilitates a passive transport from blood vessels to tissues when administrated intravenously and can enhance the extravasation in the inflamed lesions on the skin and can facilitates macrophage targeting residence in the skin (255, 256) ), with positive charge (when prepared with TPP) or negative charge (when prepared with dextran sulphate) to identify any influence of nanoparticle charge.

Therefore, this chapter aimed to evaluate:

(i) the in vitro activity of blank and amphotericin B loaded chitosan TPP or dextran sulphate nanoparticles against L. major and L. mexicana promastigotes and amastigotes
(ii) the in vivo anti-leishmanial activity of blank and amphotericin B loaded chitosan TPP or dextran sulphate nanoparticles (through intravenous route) in murine models of CL caused by L. major
(iii) the permeation of these nanoparticles and chitosan solution through BALB/c skin by a Franz diffusion study.

We did not include HMW chitosan solution in the intravenous route of the in vivo study as its diluted acid solutions were too viscous and this makes it very difficult and not suitable for mice intravenous route (HMW chitosan viscosity is 800-2000 cP, 1 wt. % in 1% acetic acid) (286, 287).
Table 4.2. Antimicrobial activity of chitosan nanoparticles (285)

<table>
<thead>
<tr>
<th>Nanoparticles</th>
<th>Microbes</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chitosan nanoparticles (Chitosan, MW 220 KDa)</td>
<td><em>Staphylococcus aureus</em></td>
<td>Chitosan nanoparticles were more effective than chitosan solution and doxycycline(288)</td>
</tr>
<tr>
<td></td>
<td><em>Escherichia coli</em></td>
<td></td>
</tr>
<tr>
<td>Chitosan nanoparticles (LMW)</td>
<td><em>Streptococcus mutans</em></td>
<td>Inhibited biofilm formation (289)</td>
</tr>
<tr>
<td>Chitosan-silver nanocomposites (HMW)</td>
<td><em>Staphylococcus aureus</em></td>
<td>A synergistic antimicrobial activity between chitosan and silver nanomaterials (290)</td>
</tr>
<tr>
<td>Chitosan nanoparticles (Chitosan, MW 310 KDa)</td>
<td><em>Candida albicans</em></td>
<td>Chitosan nanoparticles were more active than chitosan with lower MIC(50) (291)</td>
</tr>
<tr>
<td>AmB loaded chitosan nanoparticles (LMW)</td>
<td><em>Candida albicans</em></td>
<td>Chitosan nanoparticles showed similar activity to AmB with higher corneal penetration(292)</td>
</tr>
<tr>
<td>DDS</td>
<td>Preparation method</td>
<td>Nanoparticles properties</td>
</tr>
<tr>
<td>-----</td>
<td>---------------------</td>
<td>--------------------------</td>
</tr>
</tbody>
</table>
| Nanoemulsion based chitosan nanocapsule entrapping AmB | First, an oil-in-water (o/w) nanoemulsion was formulated by modified spontaneous emulsification solvent evaporation. Secondly nanocarrier was generated by coating with chitosan deposition on the water-oil surface | Size= 146 ± 9 nm Zeta potential= +29±0.8 mV | AmB DDS was significantly less toxic against the J774A cell line | In vitro: EC\textsubscript{50} for AmB DDs, AmBisome\textsuperscript{®} and Fungizone was 0.19±0.04, 0.29 ± 0.03 and 0.48 ± 0.05 μg/ml respectively against L. donovani promastigotes  
In vivo: L. donovani infected hamster model received (i.p.) AmB-loaded formulations at 1 mg/kg on 5 consecutive days. AmB DDS, AmBisome\textsuperscript{®} and Fungizone caused 86 ±2%, 70 ±3 % and 56 ±4% inhibition of amastigotes in spleen. (219) |
| Chitosan-coated solid lipid nanoparticles were developed and loaded with amphotericin B | Solvent emulsification-evaporation | Size= 159 ±25 nm | In mice model, AmB DDS was significantly ten-fold less toxic than pure AmB solution and was safe up to AmB concentration equivalent to 5 mg/kg body weight. | In vitro: EC\textsubscript{50} of AmB DDs, AmBisome\textsuperscript{®} and Fungizone was 0.046±0.02, 0.157±0.03 and 0.320±0.08 μg/ml respectively against L. donovani amastigotes infecting adherent mouse macrophage cell line J774A.1 (259) |
| AmB loaded pluronic F127 (PF 127) micelles coated with chitosan | Thin film hydration | Size= 139 ± 3 to 170 ± 53 nm Zeta potential= +11.0 ± 2 to +53 ± 5 mV | AmB DDS was ten-fold less toxic than pure AmB solution against J774A.1 cell | In vitro: EC\textsubscript{50} of AmB DDs, and AmB solution 0.05 and 0.09 μg/ml respectively against L. donovani amastigotes infecting macrophage cell line J774A.1(293) |
| AmB loaded chitosan nanoparticles | Inotropic gelation method (between chitosan positively charged and chondroitin sulphate negatively charged) | Size= 136±11 nm Zeta potential= +8.4 to +30.2 mV | Cytotoxicity against murine macrophages of AmB DDS was nearly ten-fold less compared to pure AmB solution | In vitro: EC$_{50}$ of AmB DDS and AmB was 1±0 and 0.1±0 respectively (AmB DDS was less active) against *L. amazonensis* and 0.1±0 and 0.1±0 µg/ml respectively against *L. chagasi* (AmB DDS had similar activity to AmB). AmB DDS and AmB caused 90% and 89% reduction of *L. amazonensis* internalized macrophages (%) (122) |
| AmB loaded chitosan nanoparticles | Inotropic gelation method (between chitosan positively charged and chondroitin sulphate negatively charged) | Size= 136±11 nm Zeta potential= +8.4 to +30.2 mV | Cytotoxicity against murine macrophages of AmB DDS was nearly ten-fold less compared to pure AmB | In vivo: *L. amazonensis* infected BALB/c mice received (i.v.) 1 mg of drug/kg daily for 10 days. AmB DDS treated mice showed a smaller lesion size which was sustained up to 30 days after the treatment compared with AmB treated group (83) |
| AmB loaded chitosan nanoparticles | Phase separation method by mixing chitosan - TPP nanoparticles with AmB solution under stirring for 7 days | Size= 112 nm Zeta potential= +8mV | Mortality in mice received (i.p.) AmB solution 10 mg/kg was 10% while 0% in mice received AmB nanoparticles (10 mg/kg AmB equivalent) every other day for 3 weeks | In vitro: *L. major* promastigote killing (%): 82% at 20 µg/mL. *L. major* amastigote killing (%): 78% at 20 µg/mL. In vivo: *L. major* infected BALB/c mice received (i.p.) AmB nanoparticles of 10 mg/kg while the positive control mice received AmBisome® of 50mg/kg. There was no significant difference in the efficacy of the two formulations and caused 100% reduction of lesion size. (294) and https://www.dovepress.com/comparative-analysis-between-four-model-nanoformulations-of-amphoteric-peer-reviewed-article-IJN |
4.2. Material and methods

4.2.1. Preparation of blank and AmB loaded chitosan nanoparticles

All nanoparticles in this study were prepared and characterised as described in Chapter 3 in sections 3-2-1- and 3-2-2-, within the parameters (10 ml of HMW chitosan solution (30 mg in 10 ml AC 1%), 10 mg of AmB (Purity ≥ 95%, Cambridge Bioscience, UK) dissolved in 0.5 ml of DMSO (pure AmB), 10 ml of TPP solution (6 mg in 10 ml DS water) or 10 ml of dextran solution (30 mg in 10 ml double distilled water). After freeze drying the nanoparticle suspension, the white (blank nanoparticles) or yellow (AmB loaded nanoparticles) product was reconstituted in double distilled water. After this, these nanoparticles were characterised by size, charge and AmB loading as described in Chapter 3. Additionally, the AmB loading was evaluated again after freeze drying by dissolving the yielded yellow powder in DMSO, in acidic pH 3 (by using 1% (v/v) acetic acid), and then measuring the quantity of AmB by HPLC as described previously in Chapter 3, section 3.3.6.1. There was no significant difference in the loading value between this method and the previously used one in Chapter 3, section 3-2-5- (p<0.05 by t-test).

AmB (Purity ≥ 95%, Cambridge Bioscience, UK) dissolved in DMSO at a 10 mM stock and diluter for proper concentrations in RPMI-1640 with 10% HiFCS (pure AmB).

AmBisome® (a liposomal formulation of AmB, Gilead Sciences international Ltd, UK) was prepared according to the manufacturer’s instructions. Briefly, a suspension of AmB liposome was prepared in cold sterile MilliQ water to obtain an initial concentration of 4 mg/ml. The suspension was shaken and incubated at 65°C for 10 mins and then cooled to room temperature. Further dilution to the required concentration of AmBisome® was done with 5% dextrose (w/v) (71).

4.2.2. Red blood cells haemolysis

Blood samples were obtained from two human donors (O+) (volunteers, Queen Mary, University of London) drawn directly into EDTA tubes to prevent coagulation. Blood samples were centrifuged at 500 x g for 5 min and the
plasma aspirated and discarded. The remaining red blood cells (RBCs) were then washed three times in buffered saline (10 mM Tris, 150 mM NaCl, pH 7.4) prior to the assay. The RBCs were diluted to a density of 5x10^8 cells/ml and exposed to 1000, 500, 250, 125, 65.5, 31.25, 15.62 and 7.81 μg/ml of chitosan solution (HMW chitosan), blank chitosan-TPP nanoparticles, AmB loaded chitosan-TPP nanoparticles (AmB equivalent), blank chitosan-dextran sulphate nanoparticles, AmB loaded chitosan-dextran sulphate nanoparticles (AmB equivalent) and pure AmB in 96 well plates (200 μl in each well) for 1 h at 37°C. The plate was centrifuged for 5 mins at 500 x g to pellet intact RBCs. 100 μl of supernatant from each well was transferred into a clear, flat-bottomed 96-well plate and cell lysis was determined spectrophotometrically (540 nm). Phosphate buffer was used as a negative control and 20% Triton X-100 was used as a positive control representing 100% haemolysis. The results were expressed as the mean percentage reduction in human red blood cells compared with non-treated control wells, and represented by the 50% haemolytic concentration (RBC_{50}) (295).

4.2.3. In vitro cytotoxicity assays

Cytotoxicity of chitosan formulations against KB cells was evaluated at concentrations of 2000, 1000, 500, 250, 125, 65.5, 31.25 and 15.62 μg/ml of blank chitosan-TPP nanoparticles, blank chitosan-dextran sulphate nanoparticles, AmB loaded chitosan-TPP nanoparticles (AmB equivalent) and AmB loaded chitosan-dextran sulphate nanoparticles (AmB equivalent). Cytotoxicity was evaluated in RPMI 1640 at two pH values (at normal pH of RPMI 7.5 and at a lower pH 6.5). Pure AmB, AmBisome® and chitosan solution (HMW chitosan) were included in this experiment for comparison. Podophyllotoxin (Sigma, UK) was included as a positive control at a starting concentration of 0.05 μM. Cytotoxicity was evaluated by a cell viability assay using the resazurin sodium salt solution (AlamarBlue, Sigma, UK) as described in Chapter 2, section 2.8.5.
4.2.4. *In vitro* 72-hour activity of chitosan and its derivatives against extracellular *L. major* and *L. mexicana* promastigotes

The activity of chitosan formulations against *L. major* and *L. mexicana* promastigotes was evaluated at concentrations 486, 162, 54, 18, 6, 2, 0.66, 0.22, 0.072, 0.024 and 0.008 μg/ml of blank chitosan-TPP nanoparticles, blank chitosan-dextran sulphate nanoparticles, AmB loaded chitosan-TPP nanoparticles (AmB equivalent) and AmB loaded chitosan-dextran sulphate nanoparticles (AmB equivalent). The anti-leishmanial activity was evaluated in RPMI 1640 at two pH values (7.5 and 6.5). Pure AmB, AmBisome® and chitosan solution (HMW chitosan) were included in this experiment for comparison. See Chapter 2, section 2.8.6 for full details.

4.2.5. *In vitro* 72-hour activity of chitosan and its derivatives against intracellular amastigotes of *L. major* and *L. mexicana*

The activity of chitosan formulations against *L. major* and *L. mexicana* intracellular amastigotes was evaluated at concentrations 486, 162, 54, 18, 6, 2, 0.66, 0.22, 0.072, 0.024 and 0.008 μg/ml of blank chitosan-TPP nanoparticles, blank chitosan-dextran sulphate nanoparticles, AmB loaded chitosan-TPP nanoparticles (AmB equivalent) and AmB loaded chitosan-dextran sulphate nanoparticles (AmB equivalent). The anti-leishmanial activity was evaluated in RPMI 1640 at two pH values (7.5 and 6.5). Pure AmB, AmBisome® and chitosan solution (HMW chitosan) was included in this experiment for comparison. PEMs were used as a macrophage model of intracellular amastigotes. See Chapter 2, section 2.8.7 for full details.

Similarly, the host cell dependence of the anti-*L. major* amastigotes activity of chitosan formulations (blank chitosan-TPP nanoparticles, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles) was evaluated by using two further host cell types (bone marrow macrophages (BMMs) and differentiated THP-1 cells). See Chapter 2, section 2.8.8 for full details.
4.2.6. Evaluation of the *in vivo* anti-leishmanial activity of chitosan formulations

A pre-toxic study of AmB loaded nanoparticles was done before starting the treatment. This toxic study was done by using female BALB/c mice aged 6 to 8 weeks, at 18-20g (Charles River, UK) and these mice were injected intravenously with 100 μL of AmB loaded chitosan-TPP nanoparticles or AmB loaded chitosan-dextran sulphate nanoparticles starting with concentration 20 mg/kg of AmB and then 2-fold decrease. All mice monitored closely and immediately after administration and then regularly until 48 hours post-dose for two weeks. The safest doses were chosen for the treatment; 5 mg/kg of AmB for AmB loaded chitosan-TPP nanoparticles and 10 mg/kg of AmB loaded chitosan-dextran sulphate nanoparticles.

### 4.2.6.1. *In vivo* L. major model of CL

Female BALB/c mice aged 6 to 8 weeks, at 18-20g, were purchased from Charles River Ltd. These mice were maintained under specific conditions (they were kept in controlled rooms with humidity of 55% and temperature of 26°C and fed water and rodent food ad libitum). *Luciferase-expressing L. major JISH118 (Ppy RE9H+L. major JISH118)* amastigotes were harvested and isolated from mouse skin lesions previously infected with *Leishmania* promastigotes (at a low passage number). Harvested amastigotes were transformed to promastigotes by keeping them at 26°C in Schneider’s insect medium + 10% HiFCS. Promastigotes were passaged every week and used at a low passage number (≤3) to infect experimental mice due to the potential decrease in virulence with increasing passage number and extended culture (166).

For this study, mice were shaved and then infected with 200 μL of 4x10^7 of stationary-phase luciferase-expressing *L. major JISH118 (Ppy RE9H+L. major JISH118)* promastigotes subcutaneously on the rump above the tail. After 7 days of infection, small nodules started to be visible at the site of injection and the lesion size was recorded daily by using a digital calliper; 10 days post infection the lesions measured 5 mm approximately in diameter. The infected
mice were allocated in 8 groups (5 mice in each group) with similar average lesion diameters (P >0.5, one-way-ANOVA) after which, the administration of formulations was started as described below:

*in vivo* experiment 1

**Group 1**  
Negative control: untreated, uninfected

**Group 2**  
The positive control group (G2): mice received 10 doses of 100 μL of paromomycin at a dose of 50 mg/kg intraperitoneally (i.p.) for 10 consecutive days, a regimen with proven efficacy in this CL model (77, 202)

**Group 3**  
Group 3 (G3): mice received 5 doses of 100 μL of AmBisome® (size= 70-80 nm) (272, 273), 10 mg/kg intravenously (i.v.) over 10 days, alternate day dosing on days 0, 2, 4, 6, and 8.

**Group 4**  
Group 4 (G4): mice received 5 doses of 100 μL of blank chitosan-TPP nanoparticles (equivalent to AmB-loaded) intravenously (i.v.) over 10 days, alternate day dosing

**Group 5**  
Group 5 (G5): mice received 5 doses of 100 μL of AmB loaded chitosan-TPP nanoparticles (5 mg/kg of AmB) intravenously (i.v.) over 10 days, alternate day dosing

**Group 6**  
Group 6 (G6): mice received 5 doses of 100 μL of blank chitosan-dextran sulphate nanoparticles (equivalent to AmB-loaded) intravenously (i.v.) over 10 days, alternate day dosing

**Group 7**  
Group 7 (G7): mice received 5 doses of 100 μL of AmB loaded chitosan-dextran sulphate nanoparticles (10 mg/kg of AmB) intravenously (i.v.) over 10 days, alternate day dosing

**Group 8**  
Group 8 (G8): mice received 5 doses of 100 μL of chitosan nanoparticle vehicles (water) intravenously (i.v.) for over 10 days, alternate day dosing

At day 9 (one day after the last dose was administered), the experiment was terminated, mice were humanely killed and skin samples were harvested by surgical removal from the areas containing the localized CL lesion and non-CL-infected skin on the back (control site) of the same mouse (Fig 4.1), stored
at -80°C for further experiments (the biodistribution of AmB and for qPCR and determination of burden). Treatment efficacy was evaluated by measuring the lesion size progression and parasite load (bioluminescence signal).

A second, repeated in vivo experiment was conducted with 35 mice were used (in vivo experiment 2). This experiment was performed for reproducibility and to study the dose-response effect of AmB loaded chitosan-TPP nanoparticles. Ten days post infection, the lesions measured 5 mm approximately in diameter and mice were allocated to 7 different experimental groups to ensure comparable lesion sizes in each group (5 mice in each group).

Mice were then treated for 10 days, receiving injections containing one of the following regimens:

Group 1  Control group (G1): untreated, infected mice

Group 2  The positive control group (G2): mice received 10 doses of 100 μL of paromomycin 50 mg/kg intraperitoneally (i.p.) for 10 consecutive days, a regimen with proven efficacy in this CL model (77, 202)

Group 3  mice received 5 doses of 100 μL of AmBisome® 10 mg/kg intravenously (i.v.) over 10 days, alternate day dosing on days 0, 2, 4, 6, and 8.

Group 4  mice received 5 doses of 100 μL of AmB loaded chitosan-TPP nanoparticles (5 mg/kg of AmB) intravenously (i.v.) over 10 days, alternate day dosing

Group 5  mice received 5 doses of 100 μL of AmB loaded chitosan-TPP nanoparticles (2.5 mg/kg of AmB) intravenously (i.v.) over 10 days, alternate day dosing

Group 6  mice received 5 doses of 100 μL of AmB loaded chitosan-TPP nanoparticles (1.25 mg/kg of AmB) intravenously (i.v.) over 10 days, alternate day dosing

Group 7  mice received 5 doses of 100 μL of blank chitosan-TPP nanoparticles (equivalent to 5 mg/kg AmB-loaded) intravenously (i.v.) over 10 days, alternate day dosing
After which the experiment was carried out as previously described.

Both blank and AmB loaded chitosan nanoparticles were suspended in distilled water, characterizes (size, charge, AmB loading) and used freshly for the in vivo study.

- Fungizone (a conventional deoxycholate formulation of AmB) was not included in both in vivo experiments as controls, because Wijnant et al (2017) found that the highest tolerated dose of Fungizone was 1 mg/kg/i.v. (which did not cause acute toxicity to BALB/c mice) and demonstrated that Fungizone (1 mg/kg/QAD for 10 days; i.v.) did not cause a significant reduction in lesion sizes or parasite load in murine (BALB/c) models of L. major (170).

**Figure 4.1.** Schematic representation of skin samples used in the study (166)

**4.2.6.2. Measurement of lesion size**

The lesion size was measured daily using digital calipers by determining the width and length of the lesion and then calculating the average (mm). One-way ANOVA with post-hoc Tukey test was performed to analyse the statistical differences between the average diameters per group (166).
4.2.6.3. Measurement of the bioluminescence signal

The luciferase substrate; luciferin (D-Luciferin potassium salt, Xenogen, CA and Gold Biotechnology, St. Louis, MO) was injected (sc) into the mice at 10 mins before the acquiring of bioluminescent signal. After 7 mins of injection, the mice were anaesthetized by inhalation with 3% isoflurane with 100% oxygen at a flow rate of 2.5 l/min until no movement was shown (3 mins approx.). Mice were then imaged and the images were acquired by using a Living Image software (version 4). Emitted photons were gathered by auto acquisition with a charge couple device (CCD) camera (PerkinElmer IVIS Spectrum In vivo Imaging System) using the medium resolution (medium binning) mode. A circular region of interest (ROI) encompassing the nodular area on the rump was drawn to quantify the bioluminescence, expressed as radiance and results were expressed in numbers of photons/sec (296).

4.2.6.4. Quantification of AmB in skin samples

Each frozen skin sample was cut into fine, long pieces, weighed and then inserted into microcentrifuge tubes. A spatula of 2 mm zirconium oxide beads (Next Advance, United Kingdom) (about 100 mg) was added with 1 ml of PBS to each tube. After which, the skin sample was homogenised in 3 cycles of 30 seconds of 6800 rpm using a Precellys 24 homogenizer (Bertin Technologies, France) to obtain a smooth homogenate. Then 100 ul of the homogenate was added to 250ul of a mixture of methanol: DMSO (84:16) plus 200 ng/ml tolbutamide (analytical standard; Sigma, United Kingdom) for drug extraction and tolbutamide was used for protein precipitation, in 96-well plates. Then, these 96-well plates were shaken for 10 mins at 200 rpm and centrifuged at 4°C at 6600 rpm for 15 mins. Two hundred microliters of supernatant were stored at -80°C until further analysis for quantification of AmB by HPLC as described previously in Chapter 3, in section 3.3.6.1. A calibration curve for the HPLC was prepared of AmB concentrations in untreated healthy skin homogenate (this homogenate was prepared as described by grinding the skin by using the zirconium oxide beads and the blender, shaking and centrifuging the samples) (170).
4.2.7. Skin permeation study by Franz diffusion cell (FDC) assay

25 female BALB/c mice (6 to 8 weeks old) at 18-20g, were shaved and infected with 200 μl of 4x10^7 of stationary phase *L. major JISH118* promastigotes subcutaneously on the rump above the tail. After 7 days of infection small nodules started to be visible at the site of injection and the lesion size was recorded daily by using a digital calliper. Ten days post infection, the lesions measured approximately 5 mm in diameter. The mice were humanely killed and 2 circular discs of skin (infected and uninfected skin- 15mm diameter approximately) were excised per mouse; the infected skin piece containing the *Leishmania* lesion was cut from the dorsal area above the tail and the uninfected piece above the lesion on the higher back of same mouse was collected (Fig 4.1). Forceps were used to gently remove fat and muscle from the skin samples and these samples were stretched carefully on Whatman filter papers. They were then mounted between the donor and receptor compartment of the Franz cell device (Fig 4.2.) and kept in place by a clamp. PBS with 2% hydroxypropyl-β-cyclodextrin (CD, Sigma, UK) was sonicated for 30 mins then added to the receptor compartment (as AmB is soluble in CD at 37 μg/ml) together with a small magnetic stirrer. The Franz cells were incubated in a warm water bath on a magnetic stirrer plate set at a speed of 800 rpm until the skin reached temperature 34°C. The Franz cells were examined for air bubbles and leakage (166). 100 μl of each formulation (Pure AmB as a control (3.96 ± SD mg of amphotericin B/ml), AmB loaded chitosan TPP nanoparticles (3.93 ± SD mg of amphotericin B/ml)) and AmB loaded chitosan dextran sulphate nanoparticles (3.84 ± SD mg of amphotericin B/ml)) was applied to each donor compartment. 100 μl of receptor fluid was taken at regular time intervals and was replaced with 100 μl of fresh PBS with 2% CD and stored at -80°C to be analysed later by HPLC to quantify AmB. After 24 h the experiment was terminated, and the Franz cells were disassembled. Donor chambers were washed with 1 ml of methanol: DMSO (84:16) which was then stored at -80°C for further AmB analysis by HPLC. A dry cotton swab was used to remove any residual AmB on the surface of skin. This was then stored at -80°C for further quantification of AmB. The skin samples were also stored at -80°C for further experiments. The cumulative amount of drug permeated as a
function of time was plotted and the linear portion of the graph was used to
calculate the flux and lag time (Fig 4.14). The permeability coefficient (Kp) was
calculated by using Equation 5.

Figure 4.2. Franz diffusion cell used for the permeation studies (166)

4.2.7.1. Quantification of AmB by HPLC

The amount of AmB in the wash was quantified by HPLC using parameters
described in Chapter 3 in section 3-2-7-; standard solutions of AmB were
achieved by diluting AmB in methanol: DMSO (84:16) solution.

The amount of AmB in the cotton swab was quantified by HPLC using
parameters described in Chapter 3 in section 3-2-7-. Firstly, the cotton swab
was soaked in 1 ml of methanol: DMSO (84:16) solution for 24 h and then
analysed. For the calibration curve, a dry cotton swab was soaked in 1 ml of
methanol: DMSO (84:16) solution for 24 h and then this solution was used to prepare a standard solution of AmB.

To extract AmB from the skin disc, the skin was homogenised as described below:

Each frozen skin sample was cut into fine, long pieces, weighed and then inserted into microcentrifuge tubes. A spatula-full of 2 mm zirconium oxide beads (Next Advance, United Kingdom) (about 100 mg) was added with 1 ml of PBS to each tube. The skin sample was homogenised in 3 cycles of 30 seconds at 6800 rpm using the Precellys homogeniser (Bertin Technologies) to obtain a smooth homogenate. 100 ul of homogenate was then added to a 250 µl of mixture of methanol: DMSO (84:16) plus 200 ng/ml tolbutamide (analytical standard; Sigma, United Kingdom) for drug extraction protein precipitation in 96-well plates. These 96-well plates were shaken for 10 mins at 200 rpm then centrifuged at 4°C at 6600 rpm for 15 mins. 200uL of supernatant was stored at -80°C until further analysis for quantification of AmB by HPLC as described previously in Chapter 3. A calibration curve for the HPLC was prepared with AmB concentrations in untreated healthy skin homogenate (this homogenate was prepared as described by grinding the skin using the zirconium oxide beads and the Precellys blender, shaking and centrifuging the samples) (170).

4.2.7.2. Fluorescence microscopy of skin sections post formulation application

To visualise the nanoparticles, formulations with rhodamine-labelled chitosan were prepared in a similar manner to unlabelled particles and then were characterised regarding size and zeta-potential using the Zeta-sizer and applied to infected and uninfected mouse skin using FDC (blank rhodamine-labelled chitosan-TPP nanoparticles equivalent to 3.93 ± SD mg of amphotericin B/ml loaded in AmB loaded chitosan TPP nanoparticles and blank rhodamine-labelled chitosan-dextran sulphate nanoparticles equivalent to 3.84± SD mg of amphotericin B/ml loaded in AmB loaded chitosan TPP nanoparticles) as described above. After the experiment, the cells were
dismantled and skin tissue fixed in tris-zinc fixative overnight as described by Accart et al (2014) (297). After 24 hours the skin samples were embedded in gelatin and immersed in OCT before storage at -80°C. Cryosections of 5 µm were cut using a cryostat (Leica CM1950).

For immunohistochemistry, the sections were defrosted and submerged in PBS (37°C) for 30 minutes to dissolve the gelatine after which they were submerged in PBS for 5 minutes, counterstained with DAPI and mounted in Prolong Gold (Thermofisher Scientific). Sections were examined using a Zeiss Axio Scan Z1 with a x 20 objective.

4.2.8. Statistical analysis.

For the efficacy experiment, ANOVA (1-way for parasite load and intralesional AmB levels, 2-way repeated measures for lesion size) followed by Tukey’s multiple comparison test were used. A $P$ value of <0.05 was considered statistically significant. All analyses were performed with GraphPad Prism version 7.02.
4.3. Results

4.3.1. Haemolysis activity of chitosan nanoparticles

The haemolytic activity of blank and AmB loaded chitosan TPP or dextran sulphate nanoparticles was clearly observed in a dose-dependent manner as shown in Fig 4.3. Pure AmB was significantly more haemolytic (around 18-fold) than both types of AmB loaded nanoparticles (p<0.05 by an extra sum-of-squares F test) (Table 4.4). On the other hand, AmBisome® is less toxic against RBCs than both types of AmB loaded nanoparticles (p<0.05 by an extra sum-of-squares F test) (Table 4.4).

Table 4.4. *In vitro* haemolytic activity of chitosan formulations after 1h of incubation

<table>
<thead>
<tr>
<th>Compound</th>
<th>Properties</th>
<th>RBC&lt;sub&gt;50&lt;/sub&gt; µg/ml</th>
<th>RBC&lt;sub&gt;90&lt;/sub&gt; µg/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amphotericin B (pure AmB)</td>
<td>Purity ≥95%, MW 924.1</td>
<td>11.3 ± 2</td>
<td>40.88 ± 5</td>
</tr>
<tr>
<td>AmBisome®</td>
<td>Liposomal AmB, Size= 70-80 nm</td>
<td>525.8 ± 6</td>
<td>1782 ± 8</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>MW=310-375 KDa</td>
<td>810.1 ± 7</td>
<td>3367 ± 9</td>
</tr>
<tr>
<td>Blank chitosan-TPP nanoparticles</td>
<td>Size= 67 ± 7 nm, Zeta potential= 28.5 ±1.9 mv</td>
<td>623.7 ± 6</td>
<td>3639 ± 10</td>
</tr>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td>Size= 69 ± 8 nm, Zeta potential= 25.5 ± 1 mv</td>
<td>209.5 ± 5</td>
<td>1129 ± 10</td>
</tr>
<tr>
<td>Blank chitosan-dextran sulphate</td>
<td>Size= 170 ± 9 nm, Zeta potential= -12.9 ± 3 mv</td>
<td>621.4 ± 8</td>
<td>3341 ± 16</td>
</tr>
<tr>
<td>sulphonate nanoparticles</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AmB loaded chitosan-dextran sulphate</td>
<td>Size= 174 ± 8 nm, Zeta potential= -11 ± 1 mv</td>
<td>202.8 ± 8</td>
<td>931.4 ± 8</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown). A statistically significant difference was found in RBC<sub>50</sub> values between AmB loaded chitosan nanoparticles and pure AmB (p<0.05 by an extra sum-of-squares F test).
Figure 4.3. Dose-response curves of haemolytic activity of chitosan nanoparticles (blank and AmB loaded nanoparticles) after 1h of incubation. Data are expressed as means ± SD from triplicates, statistically significant difference in RBC\textsubscript{50} values between pure AmB and AmB loaded nanoparticles (pure AmB is significantly more toxic AmB loaded nanoparticles) (P < 0.05 by an extra sum-of-squares F test).
4.3.2. Cytotoxicity of blank and AmB loaded chitosan nanoparticles against KB cells in RPMI (pH 7.5 and pH 6.5)

The cytotoxicity of blank and AmB loaded chitosan TPP or dextran sulphate nanoparticles against KB cells was clearly observed in a dose-dependent manner at two pH values (7.5 and 6.5) as shown in Fig 4.4. No significant difference in the cytotoxicity was observed for all formulations at pH of 7.5 and pH of 6.5 (pH did not have an effect on the cytotoxicity) (\(p>0.05\) by t-test) (Table 4.5). Both types of blank chitosan nanoparticles showed a significantly less cytotoxicity than AmB loaded chitosan nanoparticles (\(p<0.05\) by an extra sum-of-squares F test). AmB loaded chitosan TPP or dextran sulphate nanoparticles were significantly less toxic than pure AmB (6-fold less toxic against KB cells) (\(p<0.05\) by an extra sum-of-squares F test). However, no significant difference was observed in the cytotoxicity between AmB loaded nanoparticles and AmBisome® (\(p>0.05\) by an extra sum-of-squares F test) (Table 4.5).

<table>
<thead>
<tr>
<th>Compound</th>
<th>pH=7.5</th>
<th>pH=6.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Properties</td>
<td>LD(_{50}) µg/ml</td>
<td>LD(_{90}) µg/ml</td>
</tr>
<tr>
<td><strong>Podophyllotoxin</strong></td>
<td>0.7 ± 0.03</td>
<td>2 ± 0.3</td>
</tr>
<tr>
<td><strong>Amphotericin B (pure AmB)</strong></td>
<td>59 ± 2</td>
<td>228 ± 2</td>
</tr>
<tr>
<td><strong>AmBisome®</strong></td>
<td>401 ± 2</td>
<td>1568 ± 2</td>
</tr>
<tr>
<td><strong>HMW chitosan</strong></td>
<td>894 ± 4</td>
<td>2840 ± 3</td>
</tr>
<tr>
<td></td>
<td>Size= 67 ± 7 nm, Zeta potential= 28.5 ±1.9 mv</td>
<td>728 ± 2</td>
</tr>
<tr>
<td>----------------------</td>
<td>-----------------------------------------------</td>
<td>---------</td>
</tr>
<tr>
<td>Blank chitosan-TPP nanoparticles</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td>Size= 69 ± 8 nm, Zeta potential= 25.5 ± 1 mv</td>
<td>356 ± 5</td>
</tr>
<tr>
<td>Blank chitosan-dextran sulphate nanoparticles</td>
<td>Size= 170 ± 9 nm, Zeta potential= -12.9 ± 3 mv</td>
<td>949 ± 6</td>
</tr>
<tr>
<td>AmB loaded chitosan-dextran sulphate nanoparticles</td>
<td>Size= 174 ± 8 nm, Zeta potential= -11 ± 1 mv</td>
<td>366 ± 3</td>
</tr>
<tr>
<td>TPP</td>
<td>MW= 367.864 g/mol</td>
<td>840± 8</td>
</tr>
<tr>
<td>Dextran sulphate</td>
<td>MW= 40 KDa</td>
<td>&gt;1200</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown). Blank or AmB loaded chitosan nanoparticles had a similar toxicity at both pH values (6.5 and 7.5) toward KB-cells (p >0.05 by t-test). A statistically significant difference was found in LD$_{50}$ (50% lethal dose) values between AmB loaded chitosan nanoparticles and pure AmB (p<0.05 by an extra sum-of-squares F test).

Figure 4.4. Dose-response curves of the cytotoxicity against KB-cells. KB cells were cultured in the presence of different concentrations of chitosan formulations. The toxicity of drugs was measured after 72 hours by measuring the inhibition of metabolic activity. Values are expressed as % inhibition of KB cells relative to untreated controls. Statistically significant difference in LD$_{50}$ values between pure AmB and AmB loaded nanoparticles against KB-cells (AmB is significantly more toxic AmB loaded nanoparticles) (P <0.05 by an extrasum-of-squares F test)).
4.3.3. Activity of blank and AmB loaded chitosan nanoparticles against 
*L. major* and *L. mexicana* promastigotes in RPMI (pH 7.5 and pH 6.5)

Both chitosan solution and blank chitosan TPP nanoparticles were significantly more active at lower pH (6.5) than at higher pH (7.5) (*p* < 0.05 by t-test), chitosan solution was more active than blank chitosan-TPP nanoparticles at two pH values (*p* < 0.05 by t-test) (Fig 4.5.). Blank chitosan-dextran sulphate nanoparticles had no activity against *Leishmania* promastigotes up to a concentration of 486 µg/ml at two pH values. At both pH values (7.5 and 6.5) pure AmB, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles showed a similar anti-promastigote activity without a significant difference in their activity at these two pH values (*p* >0.05 by t-test). They were significantly more active against *Leishmania* promastigotes than AmBisome® (*p*<0.05 by an extra sum-of-squares F test) (Table 4.6). *L. major* promastigotes were more sensitive than *L. mexicana* to pure AmB, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles (*p*<0.05 by an extra sum-of-squares F test).
### Table 4.6. *In vitro* activity of chitosan formulations against promastigotes at two pH values

<table>
<thead>
<tr>
<th>Compound</th>
<th>Properties</th>
<th>pH=7.5</th>
<th>L. major**</th>
<th>L. mexicana**</th>
<th>pH=6.5</th>
<th>L. major**</th>
<th>L. mexicana**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>EC50 µg/ml</td>
<td>EC90 µg/ml</td>
<td>EC50 µg/ml</td>
<td>EC90 µg/ml</td>
<td>EC50 µg/ml</td>
<td>EC90 µg/ml</td>
</tr>
<tr>
<td>Amphotericin B (pure AmB)</td>
<td>Purity ≥95%, MW 924.1</td>
<td>0.06 ± 0.003</td>
<td>0.3 ± 0.02</td>
<td>0.2 ± 0.004</td>
<td>0.4 ± 0.03</td>
<td>0.06 ± 0.003</td>
<td>0.3 ± 0.02</td>
</tr>
<tr>
<td>AmBisome®</td>
<td>Liposomal AmB, Size= 70-80 nm</td>
<td>1 ± 0.08</td>
<td>7 ± 0.3</td>
<td>1.8 ± 0.1</td>
<td>7 ± 0.07</td>
<td>1.1 ± 0.08</td>
<td>7 ± 0.1</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>MW=310-375 KDa</td>
<td>106 ± 7</td>
<td>539 ± 31</td>
<td>141 ± 31</td>
<td>556 ± 5</td>
<td>7.1 ± 0.5</td>
<td>56 ± 4</td>
</tr>
<tr>
<td>Blank chitosan-TPP nanoparticles</td>
<td>Size= 67 ± 7 nm, Zeta potential= 28.5 ± 1.9 mv</td>
<td>164 ± 6</td>
<td>443 ± 10</td>
<td>185 ± 10</td>
<td>443 ± 0.8</td>
<td>28 ± 1.5</td>
<td>169 ± 11</td>
</tr>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td>Size= 69 ± 8 nm, Zeta potential= 25.5 ± 1 mv</td>
<td>0.08 ± 0.003</td>
<td>0.5 ± 0.02</td>
<td>0.3 ± 0.02</td>
<td>0.7 ± 0.02</td>
<td>0.06 ± 0.003</td>
<td>0.4 ± 0.02</td>
</tr>
<tr>
<td>Blank chitosan-dextran sulphate nanoparticles</td>
<td>Size= 170 ± 9 nm, Zeta potential= -12.9 ± 3 mv</td>
<td>No activity up to 486</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AmB loaded chitosan-dextran sulphate nanoparticles</td>
<td>Size= 174 ± 8 nm, Zeta potential= -11 ± 1 mv</td>
<td>0.09 ± 0.003</td>
<td>0.4 ± 0.01</td>
<td>0.5 ± 0.02</td>
<td>1 ± 0.07</td>
<td>0.06 ± 0.003</td>
<td>0.3 ± 0.02</td>
</tr>
<tr>
<td>TPP</td>
<td>MW = 367.864 g/mol</td>
<td>No activity up to 486</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dextran sulphate</td>
<td>MW = 40 KDa</td>
<td>No activity up to 486</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data not shown). *Statistically significant differences were found for the EC50 values of chitosan or blank chitosan-TPP nanoparticles at pH=6.5 and pH=7.5 (p<0.05 by using t-test). **L. major promastigotes were significantly more susceptible to pure AmB and AmB loaded chitosan nanoparticles than L. mexicana ((p<0.05 by an extra sum-of-squares F test)). Pure AmB and AmB loaded chitosan TPP or dextran sulphate nanoparticles had a similar anti-leishmanial activity.
Figure 4.5. Dose-response curves of the activity of blank and AmB loaded chitosan nanoparticles against *Leishmania* promastigotes at two pH values. A: *L. major*; B: *L. mexicana*. Promastigotes were cultured in the presence of different concentrations of chitosan nanoparticles. The activity of drugs was measured after 72h by the resazurin solution. Values are expressed as % inhibition of promastigotes relative to untreated controls. No statistically significant difference was observed in EC$_{50}$ values of AmB loaded chitosan nanoparticles and pure AmB against *L. mexicana* or *L. major* promastigotes ($p>0.05$ by t-test).
4.3.4. Activity of blank and AmB loaded chitosan nanoparticles against 
*L. major* and *L. mexicana* amastigotes infecting PEMs

Both chitosan solution and blank chitosan-TPP nanoparticles were significantly more active at lower pH (6.5) than at higher pH (7.5) (*p* < 0.05 by t-test) (Fig 4.6.). Chitosan solution was more effective against amastigotes than blank chitosan-TPP nanoparticles at higher pH of 7.5 (*p* < 0.05 by an extra sum-of-squares F test), However, both have a similar activity against *Leishmania* amastigotes at lower pH of 6.5 (*p* >0.05 by an extra sum-of-squares F test). Blank chitosan-dextran sulphate nanoparticles had no activity against *Leishmania* amastigotes to concentration up to 486 µg/ml at two pH values. Pure AmB, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles showed similar anti-amastigotes activity against both *L. major* and *L. mexicana* amastigotes at two pH values (7.5 and 6.5) without a significant difference in their activity at these two pH values (*p* >0.05 by t-test) and they were significantly more effective against *Leishmania* amastigotes than AmBisome® (*p* <0.05 by an extra sum-of-squares F test). Pure AmB and AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles showed higher anti-amastigote activity against *L. major* than *L. mexicana* (*p* <0.05 by an extra sum-of-squares F test) (Table 4.7).
Table 4.7. *In vitro* activity of chitosan formulations against intracellular amastigotes at two pH values

<table>
<thead>
<tr>
<th>Compound</th>
<th>Properties</th>
<th>pH=7.5*</th>
<th>pH=6.5*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L. major**</td>
<td>L. mexicana**</td>
<td>L. major**</td>
</tr>
<tr>
<td></td>
<td>EC_{50} µg/ml</td>
<td>EC_{90} µg/ml</td>
<td>EC_{50} µg/ml</td>
</tr>
<tr>
<td>Amphotericin B (pure AmB)</td>
<td>Purity ≥95%, MW 924.1</td>
<td>0.09 ± 0.003</td>
<td>0.5 ± 0.04</td>
</tr>
<tr>
<td>AmBisome®</td>
<td>Liposomal AmB, Size= 70-80 nm</td>
<td>1.2 ± 0.07</td>
<td>8 ± 0.3</td>
</tr>
<tr>
<td>HMW chitosan</td>
<td>MW=310-375 KDa</td>
<td>105 ± 7</td>
<td>1192±58</td>
</tr>
<tr>
<td>Blank chitosan-TPP nanoparticles</td>
<td>Size= 67 ± 7 nm, Zeta potential= 28.5 ±1.9 mv</td>
<td>162 ± 10</td>
<td>828 ± 43</td>
</tr>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td>Size= 69 ± 8 nm, Zeta potential= 25.5 ± 1 mv</td>
<td>0.14 ± 0.009</td>
<td>1 ± 0.09</td>
</tr>
<tr>
<td>Blank chitosan-dextran sulphate nanoparticles</td>
<td>Size= 170 ± 9 nm, Zeta potential= -12.9 ± 3 mv</td>
<td>No activity up to 486</td>
<td></td>
</tr>
<tr>
<td>AmB loaded chitosan-dextran sulphate nanoparticles</td>
<td>Size= 174 ± 8 nm, Zeta potential= -11 ± 1 mv</td>
<td>0.16 ± 0.008</td>
<td>1.4 ± 0.02</td>
</tr>
<tr>
<td>TPP</td>
<td>MW= 367.864 g/mol</td>
<td>No activity up to 486</td>
<td></td>
</tr>
<tr>
<td>Dextran sulphate</td>
<td>MW= 40 KDa</td>
<td>No activity up to 486</td>
<td></td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data not shown). *Statistically significant differences were found for the EC_{50} values of chitosan or blank chitosan TPP nanoparticles at pH=6.5 and pH=7.5 (p<0.05 by using t-test). ** L. major amastigotes were significantly more susceptible to pure AmB and AmB loaded chitosan nanoparticles than L. mexicana ((p<0.05 by an extra sum-of-squares F test)). Pure AmB, AmB loaded chitosan TPP and dextran sulphate nanoparticles had a similar anti-leishmanial activity.
Dose-response curves of the activity of blank and loaded chitosan nanoparticles against *Leishmania* amastigotes at two pH values. A: *L. major*, B: *L. mexicana*. PEMs were infected with stationary-phase promastigotes and exposed to various concentrations of chitosan and its derivatives, followed by microscopic counting of the number of infected macrophages*. Values are expressed as % inhibition of infection relative to untreated controls. No statistically significant difference was observed in EC\textsubscript{50} values of AmB loaded chitosan nanoparticles and pure AmB against *L. mexicana* or *L. major* amastigotes at pH=6.5 or pH=7.5 (p>0.05 by t-test).
4.3.5. Host cell dependence of the anti-leishmanial activity of chitosan nanoparticles at pH of 6.5

EC$_{50}$ and EC$_{90}$ values of blank chitosan-TPP nanoparticles, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles against amastigotes infecting three different macrophage populations are summarized in Table 4.8. There was a significant difference in the activity of chitosan formulations depending on the type of macrophage; as blank chitosan-TPP nanoparticles, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles were significantly more active against intracellular amastigotes in PEMs and BMMs compared to differentiated THP-1 cells (p<0.05 by an extra sum-of-squares F test) (Table 4.8).
Table 4.8. Activity of chitosan formulations against *L. major* amastigotes in three different macrophage cultures after 72 h at pH of 6.5

<table>
<thead>
<tr>
<th>Host cell / infection rate % at 24 h</th>
<th>Pure AmB</th>
<th>AmB loaded chitosan-dextran sulphate nanoparticles</th>
<th>AmB loaded chitosan-TPP nanoparticles</th>
<th>blank chitosan-TPP nanoparticles</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEMs / &gt; 80%</td>
<td>EC50 µg/ml</td>
<td>EC50 µg/ml</td>
<td>EC90 µg/ml</td>
<td>EC50 µg/ml</td>
</tr>
<tr>
<td>0.08 ± 0.01</td>
<td>0.4 ± 0.1</td>
<td>0.08 ± 0.004</td>
<td>0.4 ± 0.1</td>
<td>0.09 ± 0.004</td>
</tr>
<tr>
<td>BMMs / &gt; 80%</td>
<td>0.09 ± 0.02</td>
<td>0.6 ± 0.1</td>
<td>0.09 ± 0.02</td>
<td>0.6 ± 0.1</td>
</tr>
<tr>
<td>THP-1/ &gt; 80%</td>
<td>0.2 ± 0.05</td>
<td>3.4 ± 0.4</td>
<td>0.2 ± 0.06</td>
<td>3.3 ± 0.3</td>
</tr>
</tbody>
</table>

Experiments were conducted in quadruplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown), statistically significant difference in EC50 as chitosan formulations were significantly more active in PEMs and BMMs compared with THP-1 cells (*p*<0.05 by an extra sum-of-squares F test). % infection rate gives the percentage of infected macrophages.
4.3.6. *In vivo* anti-leishmanial activity (intravenous route route)

We assessed the efficacy of blank and AmB loaded chitosan nanoparticles (two types) in murine models of CL caused by *L. major*, by analysing the lesion sizes and bioluminescence signal progression among the groups.

4.3.6.1. *In vivo* experiment 1

4.3.6.1.1. Evaluation of the lesion size progression

Fig 4.7. shows the progression of the mean lesion size for each group as a function of time. Blank chitosan-dextran sulphate nanoparticles and nanoparticles vehicles did not cause any reduction in the progression of the lesion size compared to the untreated controls. Both AmBisome® (10 mg/kg/QAD for 10 days; i.v.) and blank chitosan-TPP nanoparticles reduced the lesion size at the end of the treatment with 36% and 34% respectively, reduction compared to the untreated controls without a significant different in their efficacy (*p>*0.05 by one-way ANOVA). AmB loaded chitosan-TPP nanoparticles (5 mg/kg/QAD for 10 days; i.v.) were the most effective compared with other chitosan formulations and caused a 87% reduction of lesion sizes and was significantly more effective than AmBisome® with 2.4 times greater activity (*p*<0.05 by one-way ANOVA). There was no significant difference in the anti-leishmanial efficacy between AmB loaded chitosan-TPP nanoparticles (5 doses) and paromomycin (50 mg/kg, 10 doses, positive control, 10 doses) with 87% and 93% respectively, reduction of lesion sizes (*p>*0.05 by one-way ANOVA).

Group 7 received one dose of AmB loaded chitosan-dextran sulphate nanoparticles (10 mg/kg, i.v.) for the reason that the day following this dose, mice looked unwell and showed signs of a piloerection and weight loss. Therefore, no more doses were administered. After two days, two mice had died and without any signs of potential CL-related mortality such as severe ulceration, dissemination of the lesion. We just kept monitoring the lesion sizes of the other three mice.
Figure 4.7. Amphotericin B nanoparticles efficacy in the lesion cure model in BALB/c mice infected with luciferase-expressing *L. major* parasites. Female BALB/c mice were infected with stationary-phase promastigotes in the rump above the tail (n = 5 per group). At 10 days post-inoculation, animals presenting with CL nodules were dosed with paromomycin (G2) as a positive control (50 mg/kg/QD for 10 days; i.p.), AmBisome® (G3) as a comparison group (10 mg/kg/QAD for 10 days; i.v.), blank chitosan-TPP nanoparticles equivalent to AmB loaded nanoparticles (G4) (QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G5) (5 mg of AmB/kg/QAD for 10 days; i.v.), blank chitosan-dextran sulphate nanoparticles equivalent to AmB loaded nanoparticles (G6) (QAD for 10 days; iv), AmB loaded chitosan-dextran sulphate nanoparticles (G7) (10 mg of AmB/kg/one dose; i.v.) or the nanoparticles vehicle (G8) (distilled water, QAD for 10 days; i.v.). (G1) represents untreated infected group. During treatment, lesion size was measured daily. The average lesion size represents the mean ± SD. ANOVA (1 way for parasite load and repeated measures for lesion size) followed by Turkey's multiple-comparison tests was used to compare outcomes among the groups. A p-value < 0.05 was considered statistically significant ((*) p<0.05, (**) p<0.05 and (***) p>0.05). (A) represents mean lesion size progression in function of time since the start of treatment, (B) represents the mean lesion size at day 9 (one day after the last dose was administered), (C) represents the % reduction in lesion size compared with G1 (untreated infected group) at day 9, (D) represents images of untreated group on day 9 (lesions are circled) and (E) represents images of G4 on day 9 (infection sites are circled and it is clear the healing effects of treatment on the lesions).
4.3.6.1.2. Evaluation of the parasite load (bioluminescent signal)

Bioluminescence signal progression in all treatment groups is shown in Fig 4.8. The reduction of parasite loads in the skin followed a similar trend of the lesion size with a good correlation between lesion size and bioluminescent signal identified by Pearson correlation coefficients (using GraphPad Prism). AmB loaded chitosan-TPP nanoparticles (5 mg/kg/QAD for 10 days; i.v.) were the most effective compared with other chitosan formulations with 99% reduction in parasite loads (bioluminescent signal) at the end of the treatment compared to the untreated controls and with similar reduction to the Group 2 treated with the positive control (paromomycin, 50 mg/kg, 10 doses; i.p.) ($p>0.05$ by one-way ANOVA). There was no significant difference between AmBisome® (10 mg/kg/QAD for 10 days; i.v.) and blank chitosan-TPP nanoparticles with 72% and 62% respectively, reduction ($p>0.05$ by one-way ANOVA). Blank chitosan-dextran sulphate nanoparticles did not cause any reduction in the signal at the end of treatment. We did not image Group 7 as the mice did not look healthy to be anaesthetized and imaged.
Figure 4.8. Amphotericin B nanoparticles efficacy as measured by the bioluminescence signal (parasite load) at the infection site in BALB/c mice infected with luciferase-expressing L. major parasites. Female BALB/c mice were infected with stationary-phase promastigotes in the rump above the tail (n = 5 per group). At 10 days post-inoculation, animals presenting with CL nodules were dosed with paromomycin (G2) as a positive control (50 mg/kg/QD for 10 days; i.p.), AmBisome® (G3) as a comparison group (10 mg/kg/QAD for 10 days; i.v.), blank chitosan-TPP nanoparticles equivalent to AmB loaded nanoparticles(G4) (QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G5) (5 mg of AmB/kg/QAD for 10 days; i.v.), blank chitosan-dextran sulphate nanoparticles equivalent to AmB loaded nanoparticles (G6) (QAD for 10 days; i.v.), or the nanoparticles vehicle (G8) (distilled water, QAD for 10 days; i.v.). (G1) represents untreated infected group. The bioluminescence signal was measured three times: start of treatment, after two doses of treatment and lastly on the day after the administration of the last dose. The data represents the mean ± standard error. ANOVA (1 way for parasite load and repeated measures for lesion size) followed by Turkey’s multiple-comparison tests was used to compare outcomes among the groups. A p-value < 0.05 was considered statistically significant ((*) p<0.05, (**) p<0.05 and (***) p>0.05). (A) represents the bioluminescence signal in function of time since the start of treatment, (B) represents the bioluminescence signal on the day after the administration of the last dose (day 9). (C) represents the % reduction in the signal compared with G1 (untreated infected group) at day 9. (D) represents the correlation between lesion size and the bioluminescence signal on the day after the administration of the last dose and (E) represents the bioluminescent images of mice on day 9 (24 h after the last drug dose administration). Emitted photons were gathered by auto acquisition with a charge couple device (CCD) camera (PerkinElmer IVIS Spectrum In vivo Imaging System) using the medium resolution (medium binning) mode.
4.3.6.1.3. Intralesional amphotericin B levels

We measured the levels of the active compound (AmB) within the infected lesion (rump skin) and control skin (uninfected skin, back skin) at the end of the experiment (Fig 4.9.). AmB levels were significantly higher (between 13 and 20-fold) in lesions sites (rump skin) compared to control skin (uninfected skin, back skin) in both Group 3 and Group 5 (p<0.05 by one-way ANOVA). After multiple dosing of either AmBisome® (G3, 10 mg/kg/QAD for 10 days; i.v.) or AmB loaded chitosan-TPP nanoparticles (G5, 5 mg of AmB/kg/QAD for 10 days; i.v.), intralesional AmB levels were significantly lower (6.8-fold) in Group 3 than in Group 5 (p<0.05 by one-way ANOVA). We could not detect any AmB levels as expected in samples from untreated group (G1) and positive control (G2).

Figure 4.9. Multiple dose skin pharmacokinetics of AmB loaded chitosan-TPP nanoparticles and AmBisome®. L. major-infected BALB/c mice received intravenous doses of AmBisome® (G3, 10 mg/kg/QAD for 10 days; i.v.) and AmB loaded chitosan-TPP nanoparticles (G5, 5 mg of AmB/kg/QAD for 10 days; i.v.). 24 hours after the last dosing, AmB levels in skin were determined. The CL lesion was localized on the rump, while the back skin of same mice was used as lesion-free, healthy control site. Each point represents the mean and standard error of the mean (n=5 per group). (A) represents intralesional AmB and (B) represents a comparison between infected and uninfected skin AmB concentration. The data represents the mean ± standard error. ANOVA followed by Turkey’s multiple-comparison tests was used to compare outcomes among the groups. A p-value < 0.05 was considered statistically significant ((*) p<0.05 and (**) p<0.05).
4.3.6.2. In vivo experiment 2 (dose-response effect)

4.3.6.2.1. Evaluation of the lesion size progression

Fig 4.10. shows the progression of the mean lesion size for each group as a function of time. Our data were similar and reproducible with previous in vivo experiment 1 regarding the efficacy of AmB loaded chitosan-TPP nanoparticles (5 mg of AmB/kg/QAD for 10 days; i.v.). In a mouse model of CL caused by L. major, AmB loaded chitosan-TPP nanoparticles efficacy showed a dose-response activity in reduction of lesion sizes at doses of 1.25, 2.5 and 5 mg of AmB/kg/QAD for 10 days; i.v., which caused 29%, 40% and 83% respectively, reduction in lesion sizes at the end of the treatment compared to the untreated controls. Similar to in vivo experiment 1, there was no significant difference in the efficacy of AmBisome®, blank chitosan-TPP nanoparticles and AmB loaded chitosan-TPP nanoparticles (2.5 AmB/kg/QAD for 10 days; i.v.) with 40%, 35% and 40% respectively, reduction of lesion sizes (p>0.05 by one-way ANOVA). Paromomycin (positive control) and AmB loaded chitosan-TPP nanoparticles (5 AmB/kg/QAD for 10 days; i.v.) were the most effective in reduction of lesion sizes and there was no statistically significant difference between the two treated groups with 89% and 83% respectively, reduction of lesion sizes at the end of the treatment compared to the untreated controls (p>0.05 by one-way ANOVA).
Figure 4.10. Amphotericin B nanoparticles efficacy in the lesion cure model in BALB/c mice infected with luciferase-expressing *L. major* parasites. Female BALB/c mice were infected with stationary-phase promastigotes in the rump above the tail (n = 5 per group). At 10 days post-inoculation, animals presenting with CL nodules were dosed with paromomycin (G2) as a positive control (50 mg/kg/QD for 10 consecutive days; i.p.), AmBisome® (G3) as a comparison group (10 mg/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G4) (5 mg of AmB/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G5) (2.5 mg of AmB/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G6) (1.25 mg of AmB/kg/QAD for 10 days; i.v.) and blank chitosan-TPP nanoparticles equivalent to AmB loaded
nanoparticles (5 mg/kg) (G7) (QAD for 10 days; i.v.). (G1) represents untreated infected group. During treatment, lesion size was measured daily. The average lesion size represents the mean ± standard error. ANOVA (1 way for parasite load and repeated measures for lesion size) followed by Turkey’s multiple-comparison tests was used to compare outcomes among the groups. A p-value < 0.05 was considered statistically significant (* p<0.05, ** p>0.05, *** p<0.05 and **** p>0.05). (A) represents mean lesion size progression in function of time since the start of treatment, (B) represents mean lesion size on the day after the administration of the last dose and (C) represents the % reduction in lesion size compared with G1 (untreated infected group) at day 9.
4.3.6.2.2. Evaluation of the parasite load (bioluminescent signal)

Bioluminescence signal progression in all treatment groups is shown in Fig 4.11. The reduction of parasite loads in the skin follows a similar trend of the lesion size with a good correlation between lesion size and bioluminescent signal identified by Pearson correlation coefficients (using GraphPad Prism). In a mouse model of CL caused by *L. major*, AmB loaded chitosan-TPP nanoparticles efficacy showed a dose-response activity in reduction of the parasite loads at doses of 1.25, 2.5, and 5 mg of AmB/kg/QAD for 10 days; i.v., which caused 48%, 75% and 99%, respectively, reduction in parasite loads (bioluminescent signal) at the end of the treatment compared to the untreated controls. Paromomycin and AmB loaded chitosan-TPP nanoparticles (5 mg of AmB/kg/QAD for 10 days; i.v.) were the most effective compounds with 99% reduction of the signal at the end of the treatment.

There was no significant difference in the efficacy of AmBisome® and AmB loaded chitosan-TPP nanoparticles (2.5 AmB/kg/QAD for 10 days; i.v.) in reducing parasite load with 80% and 75% respectively, reduction of bioluminescent signal (*p>*0.05 by one-way ANOVA). Blank chitosan-TPP nanoparticles caused a 65% reduction in parasite loads (bioluminescent signal).
Figure 4.11. Amphotericin B nanoparticles efficacy on the bioluminescence signal (parasite load) at the infection site in BALB/c mice infected with luciferase-expressing L. major parasites. Female BALB/c mice were infected with stationary-phase promastigotes in the rump above the tail (n = 5 per group). At 10 days post-inoculation, animals presenting with CL nodules were dosed with paromomycin (G2) as a positive control (50 mg/kg/QD for 10 consecutive days; i.p.), AmBisome® (G3) as a comparison group (10 mg/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G4) (5 mg of AmB/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G5) (2.5 mg of AmB/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G6) (1.25 mg of AmB/kg/QAD for 10 days; i.v.) and blank chitosan-TPP nanoparticles equivalent to AmB loaded nanoparticles (5 mg/kg) (G7) (QAD for 10 days; i.v.). (G1) represents untreated infected group. During treatment, lesion size was measured daily. The bioluminescence signal was measured three times: start of treatment, after two doses of treatment and lastly on the day after the administration of the last dose. The data represents the mean ± standard error. ANOVA (1 way for parasite load and repeated measures for lesion size) followed by Turkey’s multiple-comparison tests was used to compare outcomes among the groups. A p-value < 0.05 was considered statistically significant ((*) p<0.05, (**) p>0.05, (***) p<0.05 and (****) p>0.05). (A) represents the bioluminescence signal in function of time since the start of treatment, (B) represents mean the bioluminescence signal on the day after the administration of the last dose (day 9), (C) represents the % reduction in the signal compared with G1 (untreated infected group) at day 9. (D) represents the correlation between lesion size and the bioluminescence signal on the day after the administration of the last dose and (E) represents the bioluminescent images of mice on day 9 (24 h after the last drug dose administration). Emitted photons were gathered by auto acquisition with a charge couple device (CCD) camera (PerkinElmer IVIS Spectrum In vivo Imaging System) using the medium resolution (medium binning) mode.
4.3.6.2.3. Intraleional amphotericin B levels

We measured the drug levels of the active compound AmB within the infected lesion (rump skin) and control skin (uninfected skin, back skin) at the end of the experiment (Fig 4.12.). After multiple dosing of either AmBisome® (G3) or AmB loaded chitosan-TPP nanoparticles (G4 or G5 or G6), intra-leisional AmB levels were significantly lower (6.7-fold) in Group 3 (received AmBisome® at 10 mg/kg/QAD for 10 days) than in Group 4 (received AmB loaded chitosan-TPP nanoparticles at 5 mg of AmB/kg/QAD for 10 days; i.v.) (p<0.05 by one-way ANOVA). There was no significant difference in the intra-leisional AmB levels between Group 3 and Group 5 (received AmB loaded chitosan-TPP nanoparticles at 2.5 mg of AmB/kg/QAD for 10 days; i.v.) (p>0.05 by one-way ANOVA) and these levels of AmB were significantly higher in these two groups than in Group 6 (received AmB loaded chitosan-TPP nanoparticles at 1.25 mg of AmB/kg/QAD for 10 days; i.v.) (p<0.05 by one-way ANOVA). AmB levels were significantly higher in lesions sites (rump skin) compared to control skin (uninfected skin, back skin) (p<0.05 by one-way ANOVA) for all treated groups with AmB formulations. We could not detect any AmB levels as expected in samples from untreated group (G1) and positive control (G2).
Figure 4.12. Multiple dose skin pharmacokinetics of AmB loaded chitosan-TPP nanoparticles and AmBisome®. *L. major*-infected BALB/c mice received intravenous doses of AmBisome® (G3, 10 mg/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G4, 5 mg of AmB/kg/QAD for 10 days; i.v.), AmB loaded chitosan-TPP nanoparticles (G5, 2.5 mg of AmB/kg/QAD for 10 days; i.v.) and AmB loaded chitosan-TPP nanoparticles (G6, 1.25 mg of AmB/kg/QAD for 10 days; i.v.). 24 hours after the last dosing, AmB levels in skin were determined. The CL lesion was localized on the rump, while the back skin of same mice used as lesion-free, healthy control site. Each point represents the mean and standard error of the mean (n=5 per group). (A) represents intralesional AmB and (B) represents a comparison between infected and uninfected skin AmB concentration. The data represents the mean ± standard error. ANOVA followed by Turkey’s multiple-comparison tests was used to compare outcomes among the groups. A p-value < 0.05 was considered statistically significant (*) p<0.05, (**) p>0.05 and (*** p<0.05).
4.3.6.2.4. Dose concentration-response of AmB loaded chitosan-TPP nanoparticles in *L. major*-infected mice

The intralesional AmB levels were related to the dose levels of treatment with AmB loaded chitosan-TPP nanoparticles at concentrations (5 (G4) or 2.5 (G5) or 1.25 (G6) mg of AmB/kg/QAD for 10 days; i.v.) (Fig 4.13.a) and to the response (indicated by lesion size and parasite load) (Fig 4.13.b and 4.13.c, respectively). Fig 3d shows the nonlinear-fit sigmoidal dose-response curve plotting the logarithm of these intralesional AmB levels versus relative reductions in parasite load and lesion size compared to the untreated controls (0 mg/kg). Fig 4.13.e shows the % of relative reduction of lesion size and parasite load related to the doses per kg of AmB.

Correlation was strong between dose concentration and concentration response for relative reduction in parasite load and lesion size (identified by Pearson correlation coefficients (using GraphPad Prism)). We calculated ED$_{50}$ (The required dose to achieve 50% of maximum effect) and ED$_{90}$ (The required dose to achieve 90% of maximum effect) after plotting the logarithm of the dose level against percentage response (lesion size or parasite load). ED$_{50}$ and ED$_{90}$ were 2.5 and 8.9 mg/kg, respectively for lesion size. ED$_{50}$ and ED$_{90}$ were 1.3 and 3.8 mg/kg, respectively for parasite load (bioluminescent signal).
Figure 4.13. Dose concentration-response relationship of AmB loaded chitosan-TPP nanoparticles in experimental CL. *L. major*-infected BALB/c mice received intravenous doses of AmB loaded chitosan-TPP nanoparticles 0 or 1.25 or 2.5 or 5 of AmB/kg/QAD for 10 days (n = 5 per group). (a) represents the resulting intralesional amphotericin B levels, (b) lesion size, and (c) parasite load on the day after the last dose. (d) Outcomes are linked in a logarithmic-scale dose-response curve plotting drug concentrations against relative reduction in lesion size and parasite load. (e) is the relation between the dose in mg/kg and % of reduction of lesion size and parasite load. Each point represents the means ± SD (n = 5 per group). ANOVA followed by Turkey’s multiple-comparison tests was used to compare outcomes among the groups. A p-value < 0.05 was considered statistically significant.
4.3.7. *Ex vivo* permeability of *Leishmania*-infected skin in Franz diffusion cells

The permeability of uninfected and *L. major* infected skin for AmB loaded chitosan nanoparticles and fluorescence images of the nanoparticles distribution were evaluated in Franz diffusion cells. The cumulative concentration of AmB from AmB-loaded chitosan-TPP nanoparticles and AmB-loaded chitosan-dextran sulphate nanoparticles in the receptor compartment of Franz diffusion cells permeated as a function of time is shown in Fig 4.14. When applied as solution, pure AmB did not permeate through uninfected or infected skin throughout the 24 h permeation experiment. This was in contrast to the nanoparticle formulations, for which AmB could be detected in the receptor fluid. At the end of the 24 h experiment, both types of AmB loaded chitosan nanoparticles showed approximately a two-fold higher permeation of AmB through infected skin than uninfected skin (*p*<0.05 by t-test). AmB from AmB loaded chitosan-TPP nanoparticles permeated with almost two times more than from AmB loaded chitosan-dextran sulphate nanoparticles through both uninfected and infected skin (*p*<0.05 by t-test).
Figure 4.14. The cumulative amount of AmB permeated per surface area (ug/cm²) through uninfected BALB/c mouse skin (n=5) and *L. major* infected BALB/c mouse skin (n=5). Infected skin was more permeable to both types of AmB loaded chitosan nanoparticles than uninfected skin (*p*<0.05 by t-test). The use of AmB loaded chitosan-TPP nanoparticles enhanced AmB penetration through both healthy and infected skin in more amount than AmB loaded chitosan-dextran sulphate nanoparticles (*p*<0.05 by t-test).

Lag time, flux and permeability coefficients of the formulations are shown in Table 4.9. There was no significant difference in the lag time for both types of AmB nanoparticles between uninfected and infected skin (*p*>0.05 by t-test) and no significant difference was observed between AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles (*p*>0.05 by t-test). The flux was 2 times higher for both types of AmB loaded chitosan nanoparticles in infected skin compared to uninfected skin. The permeability coefficient was 1.75 and 2.5 times higher for AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles respectively in infected skin compared with uninfected skin. All the above indicated that *L. major* infection of the skin enhanced the permeation of both types of nanoparticles and the permeation of AmB nanoparticles is slow and poor.
Table 4.9. Flux, lag time and the permeability coefficient (kp) for AmB loaded chitosan nanoparticles

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Flux (μg/cm²/h)</th>
<th>Lag time (h)</th>
<th>Kp (cm/h)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Uninfected skin</td>
<td>Infected skin</td>
<td>Uninfected skin</td>
</tr>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td>0.06 ± 0.002</td>
<td>0.12 ± 0.005</td>
<td>20 ± 0.1</td>
</tr>
<tr>
<td>AmB loaded chitosan-dextran sulphate nanoparticles</td>
<td>0.04 ± 0.002</td>
<td>0.09 ± 0.002</td>
<td>20.5 ± 0.1</td>
</tr>
<tr>
<td>Pure AmB</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

Data expressed as mean +/- SD, n=5. No statistically significant difference of lag time was observed between uninfected and infected skin for both formulations (p>0.05 by t-test). Statistically significant differences of flux and kp were observed between uninfected and infected skin for both formulations (p<0.05 by t-test).
Table 4.10 shows the distribution of topical AmB from AmB loaded chitosan nanoparticles and pure AmB into healthy and *L. major* infected skin. After the 24 h permeation experiment, more than 90% of pure AmB stayed on the skin without any drug in the receptor fluid. Regarding both types of AmB loaded chitosan nanoparticles only a limited amount of AmB passed through the skin with 0.23% and 0.42% of AmB from applied AmB loaded chitosan-TPP nanoparticles through uninfected and infected skin respectively and 0.12% and 0.28% of AmB from applied AmB loaded chitosan-dextran sulphate nanoparticles through uninfected and infected skin respectively.

Table 4.10. Disposition of topically applied AmB loaded chitosan nanoparticles on healthy and *L. major* infected BALB/c mice skin using Franz diffusion cells

<table>
<thead>
<tr>
<th>Applied compounds</th>
<th>Average % recovered of AmB (±SD)</th>
<th>Uninfected skin</th>
<th><em>L. major</em> infected skin</th>
<th><em>P</em> value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure AmB</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>on skin (in wash and cotton swab)</td>
<td>94.65 ± 2</td>
<td>92.32 ± 1</td>
<td>&gt;0.05</td>
<td></td>
</tr>
<tr>
<td>in skin (extracted from skin homogenate)</td>
<td>5.35 ± 0.2</td>
<td>7.68 ± 0.2</td>
<td>&gt;0.05</td>
<td></td>
</tr>
<tr>
<td>through skin after 24h (in receptor fluid)</td>
<td>0</td>
<td>0</td>
<td>&gt;0.05</td>
<td></td>
</tr>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>on skin (in wash and cotton swab)</td>
<td>69.92 ± 1</td>
<td>61.49 ± 1</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>in skin (extracted from skin homogenate)</td>
<td>29.85 ± 1</td>
<td>38.09 ± 0.5</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>through skin after 24h (in receptor fluid)</td>
<td>0.23 ± 0.02</td>
<td>0.42 ± 0.05</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>AmB loaded chitosan-dextran sulphate nanoparticles</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>on skin (in wash and cotton swab)</td>
<td>81.65 ± 2</td>
<td>73.14 ± 2</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>in skin (extracted from skin homogenate)</td>
<td>18.23 ± 1</td>
<td>26.58 ± 1</td>
<td>&lt;0.05</td>
<td></td>
</tr>
<tr>
<td>through skin after 24h (in receptor fluid)</td>
<td>0.12 ± 0.02</td>
<td>0.28 ± 0.02</td>
<td>&lt;0.05</td>
<td></td>
</tr>
</tbody>
</table>

The total amount of AmB per Franz diffusion cell recovered at the end of the experiment was considered 100%. The amounts of AmB recovered from the different sites were expressed as a fraction of this amount. The average (±SD) percent for 5 infected mice is shown. *p* values were determined by a *t* test.

Fluorescence microscopy of skin sections showed no evidence for the penetration of rhodamine labelled chitosan-TPP nanoparticles (size= 72 ± 7 nm, Zeta potential= 22 ± 2) or rhodamine labelled chitosan-dextran sulphate nanoparticles (size= 174 ± 7 nm, Zeta potential= -14 ± 2) or rhodamine labelled chitosan solution in excised uninfected and *L. major* infected mouse skin. The
microscope study indicates that the nanoparticles act as drug delivery vehicle and release the AmB rather than permeating alongside the AmB molecules (Fig 4.15).

Figure 4.15. Fluorescence images of skin penetration (uninfected and L. major infected skin) of blank rhodamine labelled chitosan nanoparticles (A) and rhodamine labelled chitosan solution (B). We found the same scene for both types of nanoparticles and in both uninfected and infected skin. The red signals (refer to rhodamine labelled chitosan) indicated that the three formulations remained on the surface of skin.
4.4. Discussion

**In vitro haemolytic activity and cytotoxicity of chitosan formulations**

Haemolytic activity of chitosan formulations was determined by using freshly obtained human RBCs (295). Pure AmB showed a serious and significant toxic effect to RBCs after 1h of incubation. Loading the drug into both types of chitosan nanoparticles mitigated these effects, presumably by entrapping and retaining the AmB, allowing for slow release of drug. Similar findings have been reported for blank and AmB loaded chitosan- chondroitin sulphate nanoparticles (122). To evaluate the cytotoxicity of chitosan formulations in more details we found that both types of AmB loaded chitosan nanoparticles were around 6-fold less toxic than pure AmB against KB-cells and there was no significant difference in the cytotoxicity between these AmB loaded chitosan nanoparticles and AmBisome® for same reasons mentioned previously in terms of drug entrapment and slow release. Chitosan solution and blank chitosan nanoparticles (both types) showed a similar cytotoxicity against KB-cells and were significantly less toxic than AmB loaded nanoparticles. This data supports previous reports of less cytotoxicity of AmB loaded chitosan nanoparticles (136±11 nm, positive charge) compared to pure AmB against murine macrophages and the low toxicity of chitosan solution and blank nanoparticles against murine macrophages (122). Similarly, Jain et al reported that chitosan-coated AmB-loaded solid lipid nanoparticles (158.9±7.1 nm, positive charge) showed significantly less toxic effects against macrophages (J774A.1 cells in exponential growth phase) compared to amphotericin B deoxycholate (Fungizone) (259).

**In vitro anti-leishmanial activity of chitosan formulations**

Consistently with previous data in Chapter 2, lowering pH of RPMI medium from 7.5 to 6.5 increased by 7-20 times, the anti-leishmanial activity of chitosan solution and blank chitosan-TPP nanoparticles against *L. major* and *L. mexicana* promastigotes and amastigotes due to the greater ionisation at lower pH for both chitosan solution and blank chitosan-TPP nanoparticles (positive surface charge). As mentioned in the Chapter 2, increasing the
positive charge could enhance the chitosan antimicrobial activity by interacting with the negatively charged microbial membrane – in accordance with the first postulated mechanism of antimicrobial activity described in the introduction.

Regarding blank chitosan-TPP nanoparticles, they showed less anti-leishmanial activity than chitosan solution against *L. major* and *L. mexicana* promastigotes due to the lower positive charge of these nanoparticles as few numbers of the amino groups have been substituted by TPP groups. On the other hand, blank chitosan-TPP nanoparticles showed similar anti-leishmanial activity to chitosan solution against *L. major* and *L. mexicana* amastigotes at a lower pH due to the significant higher uptake of these nanoparticles by macrophages than chitosan solution (254).

Blank chitosan-dextran sulphate nanoparticles did not present any activity against *L. major* and *L. mexicana* promastigotes and amastigotes at both pH values- these nanoparticles have a negative surface charge as the positive amino groups on chitosan have been substituted by negatively charged sulphate groups.

However, AmB loaded chitosan nanoparticles (both types, positive or negative charged nanoparticles) showed a similar anti-leishmanial activity *L. major* and *L. mexicana* promastigotes and amastigotes at two pH values due to the high activity of AmB and this anti-leishmanial activity was similar to the activity of pure AmB and significantly higher than AmBisome®. Ribeiro *et al* (2014) reported that the anti-leishmanial activity of AmB loaded chitosan-chondroitin sulphate nanoparticles (136±11 nm, positive charge) was similar in comparison to pure AmB against *L. amazonensis* and *L. chagasi* promastigotes with similar EC$_{50}$ values to our study (83). Additionally, our EC$_{50}$ values against *L. major* and *L. mexicana* amastigotes were in accordance with another report that found the EC$_{50}$ values of chitosan-coated AmB-loaded solid lipid nanoparticles (158.9±7.1 nm , positive charge), AmBisome® and Fungizone were 0.022±0.07, 0.086±0.04, and 0.253±0.03 μg/ml, respectively, against *L. donovani* amastigotes infecting mouse macrophage cell line J774A.1 after 72 h of incubation (259).
Ribeiro et al (2014) showed that chitosan solution had EC$_{50}$ values of 66±1 and 71±1 µg/ml and blank chitosan nanoparticles had EC$_{50}$ values of 52 ±2 and 46 ±6 µg/ml against *L. amazonensis* and *L. chagasi* promastigotes, respectively and these values are different from EC$_{50}$ values in our study at two pH values (Table 4.8) and this could be explained as Ribeiro et al used different *Leishmania* species, 48h incubation of compounds with *Leishmania* and did not mention the pH of the experiment (83).

We were able to develop AmB loaded chitosan-TPP nanoparticles (69 ± 8 nm, positive surface charge) and AmB loaded chitosan-dextran sulphate nanoparticles (170 ± 9 nm, negative surface charge) which showed similar anti-leishmanial activity to pure AmB and higher activity than AmBisome® against promastigotes and amastigotes. These nanoparticles did not show significant haemolytic activity against RBCs and they were 6-fold less cytotoxic against KB-cells than pure AmB. This encouraged us to evaluate their *in vivo* anti-leishmanial activity using the mouse module.

**In vivo anti-leishmanial activity of chitosan formulations**

We assessed the efficacy of the chitosan formulations in murine models of CL caused by *L. major*, when administrated intravenously.

We evaluated the skin distribution of AmB following intravenous dosing with AmB loaded chitosan-TPP nanoparticles (1.25, 2.5 or 5 mg of AmB/ml/QAD for 10 days; i.v.) and AmBisome® (10 mg/kg/QAD for 10 days; i.v.). AmB accumulated in significant higher levels in the localized lesion compared to those in healthy skin tissue of the same infected mice; revealing the influence of CL skin infection on the drug accumulation. This could be explained by localized inflammatory immune response caused by *L. major* parasites multiplying within dermal macrophages of CL infected skin. Therefore, at the site of infection, the leaky vasculature could enhance permeability and retention effect of the drug and this may promote the local drug accumulation (170, 298) and these small nanoparticles could facilitate extravasation through the leaky capillaries in the inflamed lesion skin while in the healthy skin, the impairment in the extravasation (continuous endothelium with small vessel
pores of 6- to 12-nm diameter) could decrease the drug accumulation (299). Another explanation is because of the immune response to the CL, phagocytic monocytes immigrate from the bloodstream to the infection site (skin lesion) and these cells can act as drug reservoirs (16, 19, 300). Similar finding was reported by Wijnant et (2018) as AmB levels were 5- to 20-fold higher in L. major infected BALB/c mice skin than in healthy skin from same infected mice following dosing with AmBisome® or Fungizone.

However, AmB loaded chitosan-TPP nanoparticles (5 mg/ml/QAD for 10 days; i.v.) resulted in significant higher levels of AmB accumulation in infected skin than AmBisome® (10 mg/kg/QAD for 10 days; i.v.). There was no difference in the quantity of AmB in lesion skin following dosing of AmB loaded chitosan-TPP nanoparticles (2.5 mg/ml/QAD for 10 days; i.v.) and AmBisome® (10 mg/kg/QAD for 10 days; i.v.). Similarly, Sarwar et al (2017) reported that the oral administration of mannose-anchored thiolated chitosan amphotericin B nanocarriers (400 nm, positive surface charge) resulted in more AmB levels in the systemic circulation and higher pharmacokinetic parameters (AUC,t1/2 and Cmax) in comparison with same dose of AmBisome® or pure amphotericin B (301). Moreover, the same study showed mannose-anchored thiolated chitosan amphotericin B nanocarriers (400 nm, positive surface charge) promoted the cellular uptake of AmB by 70- and 23-fold in comparison to pure AmB and AmBisome®, respectively (301). This could be explained as the chitosan nanoparticles are able to retain the AmB inside the macrophages for the longer period of time compared AmBisome® and AmB and as mentioned that these macrophages could serve as reservoirs for the drug to target the infection site (301).

Blank chitosan-dextran sulphate nanoparticles did not cause any reduction in lesion size or parasite load (bioluminescent signal) of the infected mice. However, blank chitosan-TPP nanoparticles showed a similar activity in regard of lesion size and parasite load (bioluminescent signal) to AmBisome®. Ribeiro et al (2014) reported that blank chitosan-chondroitin sulphate nanoparticles (104±11 nm, positive charge) caused a significant reduction in lesion size of L. amazonensis infected BALB/c mice, when administrated intravenously (83).
AmB loaded chitosan-TPP nanoparticles (5 mg/ml/QAD for 10 days; i.v.) showed a high effectivity against CL in the mouse module and similar to the positive control (paromomycin, 50 mg/kg/QD for 10 consecutive days; i.p.), and caused a significant reduction on lesion development and parasite load (bioluminescent signal). Additionally, AmB loaded chitosan-TPP nanoparticles (2.5 mg/ml/QAD for 10 days; i.v.) resulted in a similar reduction of lesion size and parasite load (bioluminescent signal) to AmBisome® (10 mg/kg/QAD for 10 days; i.v.). The superior efficacy of AmB loaded chitosan-TPP nanoparticles (5 mg/ml/QAD for 10 days; i.v.) compared to AmBisome® (10 mg/kg/QAD for 10 days; i.v.) could be related to higher intralesional drug concentrations (described previously) and the effectivity of chitosan nanoparticles against CL.

Ribeiro et al (2014) reported that AmB loaded chitosan- chondroitin sulphate nanoparticles (136±11 nm, positive charge) caused significant reductions in the lesion size and in the parasite burden of L. amazonensis infected BALB/c mice, when administrated intravenously (1 mg/kg/day for 10 days) and were more active than pure AmB at same doses (121).

There was a good correlation between levels of intralesional AmB accumulation and the therapeutic outcomes of AmB loaded chitosan-TPP nanoparticles as the anti-leishmanial activity of AmB has a concentration-dependent response and this due to the concentration-dependency of AmB antimicrobial activity (302) and this consistent with Wijnant et al (2018) for AmBisome® in CL mouse module (170).

**Chitosan formulations for topical administration – skin penetration**

As we mentioned that topical treatment offers several advantages over systemic treatment regarding side effects, the direct target for infected lesions, less need for patient follow up and better compliance by the patients (303, 304, 305, 306). Thus, the aim was to develop topical nanoparticles formulations (positive and negative charged nanoparticles) containing AmB. There are four fundamental factors that control the efficacy of topical treatment of CL:

(i) The intrinsic efficacy of the compound against *Leishmania*
(ii) The permeability of the compound through the skin to reach the dermis where the *Leishmania* infected macrophages reside (30)

(iii) Disposition of the drug in the skin

(iv) The release of the active compound from formulation in the PV of infected macrophages in the dermis of infected skin (306)

Both AmB chitosan-TPP nanoparticles and AmB chitosan-dextran sulphate nanoparticles showed a high activity against *Leishmania* promastigotes and amastigotes. Therefore, we investigated their *in vitro* permeation characteristics through uninfected and *L. major* infected mice skin using Franz diffusion cells.

By using Franz diffusion cells, pure AmB did not permeate through uninfected or *L. major* infected skin. This is consistent with other reports (307, 308, 309, 310) and this could be explained as AmB is a big molecule (924 g/mol ) and is not soluble in water (307).

Briefly to optimise permeation, a given drug should comply with the following physicochemical properties :

- molecular weight < 500 g/mol
- log p between 1 – 3
- aqueous solubility > 1 mg/ml
- hydrogen bonding groups < 2.

Accordingly AmB is not a a good candidate for topical route, as its molecular weight is > 500 g/mol and log p of AmB is -0.66 and it is not the acceptable range for skin permeation (log P between 1-3) and has 12 H-bond donors and 18 H-bond acceptors (311, 312).This impermeability of AmB through healthy or infected skin clarified the unsuccessful treatment after the topical application of AmB on *L. major* infected mice (313). In addition, AmBisome® (liposomal AmB) was not efficient delivery topical systems for CL (314) and did not present a significant AmB skin deposition, *in vitro* study (Excised human skin from Caucasian female patients) (315).

*In vitro* permeation study showed a limited and slow permeation of AmB across healthy and infected mice skin when both types of AmB loaded chitosan
nanoparticles applied on the skin samples with a long lag time of about 20 h indicating a long time for the steady state flux to be reached (which indirectly means slow permeation across the stratum corneum). These data were confirmed by imaging the permeation of rhodamine labelled chitosan-TPP nanoparticles and rhodamine labelled chitosan-dextran sulphate nanoparticles across uninfected and L. major infected skin using laser microscope which showed that these nanoparticles stayed on the surface of skin. Our results were consistent with other reports; Vogt et al reported that most of applied 42–300 nm fluorescent silica nanoparticles stayed in the upper layers of the excised human skin using conventional fluorescence microscopy of skin sections(316).

Try et al observed a negligible penetration of poly (L-lactide-co-glycolide) nanoparticles with two sizes 70 and 300 nm in healthy male Swiss mice skin by using confocal laser scanning microscopical examination of skin biopsies while nanoparticles have been visualised in the epidermis in inflamed skin (inflammation induced by the application of oxazolone to develop atopic dermatitis like lesions) (317). Moreover, our data are in agreement with the study of Campbell et al who reported no penetration of flusospheres nanoparticles (carboxy-modified, fluorescent, polystyrene nanoparticles with three sizes 20, 100 and 200 nm) through pig skin and these nanoparticles remained in the top layers of the stratum corneum after 16 h of the application in Franz diffusion cells by using a laser scanning confocal microscopy (318). Similar observation regarding the limited permeation of chitosan nanoparticles was noticed by Nair et al, that curcumin-encapsulated chitosan nanoparticles with sizes ranged from 167.3 ± 3.8 nm to 251.5 ± 5.8 nm had a slow permeation and with low amounts using Franz diffusion cells through Strat-M® membrane (Strat-M is made of polyester sulfone arranged as multiple layers mimicking the skin structure including a tough outer layer manufactured by Merck) and the cumulative amount of curcumin permeated at 72 h was 34.3 ± 1.6 μg cm⁻² and 27.7 ± 1.7 μg cm⁻² for nanoparticles with sizes 251.5 and 167.3nm, respectively (319). Malli et al (2019) reported that the topical application of chitosan-Coated Poly (isobutyl cyanoacrylate) (size=187nm, zeta potential =53.8 mv) nanoparticles (prepared by anionic emulsion
polymerization method) gelified by pluronic F127 daily for 3 consecutive weeks to BALB/c mice infected with *L. major*, resulted in partial and not complete healing of lesion and could be due to a physical effect of the F127 hydrogel (220).

AmB loaded chitosan nanoparticles offered more permeation of AmB through infected than uninfected skin when applied topically and this was consistent with another report that showed more permeation of caffeine and ibuprofen through *L. major* infected than uninfected BALB/c mouse skin, using Franz diffusion cells (166). The same study reported no permeation of paromomycin sulphate through uninfected mice skin while a high permeation through *L. major* infected skin was observed using Franz diffusion cells (166). This could be explained as CL lesions cause a damage to the skin barrier and this alteration in skin could enhance the penetration of nanoparticles (320). Moreover, *Leishmania* infected skin is characterised by the presence of abundant inflammatory cells in the infection site and this could disarrange the consistency of the epidermal and dermal skin layers and by ulceration and necrosis (307, 310, 321). Trans-epidermal water loss (TEWL) was significantly higher in *L. major* infected skin and this reduced the barrier function of the skin and subsequently increased the accumulation of fluid in the interstitial spaces cause an oedema that could enhance the permeation of water-soluble compounds (307).

AmB loaded chitosan-TPP nanoparticles (size = 68 ± 7 nm, Zeta potential= 30 ± 2) presented more permeation of AmB than AmB loaded chitosan-dextran sulphate nanoparticles (size = 168 ± 7 nm, Zeta potential= -15.5 ± 2). Similarly, Try et al reported a higher penetration of smaller poly(L-lactide-co-glycolide) nanoparticles (70nm) than bigger ones (300 nm) in healthy male Swiss mice skin and could be explained as smaller sized nanoparticles can penetrate for more distance compared with bigger ones (317).

Another explanation of this higher penetration as the positive surface charge of chitosan-TPP nanoparticles could interact with negative charges in the skin and confirm close contact with the skin and make an occlusive barrier that enhance the hydration and this facilitates the nanoparticles permeation through the skin (322, 323).
All of the above regarding the limited and slow permeation of AmB from AmB loaded nanoparticles made these nanoparticles unsuitable candidates for topical administration. On this basis we did not pursue in vivo evaluation of the antileishmanial activity of topical route of these formulations.

In conclusion, AmB loaded chitosan-TPP nanoparticles showed efficient, stability properties and target oriented drug delivery system in an experimental model cutaneous leishmaniasis when administered by the i.v. route, these nanoparticles were significantly more active than AmBisome® against the murine model (female BALB/c mice) of L. major even with lower doses of these nanoparticles. AmB loaded chitosan-TPP nanoparticles can specifically target the CL lesions more than AmBsiome as they resulted in a higher concentration of AmB in the lesion sites in comparison to AmBisome®. However, Franz diffusion cell studies showed poor drug permeation into and through the skin of both types of AmB loaded chitosan nanoparticles suggesting that these formulations are not an appropriate candidate for topical treatment for CL. Our results indicate the need for more extensive studies using the intravenous route using different Leishmania species, different mammalian models and further extensive toxicity studies. Finally, skin samples from the in vivo study are stored for qPCR determination of parasite load and this work fell beyond the time line of this project.
5. Comparison of *in vitro* static and dynamic culture systems to evaluate the macrophages functions and the anti-leishmanial efficacy of chitosan formulations **

** research in this chapter was performed in collaboration with Alec O'Keeffe, who showed in a published paper (I am one of the co-authors) that the infection of macrophages by *L. major* was significantly reduced under slow medium flow and faster medium flow (to match the interstitial fluid flow rate in human skin) compared to macrophages under static conditions. The replication of *Leishmania* amastigotes and two functions of macrophages (phagocytosis and macropinocytosis) were also reduced under two media perfusion conditions, see publication, Appendix 2 (Paper 1).

Alec O'Keeffe completed his PhD on the development of novel predictive 2D and 3D *in vitro* models for ant-leishmanial drug testing, studying the activity and accumulation of anti-leishmanial drugs under these different flow conditions. Some of his results are included in the discussion for reference.

5.1. Media perfusion system: an introduction

The important effects of fluid flow (blood flow, interstitial flow, etc) on cell signalling and morphogenesis have been widely recognized. Cells in the mammalian body are residing in highly complex microenvironments and encounter many signals that vary in time and space. Tissues are in direct contact with moving body fluids, which encompass the haemolymphatic system, the digestive system and cerebrospinal fluid. These fluids play a significant role in the body cells such as the provision and delivery of nutrients, oxygen, cell signalling components and the removal of waste. The flow of blood and other bodily fluids within the body exerts mechanical stress on cells (324). Different rates of body fluid flow have been recorded, from fast plasma flow of 9.8 ml/min in the portal vein of the rat (325) to slow rates of 0.19 μl/min of interstitial fluid drainage from rat brains (326). 20% of the human body’s mass is estimated to be made up of interstitial fluid which is in all tissues, including skin; derived from the normal leakage of plasma from blood vessels
and has a similar structure to that of blood plasma (327). *Leishmania* amastigotes reside and survive in the phagolysosome of mammalian macrophages and in CL, these infected macrophages are exposed to interstitial fluid in the skin. The flow of interstitial fluid in uninfected human skin has been recorded to be in the order of 0.1–2 μm/s but this flow in CL infected skin has not been determined (327, 328, 329).

Most *in vitro* studies, in the *Leishmania* field (on drug discovery, host cell transport and immunology), have been conducted on macrophages in static culture, typically using 4-, 12-, 16-, 24-, 48-, and 96-well plates with a culture medium overlay. This static system does not provide the mechanical stress, and O₂ tension, amongst other things, to that of cells within a mammalian body (98). Consequently, a static system has a major limitation when evaluating cellular parameters *in vitro*, such as infection rate, drug activity, and macrophage functions such as phagocytosis and pinocytosis, offering a poor mechanistic understanding and predictive value (98, 324). Increasing the complexity of a culture system could produce, potentially, a more biologically relevant system. Additionally, the issues surrounding the use of animal models in terms of welfare, time and cost constraints, and the limits of non-human models in predicting outcomes in humans, make developing a more predictive *in vitro* culture system a high priority (324).

A first step is transforming static cultures systems to flow systems where the culture medium constantly flows, to imitate the flow conditions in the mammalian body (330). Microfluidic (Fig 5.1.) and macrofluidic systems (Fig 5.2.) are the main two types of media perfusion systems to conduct *in vitro* assays.

Many “microbioreactor” systems have been described for cell culture which range from laminar flow, membrane systems to rotating vessel systems. Most of these bioreactors require the use of particular seeding methods with narrow dimensional specifications (331, 332, 333). Microfluidic systems can be adjusted to mimic physiological conditions and deliver nutrients, dissolved gases and remove waste products. The advantage of a microfluidic system is, that less reagents are used overall helping to lower experimental costs.
However, microfluidic systems do have a number of disadvantages – they are typically very small and require significant technical ability and care when setting up (98). In microfluidic systems, just a few thousand cells can be seeded on the tiny culture surface (0.5–0.8 mm$^2$) and this low number of seeded cells cannot predict precisely the *in vivo* pathophysiology. Another drawback of these systems includes the “edge effect” in which a high proportion of cultured cells will be located on the outer circumference of the chamber. These cells will be organised differently as medium evaporates at a higher rate at the edges compared to the central area of the chamber, affecting cell seeding. An uneven cell layer can skew the results (334). Another disadvantage, when using micro systems, is that small hydrophobic molecules can be adsorbed by the material that either the chamber system or the connecting tubes are composed of (335, 336). A micro system will also have a high surface area to volume ratio and surface adsorption which will cause an increase in metabolic consumption rates and depletion of nutrition for the cells (335, 336, 337). Air bubble formation can pose problems within these systems, disrupting flow and affecting shear stress (338).

Macrofluidic systems (for example, Quasi Vivo, Kirkstall Ltd, Fig 5.2) offer many advantages over microfluidic systems - a higher volume of liquid is used which eases the preparation of low concentration compounds without wasting compounds through dilution. These systems can keep the shear stress consistently similar to the shear stress in most physiological environments (324). Moreover, macrofluidic systems can run for a longer time than micro system cultures and have a lower surface area to volume ratio, overcoming the major disadvantage of high metabolic consumption seen in micro systems (324).
Figure 5.1. Microfluidic system (A) The integrated perfusion culture micro-chamber array chip. (B) Enlarged view of a micro-chamber array unit (339).

Figure 5.2. Kirkstall LTD. Quasi Vivo 900 media perfusion system in use circulating RPMI 1640 media(340).
Our study used QV900, as described by O'Keeffe A et al (2018) with two flow rates, in which one rate mimics the interstitial tissue flow rate in the skin. Modelling of the flow rate at the cell surface and O\textsubscript{2} tension was made by a collaboration between The London School of Hygiene & Tropical Medicine (Alec O'Keeffe and Simon L Croft) and University of Glasgow (Lauren Hyndman and Sean McGinty) (100).

Here, this Chapter describes the impact of flow on host cell phagocytosis and macropinocytosis and how increasing the complexity of \textit{in vitro} model influences the anti-leishmanial activity of chitosan formulations (chitosan solution, blank chitosan-TPP -nanoparticles and AmB loaded chitosan-TPP nanoparticles) against intracellular \textit{L. major} amastigotes, these formulations showed a high \textit{in vitro} activity against \textit{L. major} amastigotes using static culture system (Chapter 4)
5.2. Material and methods

Kirkstall Ltd (Rotherham, UK), established in 2006 by Dr J Malcolm Wilkinson in 2006, has developed cell culture technology into a commercially available inter-connected cell culture system, known as Quasi Vivo®, by introducing flow into the cell cultures to increase physiological relevance and create more confidence in the data produced. The Quasi Vivo system includes QV500 (an individual chamber system) and QV900 and their specifications are summarised in Table 5.1.

Table 5.1. Specifications of QV500 and QV900 media perfusion system(340, 341, 342)

<table>
<thead>
<tr>
<th>Features</th>
<th>QV500</th>
<th>QV900</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chamber width</td>
<td>15 mm internal</td>
<td>15 mm internal</td>
</tr>
<tr>
<td>Chamber depth</td>
<td>10 mm from culture surface to top of chamber base</td>
<td>22 mm</td>
</tr>
<tr>
<td>Overall dimensions</td>
<td>23 mm height x 37 mm diameter</td>
<td>23 mm height x 37 mm diameter</td>
</tr>
<tr>
<td>Diameter of tubing</td>
<td>Inlet: 1/16” ID Outlet: 3/32” ID</td>
<td>Inlet: 1/16” ID Outlet: 3/32” ID</td>
</tr>
<tr>
<td>Volume of chamber</td>
<td>2 ml</td>
<td>4 ml</td>
</tr>
</tbody>
</table>

5.2.1. Preparation of chitosan solution and blank and AmB loaded chitosan nanoparticles

All nanoparticles in this study were prepared and characterised as described in chapter 3 in sections 3-2-1- and 3-2-2-. After freeze drying the nanoparticle
suspension, the white (blank nanoparticles) or yellow (AmB loaded nanoparticles) product was reconstituted in double distilled water (ddH₂O). The nanoparticles were then characterised by size, charge and AmB loading (see Chapter 3). A solution of HMW chitosan was prepared by dissolving 1 g in 100 ml of 1% (v/v) acetic acid solution at room temperature with continuous stirring for 24 hours until a clear solution was obtained. The pH of the solution was adjusted to ~ pH 6 by adding sodium hydroxide 2N (NaOH, Sigma, UK) solution using a pH meter (Orion Model 420A). The chitosan solutions were autoclaved (121 °C; 15 mins).

5.2.2. QV900 and media perfusion system

QV900 is a 6-chamber optical tray which can be connected together in any combination, providing a high degree of flexibility and the potential to culture cells in a defined set of conditions. QV900 is more suited to high-throughput testing than QV500. A 3D printed block (9mm) composed of Nylon 12 (Kirkstall Ltd) can be added to the chamber which will alter the depth of the chambers and can be used to adjust the level of oxygen and flow rates, the cells are subjected to. A peristaltic pump (Parker Hannifin, UK), external to the CO₂ incubator, continuously circulated culture media through the system is used. A constant flow rate of 360 μl/min of culture media was used. The cells (infected or uninfected macrophages) were cultured either at the base of a perfusion chamber or raised on 9 mm high inserts. This resulted in a cell surface flow rate of 1.33 x 10⁻⁹ at the base of the chamber or 1.17 x 10⁻⁷ (m/s) on an insert which is in line within the reported range for interstitial flow in the human skin (100).

5.2.3. Macrophages

Macrophages were plated on 12mm round glass coverslips (Bellco, US) placed in 24 well plates (Corning, UK) at a density of 4 x 10⁵ cells per well in RPMI-1640 media (PEMs and THP-1) or DMEM (BMMs) supplemented with 10% (v/v) HiFCS.

- THP-1 cells were incubated in RPMI 1640 plus 10% (v/v) HiFCS and 20 ng/ml phorbol 12-myristate 13-acetate (PMA; Sigma, UK) at 37°C
and 5% CO₂ for 72 h to induce maturation transformation of these monocytes into adherent macrophages.

5.2.4. Infection of macrophages by *L. major* promastigotes

Macrophages 4 x 10⁵/ml in RPMI-1640 media (PEMs and THP-1) or DMEM (BMMs) medium supplemented with 10% (v/v) HiFCS were plated in 24 well plates (Corning, UK) (1 ml per well) on 12mm round glass coverslips (Bellco, US) placed in 24 well and incubated for 24 hours at 37 °C in 5 % CO₂. After 24 hours, wells were washed by fresh culture medium to remove non-adherent cells. After washing, stationary phase *L. major* (MHOM/SA/85/JISH118) promastigotes were added into the wells at a ratio of 5:1 (5 parasites: 1 host). Plates were incubated for another 24 hours at 34 °C in 5 % CO₂. Subsequently, free parasites were removed by washing with the medium. One infected coverslip slide was fixed with 100 % methanol for 5 minutes and stained with 10 % Giemsa for 5 minutes. The number of infected macrophages per 100 macrophages was microscopically counted. If the initial infection was higher than 80 %, the assay was suitable for the experiments. Subsequently, two thirds of the glass coverslips were transferred to the media perfusion system (at the base of chamber or on the 9 mm insert) and maintained under flow conditions at a flow speed of 360 μl/min for 72 hours. The remaining coverslips were used for the static control.

5.2.5. Measurement of macrophage functions.

5.2.5.1. Phagocytosis

Phagocytosis by macrophages (PEMs, BMMs and THP-1) was initially evaluated using 0.5, 1 and 2 μm diameter fluorescent red labelled latex beads (carboxylate-modified polystyrene) (Sigma-Aldrich, UK) (343, 344). 2 μm beads were eventually selected as they showed maximal signal. Macrophages were infected with *L. major* promastigotes, then transferred to the three flow conditions as described above. To each well, 2μm beads (9.12 x 10⁷ latex beads/ml) were added and the cells were incubated for 0.5, 1, 2, 4 and 24 hours at 34 °C under the three different flow conditions. The experiment was terminated by washing the cells 4 times with ice-cold PBS pH 7.4 to remove
non-internalized latex beads, followed by the addition of 1 ml of 0.5% Triton X100 in 0.2 M NaOH to lyse the cells. Phagocytosis was quantified by the analysis of the cell lysate using a fluorescence plate reader (Spectramax M3, at excitation and emission wavelengths set at 575 and 610 nm), calibrated with standard solutions containing different number of latex beads in a cell lysate mixture. Uptake was expressed as the number of latex beads associated per mg of cellular protein, the protein content of the cell lysate being measured using a Micro BCA protein kit (Thermo Fisher, UK) assay as per supplier’s instructions. For control studies, 1 μg/ml cytochalasin D was used as a phagocytosis inhibitor (Sigma-Aldrich, UK) by incubation with macrophages for 2 hours prior to addition of the latex beads. Phagocytosis was completely inhibited after 0.5, 1, 2 and 4 hours of incubation with cytochalasin D and 90% after 24 hours.

5.2.5.2. Macropinocytosis

Macropinocytosis was measured using a fluorescence-labeled dextran dye (pHrodo Red dextran, average molecular weight of dextran 10,000 MW, Thermo Fisher, UK) (345). This dye has a pH-sensitive fluorescence emission that increases in intensity with increasing acidity while exhibiting a minimal fluorescence at neutral pH. Macrophages (PEMs, BMMs and THP-1) were infected with *L. major* promastigotes and then transferred to the three flow conditions as described above. Macrophages were washed 3 x by Live Cell Imaging Solution (Thermofisher, UK) and the cells were returned to RPMI 1640 + 10% hiFCS containing 40 μg/ml pHrodo Red dextran (1 ml for each well) and incubated at 34 °C / 5% CO₂ for 0.5, 1, 2, 4 and 24 hours under the three different flow conditions. At each time point, the cells were washed with Live Cell Imaging Solution and macropinocytosis was analysed by a Spectramax M3 at excitation and emission wavelengths set at 560 and 585 nm respectively. Chlorpromazine hydrochloride 10 µg/ml, a known inhibitor (Sigma-Aldrich, UK), was used as a control and was incubated with macrophages for 2 hours prior to addition of fluorescence-labeled dextran dye. Macropinocytosis was completely inhibited after 0.5, 1, 2 and 4 hours of incubation with chlorpromazine hydrochloride and by 90% after 24 hours.
5.2.6. Evaluation of the anti-leishmanial activity of chitosan solutions, blank and AmB loaded chitosan TPP nanoparticles in the media perfusion system at pH 6.5

PEMs were infected with *L. major* promastigotes, then transferred to the three flow conditions as described above. This experiment was conducted at pH 6.5. After 72 hours, the coverslips were fixed using methanol and stained with Giemsa and drug activity was evaluated by microscopically counting the number of infected and uninfected cells per 100 macrophages comparing with the control (Fig 5.3.) (324). The anti-leishmanial activity of compounds was expressed as percentage reduction in infected macrophages compared to untreated control wells.

**Figure 5.3.** Schematic overview of evaluation of the anti-leishmanial activity in static and flow culture systems.
5.3. Results

As previously mentioned, O’Keeffe A et al (2018) have described the Quasi Vivo QV900 macro-perfusion system and briefly, found that a $85 \pm 3\%$ infection rate of macrophages at 72 hours in static cultures decreased to $62 \pm 5\%$ for cultures under slow medium flow and $55 \pm 3\%$ under fast medium flow and media perfusion also decreased amastigote replication and both macrophage phagocytosis (by $44 \pm 4\%$ under slow flow and $57 \pm 5\%$ under fast flow compared with the static condition) and macropinocytosis (by $40 \pm 4\%$ under slow flow and $62 \pm 5\%$ under fast flow compared with the static condition). Mathematical and computational modelling were used to estimate the effect of speed of medium flow on infection rate, shear stress and oxygen concentration. For further details see publication Annex 1.

5.3.1. Macrophage functions

5.3.1.1. Phagocytosis.

Phagocytosis of latex beads by uninfected and infected macrophages (PEMs, BMMs or THP-1) showed a clear time dependent response (Fig 5.4.), with phagocytosis increasing with duration of incubation. Phagocytosis was significantly higher in infected cells (infection rate of $> 80\%$) compared to uninfected ones after 24 hours under static conditions ($p<0.05$ by t-test) (Table 5.2 and Fig 5.4). PEMs and BMMs showed significantly higher phagocytosis of latex beads than THP-1 ($p<0.05$ by one-way ANOVA).
Table 5.2. Phagocytosis of fluorescent latex beads (2 μm) by uninfected and infected PEMs, BMMs and THP-1 in static culture system.

<table>
<thead>
<tr>
<th>Time/Hour</th>
<th>Uninfected cells - static system</th>
<th>Infected cells - static system</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PEMs</td>
<td>BMMs</td>
</tr>
<tr>
<td>0.5</td>
<td>2.42 ± 0.2</td>
<td>2.3 ± 0.2</td>
</tr>
<tr>
<td>1</td>
<td>6.93 ± 0.8</td>
<td>6.2 ± 0.8</td>
</tr>
<tr>
<td>2</td>
<td>61.18 ± 1.5</td>
<td>60 ± 1</td>
</tr>
<tr>
<td>4</td>
<td>106.74 ± 7.7</td>
<td>95 ± 5</td>
</tr>
<tr>
<td>24</td>
<td>421.27 ± 30</td>
<td>396 ± 27</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data not shown). Phagocytosis was significantly higher (p<0.05 by t-test) in infected macrophages compared to uninfected ones. Phagocytosis was significantly higher (p<0.05 by t-test) in infected macrophages compared to uninfected ones. Initial macrophage infection rate was >80% after 24 h.
Figure 5.4. Phagocytosis of fluorescent latex beads (2 μm) by uninfected and infected PEMs (A), BMMs (B) and THP-1 (C) in static culture system. There is a significant increase in phagocytosis by infected macrophages compared to uninfected ones (p<0.05 by t-test). The data show means ± standard deviations (SD), N = 3. Infection rate was > 80%.

After which, the effects of media perfusion systems on phagocytosis function of *L. major*-infected macrophages were evaluated. Flow conditions caused a significant reduction in phagocytosis by infected macrophages as shown in Fig 5.5 - after 24 h of incubation, phagocytosis had significantly decreased from 530± 30 x 10^5, 519± 30 x 10^5 and 398± 22 x 10^5 beads/mg protein by PEMs, BMMs and THP-1, respectively in static cultures to 304± 32 x 10^5, 299.9± 24 x 10^5 and 200± 30 x 10^5 beads/mg protein by PEMs, BMMs and THP-1, respectively at slow flow speed (1.45 x 10^{-9} m/s) and this phagocytosis decreased more at faster flow speed (1.23 x 10^{-7} m/s) to 231± 28 x 10^5, 227.6± 25 x 10^5 and 144± 18 x 10^5 beads/mg protein by PEMs, BMMs and THP-1, respectively (p<0.05 by one-way ANOVA) (Table 5.3).
Table 5.3. Phagocytosis of fluorescent latex beads (2 μm) by infected PEMs, BMMs and THP-1 in the three culture systems (static, slow flow rate 1.45 x 10^{-9} m/s and fast flow rate 1.23 x 10^{-7} m/s).

Number of latex beads ± SD *10^5/mg protein

<table>
<thead>
<tr>
<th>Time/Hour</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>infected cells - static system</td>
<td>Infected cells - 1.45 x 10^{-9} m/s</td>
<td>Infected cells - 1.23 x 10^{-7} m/s</td>
<td>Infected cells - static system</td>
<td>Infected cells - 1.45 x 10^{-9} m/s</td>
<td>Infected cells - 1.23 x 10^{-7} m/s</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.5</td>
<td>3.45 ± 0.04</td>
<td>3 ± 0.04</td>
<td>1.8 ± 0.04</td>
<td>1.06 ± 0.02</td>
<td>1 ± 0.02</td>
<td>1 ± 0.02</td>
<td>0.54 ± 0.1</td>
<td>0.45 ± 0.1</td>
<td>0.3 ± 0.1</td>
</tr>
<tr>
<td>1</td>
<td>11.56 ± 0.02</td>
<td>10.9 ± 0.02</td>
<td>8 ± 0.02</td>
<td>6.59 ± 0.1</td>
<td>5.9 ± 0.1</td>
<td>3 ± 0.1</td>
<td>3.92 ± 0.06</td>
<td>3.89 ± 0.06</td>
<td>1.5 ± 0.06</td>
</tr>
<tr>
<td>2</td>
<td>76.58 ± 0.4</td>
<td>74 ± 0.3</td>
<td>59 ± 0.2</td>
<td>40.24 ± 0.4</td>
<td>39 ± 0.25</td>
<td>22 ± 0.25</td>
<td>28.18 ± 0.2</td>
<td>27 ± 0.2</td>
<td>15 ± 0.2</td>
</tr>
<tr>
<td>4</td>
<td>142.96 ± 3.9</td>
<td>139 ± 3</td>
<td>90 ± 2</td>
<td>75.92 ± 5.5</td>
<td>73.9 ± 5</td>
<td>49 ± 1</td>
<td>53.55 ± 4.9</td>
<td>50 ± 4</td>
<td>33 ± 3</td>
</tr>
<tr>
<td>24</td>
<td>530.05 ± 32.9</td>
<td>519 ± 30</td>
<td>398 ± 22</td>
<td>303.88 ± 27.5</td>
<td>299.9 ± 24</td>
<td>200 ± 30</td>
<td>231.11 ± 30</td>
<td>227.6 ± 25</td>
<td>144 ± 18</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data not shown). Flow conditions caused a significant reduction in phagocytosis by infected macrophages (p>0.05 by one-way ANOVA). Initial macrophage infection rate was >80% after 24 h.
Figure 5.5. Phagocytosis of fluorescent latex beads (2 μm) by infected PEMs (A), BMMs (B) and THP-1 (C) in the three culture systems (static, slow flow rate $1.45 \times 10^{-9}$ m/s and fast flow rate $1.23 \times 10^{-7}$ m/s). Phagocytosis is significantly higher in static than in flow system ($p<0.05$ by one-way ANOVA). The data are means ± standard deviations (SD), $N = 3$. Infection rate > 80%.

5.3.1.2. Macropinocytosis

Macropinocytosis of pHrodo Red dextran by uninfected and infected macrophages (PEMs, BMMs or THP-1) showed a clear time dependent response with macropinocytosis increasing with duration of incubation (Fig 5.6.). Macropinocytosis was significantly increased in infected macrophages,
from 19.02 ± 1.1, 16.5± 1.1 and 8±1.1 μg/mg protein of pHrodo Red dextran by uninfected PEMs, BMMs and THP-1, respectively to 25.3 ± 0.9, 23±0.8 and 13.5±0.8 μg/mg protein of pHrodo Red dextran in infected PEMs, BMMs and THP-1, respectively after 24h in static conditions (p<0.05 by t-test) (Table 5.4).
Table 5.4. Macropinocytosis of pHrodo™ Red dextran by uninfected and infected PEMs, BMMs and THP-1 in static culture system.

<table>
<thead>
<tr>
<th>Time/Hour</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>0.43 ± 0.01</td>
<td>0.3 ± 0.01</td>
<td>0.15 ± 0.01</td>
<td>0.92 ± 0.1</td>
<td>0.6 ± 0.1</td>
<td>0.3 ± 0.1</td>
</tr>
<tr>
<td>1</td>
<td>1.28 ± 0.3</td>
<td>1.1 ± 0.3</td>
<td>0.55 ± 0.3</td>
<td>2.8 ± 0.2</td>
<td>2.2 ± 0.1</td>
<td>1.6 ± 0.1</td>
</tr>
<tr>
<td>2</td>
<td>2.77 ± 0.5</td>
<td>2.5 ± 0.5</td>
<td>0.99 ± 0.5</td>
<td>3.78 ± 0.5</td>
<td>3.4 ± 0.3</td>
<td>1.8 ± 0.3</td>
</tr>
<tr>
<td>4</td>
<td>4.83 ± 0.9</td>
<td>4.1 ± 0.9</td>
<td>2.5 ± 0.9</td>
<td>7.1 ± 0.8</td>
<td>5.9 ± 0.7</td>
<td>3.9 ± 0.7</td>
</tr>
<tr>
<td>24</td>
<td>19.02 ± 1.1</td>
<td>16.5 ± 1.1</td>
<td>8 ± 1.1</td>
<td>25.3 ± 0.9</td>
<td>23 ± 0.8</td>
<td>13.5 ± 0.8</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data not shown). Macropinocytosis was significantly higher \( p<0.05 \) by t-test in infected macrophages compared to uninfected ones. Macropinocytosis was significantly higher \( p<0.05 \) by t-test in infected macrophages compared to uninfected ones. Initial macrophage infection rate was >80% after 24 h.
Figure 5.6. Macropinocytosis of pHrodo Red dextran by uninfected and infected PEMs (A), BMMs (B) and THP-1 (C) in static culture system. There is a significant increase in macropinocytosis by infected PEMs compared to uninfected ones ($p<0.05$ by t-test). The data are means ± standard deviations (SD), N = 3. Infection rate was > 80%.

After which, the effects of media perfusion systems on macropinocytosis function of $L.\ major$-infected macrophages were evaluated. Macropinocytosis was significantly reduced under flow conditions (Fig 5.7.), with higher speed of culture medium flow causing the greatest reduction, as after 24 hours of incubation with pHrodo Red dextran, macropinocytosis was reduced from 25.3 ± 0.9, 23± 0.8 and 13.5± 0.8 μg of pHrodo Red dextran /mg protein by PEMs, BMMs and THP-1, respectively under static to 15.1 ± 1, 14.99± 0.3 and 9± 0.3 μg/mg protein by PEMs, BMMs and THP-1, respectively under low flow ($1.45 \times 10^{-9}$ m/s) and more reduction occurred by higher flow ($1.23 \times 10^{-7}$ m/s) to
9.54 ± 1.2, 9± 1 and 5.5± 1 by PEMs, BMMs and THP-1 μg/mg protein, respectively (p<0.05 by one-way ANOVA) (Table 5.5).
Table 5.5. Macropinocytosis of pHrodo™ Red dextran by infected PEMs, BMMs and THP-1 at the three culture systems (static, slow flow rate $1.45 \times 10^{-9}$ m/s and fast flow rate $1.23 \times 10^{-7}$ m/s).

<table>
<thead>
<tr>
<th>Time/Hour</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
<th>PEMs</th>
<th>BMMs</th>
<th>THP-1</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>0.92 ± 0.1</td>
<td>0.6 ± 0.1</td>
<td>0.3 ± 0.1</td>
<td>0.29 ± 0.01</td>
<td>0.2 ± 0.01</td>
<td>0 ± 0.01</td>
<td>0 ± 0.0</td>
<td>0 ± 0.02</td>
<td>0 ± 0.02</td>
</tr>
<tr>
<td>1</td>
<td>2.8 ± 0.2</td>
<td>2.2 ± 0.1</td>
<td>1.6 ± 0.1</td>
<td>0.68 ± 0.5</td>
<td>0.55 ± 0.5</td>
<td>0.25 ± 0.5</td>
<td>0.13 ± 0.06</td>
<td>0.1 ± 0.05</td>
<td>0 ± 0.05</td>
</tr>
<tr>
<td>2</td>
<td>3.78 ± 0.5</td>
<td>3.4 ± 0.3</td>
<td>1.8 ± 0.3</td>
<td>1.75 ± 0.5</td>
<td>1.5 ± 0.5</td>
<td>0.7 ± 0.5</td>
<td>1.32 ± 0.2</td>
<td>1.1 ± 0.1</td>
<td>0.35 ± 0.1</td>
</tr>
<tr>
<td>4</td>
<td>7.1 ± 0.8</td>
<td>5.9 ± 0.7</td>
<td>3.9 ± 0.7</td>
<td>3.17 ± 0.9</td>
<td>3 ± 0.7</td>
<td>1.5 ± 0.7</td>
<td>2.29 ± 0.7</td>
<td>2 ± 0.55</td>
<td>0.9 ± 0.5</td>
</tr>
<tr>
<td>24</td>
<td>25.3 ± 0.9</td>
<td>23 ± 0.8</td>
<td>13.5 ± 0.8</td>
<td>15.1 ± 1.0</td>
<td>14.9 ± 0.3</td>
<td>9 ± 0.3</td>
<td>9.54 ± 1.2</td>
<td>9 ± 1</td>
<td>5.5 ± 1</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data not shown). Flow conditions caused a significant reduction in macropinocytosis by infected macrophages ($p>0.05$ by one-way ANOVA). Initial macrophage infection rate was >80% after 24 h.
Figure 5.7. Macropinocytosis of pHrodo Red dextran by infected PEMs (A), BMMs (B) and THP-1 (C) at the three culture systems (static, slow flow rate $1.45 \times 10^{-9}$ m/s and fast flow rate $1.23 \times 10^{-7}$ m/s). Macropinocytosis is significantly higher in static than in flow systems ($p < 0.05$ by one-way ANOVA). The data are means ± standard deviations (SD), N = 3. Infection rate was $> 80$.

5.3.2. Effects of media perfusion system on the anti-leishmanial activity of chitosan formulations

Dose-dependent anti-leishmanial activity (Fig 5.8.) was observed for all formulations (chitosan solution, blank chitosan-TPP nanoparticles and AmB loaded chitosan-TPP nanoparticles) across two media velocities and static culture. In the 72 h assays, the data showed that the addition of media perfusion reduced the anti-leishmanial activity of these three chitosan formulations. Chitosan solution, blank chitosan-TPP nanoparticles and AmB loaded chitosan-TPP nanoparticles showed a significantly higher activity in static culture (flow of 0 m/s) than in the QV900 system both at the base of the chamber (flow of $1.45 \times 10^{-9}$ m/s) and on an insert (flow of $1.23 \times 10^{-7}$ m/s).
The three formulations, chitosan solution, blank chitosan-TPP nanoparticles and AmB loaded chitosan-TPP nanoparticles, were 2.08 times, 2 times and 4 times respectively, more active against intracellular *L. major* amastigotes in static culture in comparison with the flow of $1.45 \times 10^{-9}$ m/s. Similarly, increasing the velocity of culture media from flow of $1.45 \times 10^{-9}$ m/s to flow of $1.23 \times 10^{-7}$ m/s by using the insert reduced the activity of chitosan solution, blank chitosan-TPP nanoparticles and AmB loaded chitosan-TPP nanoparticles against *L. major* amastigotes by 2.4 times, 1.8 times and 2.75 times respectively (Table 5.6). Regarding pure AmB, we did not find a significant difference in EC50 values between the three culture systems ($p>0.05$ by an extra sum-of-squares F test). In contrast, a significant difference was observed in EC90 values of pure AmB as increasing the media perfusion decreased the effectivity of AmB against 90% of amastigotes ($p<0.05$ by an extra sum-of-squares F test) (Table 5.6).

**Table 5.6.** *In vitro* activity of chitosan solution and nanoparticles against *L. major* amastigotes in RPMI medium (pH=6.5) at different flow rates

<table>
<thead>
<tr>
<th>Compound</th>
<th>Static- 0 m/s</th>
<th>Flow - $1.45 \times 10^{-9}$ m/s</th>
<th>Flow - $1.23 \times 10^{-7}$ m/s</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EC50</td>
<td>EC90</td>
<td>EC50</td>
</tr>
<tr>
<td>Chitosan solution</td>
<td>10.9</td>
<td>± 1</td>
<td>165</td>
</tr>
<tr>
<td>Blank chitosan-TPP nanoparticles</td>
<td>14.6</td>
<td>± 4</td>
<td>241</td>
</tr>
<tr>
<td>AmB loaded chitosan-TPP nanoparticles</td>
<td>0.1</td>
<td>± 0.01</td>
<td>1</td>
</tr>
<tr>
<td>AmB solution (Pure)</td>
<td>0.09</td>
<td>± 0.01</td>
<td>0.5</td>
</tr>
</tbody>
</table>

Experiments were conducted in triplicate cultures, data expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data not shown). *Statistically significant differences were found for the EC50 values of chitosan solution, blank chitosan-TPP nanoparticles and AmB loaded chitosan-TPP nanoparticles at static culture (flow of 0 m/s), flow of $1.45 \times 10^{-9}$ m/s and flow of $1.23 \times 10^{-7}$ m/s ($p<0.05$ by an extra sum-of-squares F test). Initial macrophage infection rate was >80% after 24 h.
Figure 5.8. Dose-response curve of the activity of chitosan solution (A), blank chitosan-TPP nanoparticles (B), AmB loaded chitosan-TPP nanoparticles (C) and AmB solution (pure) (D) against L. major amastigotes infecting PEMs in pH=6.5 under different flow conditions. Quasi Vivo QV900 system has been used as a flow culture system. Values are expressed as % amastigotes inhibition relative to untreated controls. Data are representative of one experiment in in triplicate cultures, data is expressed as mean +/- SD (experiment was reproduced further two times with confirmed similar data and data not shown).
5.4. Discussion

The QV900 culture flow system was used to overcome some fundamental limitations of *in vitro* static culture system when investigating cellular responses and anti-leishmanial activity of compounds and formulations. Static culture systems are unable to provide dynamic chemical or physical stimuli to cells, such as concentration gradients, flow, pressure, or mechanical stress caused by movement of fluids around them, which are physiologically relevant (100).

This study found a significant increase in cell functions (phagocytosis and macropinocytosis) in *L. major*-infected macrophages (PEMs, BMMs and THP-1) compared to uninfected cells - consistent with results described elsewhere, for example macrophages infected with either *L. donovani* or *L. mexicana* had greater pinocytic rates than uninfected macrophages, as measured by a fluorescent probe (fluorescein isothiocyanate dextran) (204). Similar observations have been reported with RAW 264.7 macrophages infected with *L. major* showing increased uptake of fluorescently labelled liposomes (204). This might be due to morphological changes of the infected cells or the parasitic infection may alter both the metabolic activity of the macrophages and their ability to ingest particulate material (346).

This study found that PEMs and BMMs showed significantly higher phagocytosis and macropinocytosis than THP-1, and this could be explained as BMMs and PEMs are more homogenous than THP-1, and they are characterised with their homogeneity and long lifespan (230).

We evaluated the effects of media perfusion rates on host cell phagocytosis and macropinocytosis. We found that phagocytosis and macropinocytosis were significantly decreased by media flow and increasing the media flow speed caused a further reduction in the uptake. This is consistent with previous reports of decreased uptake of fluorescein isothiocyanate (FITC)-poly (ethylene glycol) diacrylate particles (200 nm diameter) by human umbilical vein endothelial cells in a dynamic cell culture system exposed to shear stress.
of 10 dynes/cm² compared to the uptake in static cultures (347). Similar findings were also seen with a lower cellular uptake of solid silica particles (350 nm) by RAW 264.7 macrophages under dynamic condition compared to the uptake in static cultures (348). One explanation given was that the static system conditions might cause a sedimentation of the latex beads on the cell surface or exposure to higher concentrations of pHrodo Red dextran resulting in a local increase in their concentrations (349). In contrast, medium flow prevents such localization of materials with subsequently reduced uptake (350).

We also showed that the media perfusion system had a significant influence on the anti-leishmanial activity of chitosan solution, blank chitosan-TPP nanoparticles and AmB loaded chitosan-TPP nanoparticles- increasing the flow rates caused a significant decrease in their activity. Similarly, O'Keeffe reported that the anti-leishmanial activity of miltefosine and paromomycin against *L. major* amastigotes was reduced under these two flow rates (high and slow) (324).

This decrease in the anti-leishmanial activity of chitosan formulations under flow system could be attributed to a number of factors: (i) in a static system, waste products (because of catabolic and xenobiotic metabolism) accumulate in the culture medium and can cause an oxidative stress and lead to the loss of cellular function and viability during the culture time *in vitro*. On the other hand, culture under dynamic conditions can overcome these issues by the distribution of nutrients, waste products, and tested substances within the cell culture (351, 352, 353). (ii) It has been reported that static system conditions can cause a sedimentation of the drug on the cell surface resulting in a local increase in the drug concentrations (Fig 5.9). However, a flow method for the exposure of cells to the drugs can overcome this problem and leads to homogenous dispersion of the drugs and prevention of sedimentation (351, 352, 353).
(iii) The effects of the two media perfusion conditions used in our study on the accumulation of anti-leishmanial drugs (amphotericin B and miltefosine) have been previously reported by O'Keeffe et al (2017) - the accumulation of both drugs was significantly higher in the static system compared to the media perfusion system (Fig 5.10), after 24 hours and this could be due to a reduction in the rate of drug uptake (324).

The study described here also showed that cell uptake (phagocytosis and micropinocytosis) is reduced significantly by the application of flow compared with static culture conditions. Therefore, this reduction in drug accumulation and macrophage functions (phagocytosis and micropinocytosis) are contributing factor to the reduced anti-leishmanial activity seen (Fig 5.5, 5.7 and 5.8)
Figure 5.10. Accumulation of amphotericin B (left) and miltefosine (right) in peritoneal macrophages at three culture systems using the QV900 over time. Static and two flow rates \((1.33 \times 10^{-9})\) at the base of the chamber or \(1.17 \times 10^{-7}\) (m/s) on an insert) (324).

Broussou et al (2019) reported in vitro time-kill studies for a combination of amikacin and vancomycin against Staphylococcus aureus in static conditions and dynamic conditions (fluctuating antibiotic concentrations, by using A Hollow-Fibre model (Fig 5.11)) and reported a significant difference in the efficacy of the combination between static and dynamic conditions (354).

Both EC\(_{50}\) and EC\(_{90}\) values of AmB loaded chitosan-TPP nanoparticles against intracellular amastigotes significantly increased as the speed of media
perfusion increases and this pattern was not obvious when comparing EC\textsubscript{50} values of pure AmB however, the EC\textsubscript{90} values diminished with increasing flow rate. Similar finding was reported by O’Keeffe \textit{et al} (2017) in terms of EC\textsubscript{50} and EC\textsubscript{90} values of AmB solution at these three culture systems, could be due to the high activity of AmB against \textit{Leishmania} amastigotes (324).

The difference between AmB nanoparticles and AmB solution (EC\textsubscript{90} values, at three culture systems) could be due to that nanoparticles are uptaked at higher rates compared with solution and therefore a significant less amount of AmB nanoparticles entered the macrophages under flow system compared with AmB solution (353).

In conclusion, in the media perfusion culture system, flow speed was observed to influence the anti-leishmanial activities of the tested formulations. This could influence the development of new drugs for cutaneous leishmaniasis particularly by considering the possible higher flow rates in inflammatory sites.

The collateral effects of flow on pathogen replication rate and on host cell metabolism, as indicated by reduction in phagocytosis and macropinocytosis, introduces new avenues of research and how these models could be used in studies on immune response and drug and vaccine discovery. This combined experimental and modelling approach permits future hypothesis testing and development of more complex/advanced/predictive models for drug discovery and development.
6. General discussion

6.1. Discussion and conclusion

Although cutaneous leishmaniasis (CL) is not fatal, it does have a significant impact on the health and well-being impact of those infected. The large numbers involved, in at least 149 tropical and sub-tropical countries, have a detrimental impact on the economy of low- and middle-income countries where this disease is found (356, 357).

The available therapies for CL have acknowledged limitations which include adverse side effects/toxicity, are poorly tolerated, variable effectiveness against Leishmania species and are expensive in terms of both cost of drugs and care and other associated costs. Despite the clear need, new treatments for CL have not been forthcoming (51, 52). Drug discovery is a long and costly process which can take 10 to 20 years from a molecule to a usable drug, with an associated investment of a possible 2.6 billion USD$ before a new active compound is identified, developed for clinical applications and brought to the market (358, 359). CL could be regarded as one of the more neglected of the NTDs, typified by a general lack of interest in pursuing and funding drug development, both by Pharma and other actors, for this disease. Some antileishmanial drugs developed for VL that are in the current pipeline may be considered for the treatment of CL in the future (360).

One of the strategies to address the barriers of high cost and long developmental time-lines is the employment of drug delivery systems with an already known effective drug with established clinical activity. Drug delivery systems give an opportunity to manage the solubility and other pharmacokinetic parameters of a drug, such as bioavailability, half-life and biodistribution, and can serve to protect a drug from degradation. All this can result in both reducing toxicity and enhancing efficacy.

Amphotericin B (AmB), a polyene antibiotic, is considered the second most common treatment for leishmaniasis and is very effective against different Leishmania species experimentally, but its clinical use is limited due to its
inherent acute toxicity. AmB is one of the most-studied drugs for the development of new drug delivery strategies in the field of leishmaniasis (118, 361).

One of these promising drug delivery systems is AmBisome® (a liposomal formulation of AmB) which is effective against VL and CL. AmBisome® is less toxic than free AmB and the clinically used amphotericin B deoxycholate formulations (Fungizone®) and is recommended by the World Health Organization for the treatment of VL (60). AmBisome® has some limitations which include (i) the high cost (200 $ per vial of 50 mg, and is donated free for VL in endemic countries, not for CL), (ii) is the need for a cold chain (unstable over 25°C) and (iii) some renal toxicity and infusion-related reactions. In a recent study of CL and MCL in travellers coming back from both Old- and New-World countries AmBisome® treatment showed only 63% positive outcome and 53% of them experienced renal toxicity and infusion-related reactions (59) and (iv) higher rates of relapse have been observed in immunocompetent patients with VL treated with AmBisome® (60, 61).

There is an urgent need for new treatments which can eliminate the parasites, improve the healing process, are safe, reliable and also field-adaptable for use in diverse healthcare systems.

Chitosan has shown promising features in effective therapeutic delivery systems due to its cationic structure, biocompatibility, biodegradability, controlled drug release, mucoadhesive, wound healing and antimicrobial properties. Both chitosan in solution and nanoparticles showed interesting antimicrobial and antileishmanial activity with variable effective values across different published studies. These properties make chitosan an appropriate candidate for further studies to evaluate its suitability for the treatment of CL.

In Chapter 2, pH was demonstrated to play a critical role in the anti-leishmanial activity of chitosan and its derivatives (except carboxymethyl chitosan which showed no activity at both pH values), as all showed a higher anti-leishmanial effectivity at a lower pH. To date, there is no literature available on the anti-leishmanial activity of all of these derivatives or on the role of pH on the anti-leishmanial activity of chitosan. In this chapter, HMW chitosan demonstrated
a higher anti-leishmanial activity against *L. major* and *L. mexicana* promastigotes and amastigotes than other types and derivatives of chitosan. Accordingly, HMW chitosan was chosen for further studies. After which, the aim was to investigate whether the anti-leishmanial efficacy of HMW chitosan is related to indirect activity (through the activation of macrophages M1 pro-inflammatory phenotype) or via a direct way (through direct uptake of chitosan into the parasitophorous vacuole (PV) where the *Leishmania* amastigotes reside). Interestingly, it was shown that HMW chitosan acted by direct effect on the intracellular amastigotes; this has not been reported previously in any other literatures.

The results pointed towards the possibility of using HMW chitosan as a drug delivery component for CL treatment, harnessing the benefits of both anti-leishmanial activity of chitosan itself and to improve the therapeutic window of AmB (enhancing AmB anti-leishmanial activity and reducing its toxicity). AmB encapsulated in different types of chitosan nanoparticles has shown a promising *in vitro* and *in vivo* anti-leishmanial activity, see Table 4.3. Most of these studies used positively charged nanoparticles with a size greater than 100 nm. Therefore, in Chapter 3, we endeavoured to prepare two types of AmB-loaded chitosan nanoparticles; a positively charged type with TPP and a negatively charged type with dextran sulphate with the smallest possible sizes. The goal was to obtain the smallest sizes in an attempt to improve the topical delivery of AmB into the dermal layer of the skin. On the other hand, smaller nanoparticles when administrated intravenously, show a higher permeation through body membranes compared to larger nanoparticles, and smaller size of nanoparticles facilitates a passive transport from blood vessels to tissues (255).

The nanoparticle preparation parameters were optimised and two types of spherical blank and AmB loaded nanoparticles using the inotropic gelation method were successfully produced. One type of chitosan nanoparticles with a positive charge by using TPP as a crosslinker and this resulted in blank chitosan-TPP nanoparticles (size= 67 ± 7 nm, zeta potential= 28.5 ±1.9 mv) and AmB loaded chitosan-TPP nanoparticles ( size= 69 ± 8 nm, zeta potential= 25.5 ± 1 mv). The other type with a negative charge by using dextran sulphate
as a crosslinker and this resulted in blank chitosan-dextran sulphate nanoparticles (size= 170 ± 9 nm, zeta potential= -12.9 ± 3 mv) and AmB loaded chitosan-dextran sulphate nanoparticles (size= 174 ± 8 nm, zeta potential= -11 ± 1mv). Also, the importance of using cryoprotectants and the advantage of sucrose over D-mannitol in protecting the nanoparticles were identified during the freeze drying process. Then, the encapsulation efficacy and AmB loading were approximately 90% and 25%, respectively of both types of nanoparticles. In addition, these nanoparticles showed a high stability in terms of size and charge, in different conditions (different media (water, PBS, RPMI and mouse plasma) and at different temperatures (4, 34 or 37 °C)). Both types of nanoparticles displayed a slow release of AmB in PBS or mouse plasma. All previous promising properties of our nanoparticles made them suitable candidates for further studies in terms of evaluating the anti-leishmanial efficacy of blank chitosan nanoparticles or AmB loaded chitosan nanoparticles (as delivery vehicles) and the possibility of using them in CL mouse model either topically or intravenously.

The fourth chapter investigated the anti-leishmanial activity of chitosan formulations in vitro and in vivo. Firstly, both types of blank nanoparticles showed neither a significant haemolytic activity against human RBCs nor cytotoxicity against KB-cells. With regard to AmB loaded chitosan nanoparticles, both produced around 18-fold less haemolytic activity and 6-fold less toxicity against KB cells than pure AmB. Blank, positively surfaced charged, nanoparticles showed an in vitro activity against L. major and L. mexicana promastigotes and amastigotes at two pH’s of 7.5 and 6.5, with a higher activity at the lower pH. Encouragingly, AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran sulphate nanoparticles presented a similar anti-leishmanial activity to pure AmB against L. major and L. mexicana promastigotes and amastigotes, and a higher activity than AmBisome®. The little in vitro cytotoxicity and high effectivity against in vitro Leishmania parasites led to the evaluation of the anti-leishmanial activity of chitosan formulations in vivo L. major model of CL via the intravenous route of administration. A safe dosing regimen was established in BALB/c mice of AmB loaded chitosan-TPP nanoparticles and AmB loaded chitosan-dextran
sulphate nanoparticles via i.v. route - 5 mg/kg (AmB equivalent) and 10 mg/kg (AmB equivalent), respectively. Promisingly, AmB loaded chitosan-TPP nanoparticles (5 mg of AmB/kg/QAD for 10 days, i.v.) showed a higher in vivo anti-leishmanial effectivity than AmBisome® (10 mg of AmB/kg/QAD for 10 days, i.v) and was similar to the activity of paromomycin used as the positive control (50 mg/kg/QD for 10 consecutive days; i.p.) in terms of reducing lesion size and bioluminescence signal (parasite load). This anti-leishmanial activity of AmB loaded chitosan-TPP nanoparticles was in a dose-response manner. Levels of AmB within the infected lesion (rump skin) and control skin (uninfected skin, back skin) were assessed at the end of the experiment and a good correlation between the doses of AmB loaded chitosan-TPP nanoparticles and the intrallesional AmB and the relative reduction in parasite load and lesion size was found. Additionally, AmB loaded chitosan-TPP nanoparticles resulted in higher drug accumulation in the lesions in comparison with a higher dose of AmBisome®. Parasite load was determined via in vivo imaging (by using bioluminescent L. major strain) and compared with untreated controls. Previous studies have strongly correlated parasite load determined by both quantitative PCR and bioluminescent signal (199). qPCR determination of parasite load will be determined on the harvested and stored tissues from this study – this work fell beyond the time line of this project. To conclude, AmB loaded chitosan-TPP nanoparticles were more stable than AmBisome® and had a more sustainable drug release than AmBsiome (The release of AmB was 5% from AmB loaded chitosan-TPP nanoparticles and 75% from AmBisome® (362) in 24 h). Moreover, AmB loaded chitosan-TPP nanoparticles were significantly more active than AmBisome® against L. major in mice even though with lower doses of these nanoparticles than AmBisome®.

The possibility of using these nanoparticles as topical formulations was evaluated. The permeability of the nanoparticles (blank and AmB loaded nanoparticles) through uninfected and L. major infected mouse skin performing in vitro Franz cell diffusion studies was determined. Both types of nanoparticles acted as a drug delivery vehicle and released the AmB rather than permeating alongside the AmB molecules. For both types of
nanoparticles, AmB permeation was limited and slow, but interestingly higher in infected skin than uninfected, albeit in low concentrations (Kat ref). These outcomes in the permeation study indicate the poor suitability of these particular formulations as credible topical formulations to treat CL.

The effect of media perfusion on macrophage functions and on the anti-leishmanial activity of chitosan formulations was assessed in Chapter 5 in an attempt to simulate some of the more complex interactions between the parasite and macrophages in the mammalian host. For this purpose, a QV900 media perfusion system was used, as described by O'Keeffe et al (2017), with similar flow rates to mimic the interstitial tissue flow rate in the skin. Media perfusion significantly decreased both phagocytosis and macropinocytosis of different types of macrophages (PEMs, THP-1 and BMMs). This described how the additional complexity of each in vitro model could improve the predictivity of the assay and how drug properties based on static assays can give rise to misleading data. The aim of this perfusion model was to develop a more predictive in vitro model (compared to the current static 2D one), which could ultimately lead to a reduction in animal use and save both time and expenditure evaluating poor compounds. Interestingly, the anti-leishmanial activity of chitosan formulations was significantly less in the media perfusion systems compared to the static culture system.

6.2. Future work

AmB loaded chitosan-TPP nanoparticles were effective in the murine model (female BALB/c mice) of L. major, when administrated intravenously.

Many parts are associated with the scale-up of these nanoparticles from bench to the market. For instance, nature of material, procedure of nanoparticle development, cost, in vivo biodegradability of nanoparticles and acceptability of finished product both by clinicians and patients. On account of their economic feasibility, AmB loaded chitosan-TPP nanoparticles are better because they are made of chitosan and TPP whose production scale up is significantly less expensive than phospholipids in liposomal AmB.
The evaluation of anti-leishmanial activity of AmB loaded chitosan-TPP nanoparticles in vivo using a New World species (for example L. mexicana) might be of interest for future work. Assessment of the activity of these nanoparticles in other models of Leishmania infection, such as self-curing model would be interesting (363). Further extensive toxicity studies in animals would also be required.

The therapeutic index of these nanoparticles could be improved by either loading two active drugs into the nanoparticles, e.g. miltefosine (or other known active anti-leishmanials) and AmB, or by using a combination of therapy, e.g. using these nanoparticles via the i.v. route and other topical treatment (including thermotherapy or cryotherapy or paromomycin ointment) or other commercially available drugs. Both of these ways could develop more effective, lower-dose, and shorter treatments. It would be interesting to evaluate the in vitro and in vivo efficacy of AmB loaded chitosan-TPP nanoparticles in the treatment of VL. Another important experiment would be evaluating the distribution of the nanoparticles among different organs and study their uptake by lymphocytes, APCs and neutrophils.
7. References

13. CDC. Parasites - Leishmaniasis.


58. WHO. 2016. WHO and Gilead Sciences extend collaboration against visceral leishmaniasis.


73. CDC. 2018. Diagnosis and treatment of leishmaniasis: clinical practice guidelines by the Infectious Diseases Society of America (IDSA) and the American Society of Tropical Medicine and Hygiene (ASTMH). Centers for Disease Control and Prevention,


121. Ribeiro TG, Franca JR, Fuscaldi LL, Santos ML, Duarte MC, Lage PS, Martins VT, Costa LE, Fernandes SOA, Cardoso VN, Castilho RO, Soto M, Tavares CAP, Faraco AAG, Coelho EAF, Chávez-Fumagalli MA. 2014. An optimized nanoparticle delivery system based on chitosan and chondroitin sulfate molecules reduces the toxicity of
amphotericin B and is effective in treating tegumentary leishmaniasis. International journal of nanomedicine 9:5341-5353.


166. Van Bocxlaer K, Murdan S, Croft S L., Yardley V. 2015. CUTANEOUS LEISHMANIASIS -SKIN BARRIER PROPERTIES AND DRUG DELIVERY STRATEGIES-. PhD. School of Pharmacy.


176. FDA. 20002. GRAS Notices.


205. Miguel DC, Yokoyama-Yasunaka JK, Andreoli WK, Mortara RA, Uliana SR. 2007. Tamoxifen is effective against Leishmania and induces a rapid alkalinization of
parasitophorous vacuoles harbouring Leishmania (Leishmania) amazonensis amastigotes. J Antimicrob Chemother 60:526-34.


234. WHO. 2014. National Strategic Guideline on Kala-azar Elimination Program in Nepal


239. FDA. 2018. FOOD AND DRUGS.


303. Aguiar MG, Pereira AMM, Fernandes AP, Ferreira LAM. 2010. Reductions in Skin and Systemic Parasite Burdens as a Combined Effect of Topical Paromomycin and Oral Miltefosine Treatment of Mice Experimentally Infected with "Leishmania".


O’Keeffe A, Murdan S, Croft SL. 2017. DEVELOPMENT OF NOVEL PREDICTIVE 2D AND 3D IN VITRO MODELS FOR ANTI-LEISHMANIAL DRUG TESTING. PhD. London School of Hygiene and Tropical Medicine.


8. Appendix

8.1. Appendix 1: Validation of HPLC methods

Table 8.1. HPLC validation parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy</td>
<td>100.15 ± 0.22</td>
</tr>
<tr>
<td>Slope</td>
<td>108.11</td>
</tr>
<tr>
<td>Intercept</td>
<td>0.31</td>
</tr>
<tr>
<td>Linearity range</td>
<td>0.5-300 µg/ml</td>
</tr>
<tr>
<td>SE of intercept</td>
<td>0.2</td>
</tr>
<tr>
<td>SD of intercept</td>
<td>0.52</td>
</tr>
<tr>
<td>LOD (limit of detection) =3.3*(SD of intercept/Slope)</td>
<td>0.015 µg/ml</td>
</tr>
<tr>
<td>LOQ (limit of quantification) =10*(SD of intercept/Slope)</td>
<td>0.048 µg/ml</td>
</tr>
</tbody>
</table>

- Precision

Accuracy can be defined as the degree to which a measured value conforms to the true value. In pharmaceutical analysis, an assay is said to be accurate if the mean result is the same as the true value. On the other hand, precision is described as the variability of a set of measurements. Unlike accuracy, this does not provide any indication of the closeness of the obtained results from the true value. High precision is indicative of low variability in measurements usually demonstrated by low standard deviation values. This is usually reported as a percentage relative standard deviation.

(%RSD): SD/Drug*100

The precision of the method was determined by repeatability (intra-day) and intermediate precision (inter-day). Repeatability was determined by performing three repeated analysis of the same standard solution on the same day, under the same experimental conditions. The intermediate precision of the HPLC methods was assessed by carrying out the analysis on three different days (inter-day). For each drug, the percentage relative standard
deviation (%RSD) and the percentage recovery of the standard solutions are reported for each drug.

**Table 8.2. The precision of AmB HPLC assay**

<table>
<thead>
<tr>
<th>Standard concentration µg/ml</th>
<th>Intra-day calculated concentration (µg/ml)</th>
<th>Inter-day Calculated concentration (µg/ml)</th>
<th>Intra-day % RSD</th>
<th>Inter-day % RSD</th>
</tr>
</thead>
<tbody>
<tr>
<td>300</td>
<td>300.03± 0.21</td>
<td>300.33± 0.15</td>
<td>0.05</td>
<td>0.07</td>
</tr>
<tr>
<td>100</td>
<td>99.91± 0.16</td>
<td>100.10± 0.17</td>
<td>0.17</td>
<td>0.16</td>
</tr>
<tr>
<td>33.3</td>
<td>33.26± 0.05</td>
<td>33.29± 0.04</td>
<td>0.11</td>
<td>0.14</td>
</tr>
<tr>
<td>11.1</td>
<td>11.10± 0.09</td>
<td>11.27± 0.14</td>
<td>1.26</td>
<td>0.81</td>
</tr>
<tr>
<td>3.7</td>
<td>3.71± 0.02</td>
<td>3.70± 0.05</td>
<td>1.28</td>
<td>0.41</td>
</tr>
<tr>
<td>1.23</td>
<td>1.24± 0.02</td>
<td>1.23± 0.02</td>
<td>1.24</td>
<td>1.24</td>
</tr>
<tr>
<td>0.4</td>
<td>0.39± 0.02</td>
<td>0.4± 0.02</td>
<td>5.25</td>
<td>3.88</td>
</tr>
</tbody>
</table>

Figure Calibration curve of amphotericin B

\[
y = 96.6337x - 4.8347 \\
R^2 = 0.9972
\]
Development of an *in vitro* media perfusion model of *Leishmania major* macrophage infection

Alec O’Keeffe1,2, Lauren Hyndman1, Sean McGinity3, Alan Rieck1, Sudaxshina Murdan2, Simon L. Crofts2

1 Department of Infection and Immunology, London School of Hygiene and Tropical Medicine, London, United Kingdom, 2 Department of Pharmaceutics, UCL School of Pharmacy, University College London, London, United Kingdom, 3 Division of Biomedical Engineering, University of Glasgow, Glasgow, United Kingdom

* simon.crofts@ucl.ac.uk

Abstract

Background

In *in vitro* assays are widely used in studies on pathogen infectivity, immune responses, drug and vaccine discovery. However, most *in vitro* assays display significant differences to the *in vivo* situation and limited predictive properties. We applied medium perfusion methods to mimic interstitial fluid flow to establish a novel infection model of *Leishmania* parasites.

Methods

*Leishmania* major infection of mouse peritoneal macrophages was studied with the Quasisi Vivo QU9000 macro-perfusion system. Under a constant flow of culture media at a rate of 360μl/min, *L. major* infected macrophages were cultured either at the base of a perfusion chamber or raised on 9mm high inserts. Mathematical and computational modellling was conducted to estimate medium flow speed, shear stress and oxygen concentration. The effects of medium flow on infection rate, intracellular amastigote division, macrophage phagocytosis and macrophagocytosis were measured.

Results

Mean fluid speeds at the macrophage cell surface were estimated to be 1.45 x 10⁻⁵ m/s and 1.23 x 10⁻⁵ m/s for cells at the base of the chamber and cells on an insert, respectively. *L. major* macrophage infection was significantly reduced under both media perfusion conditions compared to cells maintained under static conditions; a 85±5% infection rate of macrophages at 72 hours in static cultures compared to 62±5% for cultures under slow medium flow and 55±3% under fast medium flow. Media perfusion also decreased amastigote replication and both macrophage phagocytosis (by 44±4% under slow flow and 57±5% under fast flow compared with the static condition) and macrophagocytosis (by 40±4% under slow flow and 52±5% under fast flow compared with the static condition) as measured by uptake of latex beads and pHrodo Red dextran.

PLOS ONE | [https://doi.org/10.1371/journal.pone.0219889](https://doi.org/10.1371/journal.pone.0219889) | July 24, 2019
Conclusions

Perfusion of culture medium in an in vitro L. major macrophage infection model (simulating in vivo lymphatic flow) reduced the infection rate of macrophages, the replication of the intracellular parasite, macrophage phagocytosis and macrocytophagocytosis with greater reductions achieved under faster flow speeds.

Introduction

Traditional cell culture methods typically rely on either immortalized cell lines or primary isolated cells grown in designed nutritious media on non-physiological substrates, such as functionalized plastic and glass. Although these methods have been at the core of in vitro studies on many basic biological processes, they provide a limited platform owing to both their inadequate representation of key physiological characteristics and their relevance to disease models [1]. One area that is often overlooked in cell culture models is the transport and movement of nutrients around cells, which occurs through fluid flow in the body. This could impact on the growth and survival of pathogens in intracellular models as infection is reliant on nutrients provided by the host cell and cell-cell interactions. Within the mammalian body, rates of fluid flow vary from the rapid plasma flow of 9.8 ml/min in the portal vein of the rat [2] to the slower 0.19 pl/min rate of interstitial fluid drainage from rat brains [3]. Interstitial fluid in tissues, including skin, arises from the normal leakage of plasma from blood vessels and has a composition that is similar to that of blood plasma [4]. It is estimated that up to 20% of the body’s mass is made up of interstitial fluid [4].

Leishmania is an infectious disease caused by protozoan parasites of the genus Leishmania, which have two distinct life cycle stages: an extracellular malleus promastigote form in the sandfly vector and an intracellular amastigote form that survives and multiplies in the phagocytes of mammal macrophages [5]. Two predominant forms of the disease result from infection by Leishmania parasites, the potentially fatal visceral leishmaniasis (VL) and the self-curing, cutaneous leishmaniasis (CL). Although macrophages of the liver and spleen infected with Leishmania donovani, the cause of VL, are exposed to plasma flow rates, in the skin sites of infection in CL, infected macrophages are exposed to interstitial fluid. While the exact speed of interstitial fluid flow through the CL lesion is not known, measurements have shown that interstitial fluid flow in uninfected human skin is of the order of 0.1–2 μm/s [6, 7]. Most in vitro studies on invasion, infection, immunology and drug discovery within the Leishmania field have so far been performed using macrophages grown under static culture conditions [8–10]. To simulate some of the more complex interactions between the parasite and macrophages in the host we selected the Quasi Vivo 900 media perfusion system (QV900) with a 6-channel optical tray to enable the imaging of cultures in situ at a flow rate similar to that of interstitial fluid. Here we describe the effect of media perfusion on the infection of mouse peritoneal macrophages with Leishmania major and use mathematical modelling to estimate the flow speed, shear stress and oxygen tension at the host cell surface. In addition, we have determined the impact of flow on intracellular amastigote division, and host cell phagocytosis and macrocytophagocytosis.

Material and Methods

Media perfusion system

Quasi Vivo media perfusion systems (Kirkstall Ltd, Rotherham, UK) were selected as they enable the direct observation of infected cells exposed to different medium perfusion rates and
the continuous monitoring of infection. The Quasi Vivo systems include the QV900, an individual chamber system, and the QV900, a six chamber optical tray which permits connecting of chambers in series. We selected the QV900 given that it is more suited to high throughput testing. Mathematical and computational modelling of the QV500 [11] has shown that the speed of media at the surface of cells cultured at the base of the chambers is within the range of interstitial fluid flow rates (μL/cm²) in humans for a flow rate of 30μL/min. However, the QV900 chambers differ in geometry and in particular are significantly deeper, having a depth of 22 mm compared with 12 mm in the QV500. As a result of this difference in chamber geometry, the fluid environment in the QV900 is markedly different from that in the QV500 at the same flow rate. Therefore, we inserted a 3D printed block composed of Nylon 12 (Sinterit Ltd) in selected chambers to enable us to study cells cultured at different depths in the QV900 chamber. Mathematical and computational modelling (see sections below) were utilized to calculate the insert height that would ensure the cell surface flow speeds would fall within the reported range for interstitial flow in the skin. All six chambers of the QV900 were connected in series with the last three of the chambers containing inserts. A peristaltic pump (Parker Hannifin, UK), external to the CO₂ incubator, continuously circulated culture media through the system.

Modelling fluid flow and oxygen transport in the QV900 system

COMSOL Multiphysics, a commercially available finite element analysis software, was used to perform simulations in this study. Initial modelling focused on single chamber studies to establish the size of the insert required to achieve the desired cell surface flow speeds. Subsequently, simulations were conducted for six chambers connected in series, matching the experiments. Fig 1 illustrates the computational domains for cells placed at the base of the chamber (left) and on a 9mm insert (right). Note that in both cases the chambers are identical in dimensions, but since there is assumed to be no fluid flow beneath the insert, the depth of this computational domain is reduced.

The fluid flow was modelled using the Navier-Stokes equations, assuming that the media is an incompressible Newtonian fluid. The transport of oxygen throughout the media was modelled by convection and diffusion. The cells were assumed to reside at the base of each computational domain on circular coverslips of diameter 12mm. Oxygen consumption was described using Michaelis-Menten kinetics and implemented through a flux boundary condition. The equations and parameter values used in the simulations are detailed in the supplementary material S1 File.

Culture systems

Leishmania parasites. L. major (MHOM/SA/85/JSH118) amastigotes were obtained and isolated from mouse skin lesions. They were allowed to transform to promastigotes and were maintained in Schneider’s insect medium (Sigma Aldrich, UK) supplemented with 10% heat inactivated foetal calf serum (HIFCS) (Harlan, UK) at 26°C. The parasites were routinely passed through BALB/c mice (Charles River, UK) and low passage number promastigotes (< passage number 3) were used for experiments as infectivity has been shown to decrease with time of parasite cultivation [12]. All animal experiments were conducted under licence (project license 706997 or X20014454) in accordance with UK Home Office approval, EU regulations, EU directive 2010/63/EU. Protocols followed in these studies for the isolation of peritoneal macrophages was approved by the ISHMT Animal Welfare and Ethics Review Board. The mice are housed in green line I.V.Cs. 5 mice per cage, with access to food and water ad libitum. At all stages the 3Rs (replacement, reduction and refinement) were taken into consideration.
Macrophages. Mouse peritoneal macrophages (PEM) were isolated from CD-1 mice (Charles River, Margate, UK) by abdominal lavage [1,3] with RPMI-1640 medium containing 1% penicillin and 1% streptomycin (Sigma, UK).

THP1 cells (ATCC TIB-202, UK) were maintained in RPMI-1640 containing 10% FBS (Harlan, UK) and passaged by a 1:10 split weekly.

Infection of macrophages by L. major promastigotes. Macrophages were plated on 12mm round glass coverslips (Bellco, US) placed in 24 well plates (Corning, UK) at a density of 4 x 10^5 cells per well in RPMI-1640 media supplemented with 10% FBS. The plates were incubated at 37°C in 5% CO2 for 24 hours. L. major stationary phase promastigotes were counted and diluted of different concentrations of parasite (2 x 10^7 to 6 x 10^7) were pre-prepared in media to give initial parasite:macrophage ratios within the range of 0.5:1-1:10:1. Promastigotes were added to the macrophage cultures. The plates were placed in an incubator maintained at 34°C (temperature relevant for CL, [14]) at 95% CO2 for 24 hours. Subsequently, two-thirds of the glass coverslips were transferred to the media perfusion system and maintained under flow conditions at a flow speed of 360 ul/min for 72 hours in a 34°C, 5% CO2 incubator. The remaining coverslips were used as the static control, with macrophages maintained in the same culture medium without flow. The cells were euthanized (Sigma, UK) fixed and stained with Giemsa’s stain (Sigma, UK). The infection rate of the macrophages was assessed visually using an oil immersion microscope (100x magnification Zeiss, UK) by counting the number of infected cells per 100 macrophages. Values for percentage infection throughout are shown as mean ± standard deviation.

Incorporation of 5-ethyl-2'-deoxyuridine (EdU) into dividing amastigotes. Invitrogen Click-iT EdU Imaging Kit (Invitrogen, UK) was used to measure 5-ethyl-2'-deoxyuridine incorporation as a measure of proliferation. Only the dividing parasites should incorporate the EdU as the macrophage populations used are fully differentiated non dividing cells. The kit comprised of a fluorescently labelled DNA base, which is incorporated into DNA synthesized during amastigote division. Experiments, based on the methodology of Taggarini et al. [9], were conducted as before except that PEMs were infected with a ratio of 3 L. major promastigotes:1 macrophage and maintained at 34°C, 5% CO2 in an incubator for 24 hours.

Media used contained 56 μM EdU. After 24 hours, cells were placed in a new 24 well plate and were fixed in 4% Paraformaldehyde (PFA) (Sigma, UK) for 15 minutes at room temperature. The samples were treated with 0.2% Triton X-100 (Sigma, UK) in PBS (Sigma, UK) for 20 minutes and then 1% BSA (Sigma, UK) in PBS for 10 minutes. Click-iT reaction cocktail was prepared according to instructions in Invitrogen Click-iT EdU Imaging Kit. Click-iT reaction
cocktail (0.5 mL) was added to each well containing a coverslip, and plates were incubated for 30 minutes at room temperature, protected from light. Cells were then washed with 1 mL of 3% BSA in PBS, then incubated with 300 mM DAPI stain (Sigma, UK) for 10 minutes to stain the nuclei of the cell. Coverslips were mounted onto slides and imaged using a confocal microscope (Zeiss LSM510 Axiovert, Germany). The lasers used were Laser Diode 405 nm for DAPI excitation and Argon laser: 458, 488, 514 nm for FITC excitation. Images were captured at 40x magnification and analysed using Velocity software (PerkinElmer, USA) to automatically count the total number of nuclei in each field of view and this is proportional to the total cell number. A minimum of 100 macrophages were counted microscopically from each coverslip. Images were manually viewed to count the number of fluorescent and non-fluorescent parasites within each cell. The results were exported and analysed with Graphpad Prism.

Measurement of macrophage functions. Phagocytosis by macrophages was initially evaluated using 0.5, 1, and 2 μm diameter fluorescent red labelled latex beads (carboxylate-modified polystyrene) (Sigma-Aldrich, UK) [15,16]. 2 μm beads were eventually selected as they showed maximal signal. Macrophages were infected with parasites, then transferred to the three flow conditions as described above. To each well, 2 μm beads (9.12 x 10^5 latex beads/mL) were added and the cells were incubated for 0.5, 1, 2, 4, and 24 hours at 34 °C under the three different flow conditions. The experiment was terminated by washing the cells 4 times with ice-cold PBS pH 7.4 to remove non internalized latex beads, followed by the addition of 1 mL of 0.5% Triton X-100 in 0.2 M NaOH to lyse the cells. Phagocytosis was quantified by the analysis of the cell lysate using a fluorescence plate reader (Spectramax M3, at excitation and emission wavelengths set at 575 and 610 nm), calibrated with standard solutions containing different amount of latex beads in a cell lysate mixture. Uptake was expressed as the number of latex beads associated per mg of cellular protein, the protein content of the cell lysate being measured using a Micro BCA protein kit (Thermo Fisher, UK) assay as per supplier’s instructions. For control studies, 1 μg/ml cycloheximide D was used as a phagocytosis inhibitor (Sigma-Aldrich, UK) by incubation with macrophages for 2 hours prior to addition of the latex beads. Phagocytosis was completely inhibited after 0.5, 1, 2, and 4 hours of incubation with cycloheximide D and 90% after 24 hours.

Macropinocytosis. Macropinocytosis was measured using a fluorescence-labeled dextran dye [17] (pHrodo Red dextran, average molecular weight of dextran 10,000 MW, Thermofisher, UK). This dye has a pH-sensitive fluorescence emission that increases in intensity with increasing acidity while exhibiting minimal fluorescence at neutral pH. Macrophages were infected with parasites and then transferred to the three flow conditions as described above. Macrophages were washed 3 x by Live Cell Imaging Solution (Thermofisher, UK) and the cells were returned to RPMI 1640 + 10% FCS containing 40 μg/mL pHrodo Red dextran (1 μl for each well) and incubated at 34 °C / 5% CO2 for 0.5, 1, 2, 4 and 24 hours under the three different flow conditions. At each time point, the cells were washed with Live Cell Imaging Solution and macropinocytosis was analysed by a Spectramax M3 at excitation and emission wavelengths set at 560 and 585 nm respectively. Chlorpromazine hydrochloride 10 μg/mL, a known inhibitor (Sigma-Aldrich, UK), was used as a control and was incubated with macrophages for 2 hours prior to addition of fluorescence-labeled dextran dye. Macropinocytosis was completely inhibited after 0.5, 1, 2, and 4 hours of incubation with chlorpromazine hydrochloride and by 90% after 24 hours.

Results
Establishment of infected macrophages in Quasi Vivo systems
Initial experiments using the Quasi Vivo systems involved the adaptation of the QV900 for our experimental purposes and the establishment of media perfusion within the system with a
focus on the optimization of conditions to maintain viable cells within the system. A second objective was to ensure that an infection with *Leishmania* parasites could be sustained, as shown in subsequent experiments.

**QV900 media perfusion system modelling**

Initially, single chamber simulations were carried out to estimate the height of the insert required to ensure the cell surface flow speed would be within the reported range for interstitial flow in the skin. Table 1 shows the estimated speed of the culture medium on the cell surface for various insert heights. It is clear that a 9mm insert would enable a culture medium flow speed in line with the speed of interstitial fluid flow in the skin, and therefore this height was chosen for subsequent modelling and experiments.

Subsequent mathematical and computational modelling was carried out to match the experimental set up, i.e. we simulated six chambers connected in series, with the first three chambers having cells residing at the base and the next three chambers having cells raised 9mm. Fig 2 illustrates results which are representative of the first three chambers in the series. All plots in Fig 3 show the results for chamber one, with the exception of the lower right plot which shows results for the first three connected chambers. The simulated oxygen concentration decreases from the inlet of the chamber, where oxygen is supplied, to the base of the chamber, where oxygen is consumed by the cells (Fig 2, upper left). At the base of the chamber, the oxygen concentration is highest at the inlet side, reducing towards the centre, before rising again at the outlet side of the chamber (Fig 2, middle left). This gradient, clearly highlighted in the lower left plot of Fig 2, is a combined result of the complex flow field and the fact that oxygen consumption only occurs on the part of the base where the cells reside. Similar results for chambers two and three are shown in the lower right plot of Fig 2. The oxygen concentrations are lower in each consecutive chamber as a result of consumption, but the pattern of oxygen concentrations across the base is consistent between each chamber. The overall gradient of oxygen at the base of the first three connected chambers ranges from a maximum of 0.2059 mol/m³ in chamber one to a minimum of 0.2029 mol/m³ in chamber three.

The upper right plot of Fig 2 illustrates the flow speed and streamlines (the trajectories that particles would follow), demonstrating how the media flows through the chamber. The media flow is fastest at the inlet and outlet, and flow recirculation zones are observed beneath the inlet and at the base of the chamber. In these areas, the media is recirculated which could result in parasites and oxygen/drug molecules being trapped. The flow speed of the media at the base of the first chamber is consistent with the second and third chambers and has a mean value of 1.45 x 10⁻⁹ m/s. We note that this is slightly higher than the mean flow speed obtained in a single chamber (Table 1) as a result of the altered fluid dynamics due to connecting the chambers in series. A 2D representation of the magnitude of the shear stress the cells are under at the

<table>
<thead>
<tr>
<th>Insert height (mm)</th>
<th>Mean cell surface flow speed (m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>9mm</td>
<td>1.33 x 10⁻⁹</td>
</tr>
<tr>
<td>7mm</td>
<td>2.69 x 10⁻⁹</td>
</tr>
<tr>
<td>5mm</td>
<td>3.03 x 10⁻⁹</td>
</tr>
<tr>
<td>3mm</td>
<td>5.01 x 10⁻⁹</td>
</tr>
<tr>
<td>1mm</td>
<td>7.77 x 10⁻⁹</td>
</tr>
<tr>
<td>0mm</td>
<td>1.17 x 10⁻⁹</td>
</tr>
</tbody>
</table>

https://doi.org/10.1371/journal.pone.0219985.t001
base of the chamber is shown in the middle right plot of Fig. 2. The shear stress values range from a minimum of $2.09 \times 10^{-9}$ Pa to a maximum of $1.06 \times 10^{-7}$ Pa which is consistent with the second and third chambers.

Fig. 3 illustrates results which are representative of the last three chambers in the series i.e. where the cells are placed on a 9mm insert. All plots show the results for chamber four (the first chamber in the series which has the cells raised by 9mm), with the exception of the lower right plot which shows results for the last three connected chambers (chambers 4, 5 and 6).

The inclusion of the 9mm insert has an impact on both the pattern and magnitude of the oxygen concentration and fluid flow. Higher oxygen concentrations are observed throughout the whole chamber when compared to the chambers without an insert (Fig. 3, upper left), and the minimum oxygen concentration at the base of the chamber occurs closer to the outlet side than when compared to the chambers without an insert (Fig. 3, middle). The oxygen concentration gradient across the base of the chamber is clearly highlighted in the lower left plot of Fig. 3. Again, this pattern is a combined result of the complex flow field and the fact that oxygen consumption only occurs on the part of the base where the cells reside. Similar results for chambers 5 and 6 are shown in the lower right plot of Fig. 3. As before, the oxygen concentration decreases between consecutive chambers due to consumption but the pattern remains
Fig. 3. Simulation results for cells on top of the 9mm insert. Upper left: Oxygen concentration in chamber 4. Upper right: Flow profile in chamber 4. Middle left: Oxygen concentration at the base of chamber 4. Middle right: Magnitude of the shear stress at the base of chamber 4. Lower left: Oxygen concentration across the center of the base of chamber 4. Lower right: Oxygen concentration across the center of the base of chambers 4, 5, and 6.

the same. The overall gradient of oxygen at the base of the last three connected chambers ranges from a maximum of 0.2093 mol/m³ in chamber 4 to a minimum of 0.2069 mol/m³ in chamber 6.

The depth of the last three chambers in the series is dramatically reduced due to the 9mm insert which has a large impact on the pattern of flow (Fig. 3, upper right). In this case, the only flow recirculation zone is observed beneath the inlet to the chamber. The mean flow speed of the media at the cells on top of the insert in the fourth chamber is 1.23 x 10⁻⁵ m/s—two orders of magnitude higher than in the chambers without the insert. This is consistent with the fifth and sixth chambers where the mean flow speed is also 1.23 x 10⁻⁵ m/s. Due to the difference in the flow profile, the pattern of shear stress at the base of the chamber is also noticeably different when compared to the chambers without an insert (Fig. 3, middle right). The shear stress values range from a minimum of 5.75 x 10⁻⁵ Pa at the edges of the base of the chamber to a maximum of 1.58 x 10⁻⁴ Pa at the center of the base of the chamber. This is again consistent with the fifth and sixth chambers.

Determination of optimal experimental conditions

Initially, we used both THP1 cells and PMCs at different concentrations from 1 x 10⁵ to 4 x 10⁵ cells per chamber, to establish a viable, reproducible and measurable system. After preliminary
work using THP-1 cells PEMs were selected for further studies as in this macrophage type infections with L. major were easier to establish and to sustain. Peritoneal macrophages at 4 x 10^5 cells per well were chosen as this concentration gave the most reproducible results following a series of studies at different conditions that were investigated (Fig. 5). The initial studies showed that:

1. the addition of parasites in the medium during perfusion, at parasite:macrophage ratios from 0.5:1 to 10:1, resulted in zero macrophage infection after 72 hours and was therefore not pursued.

2. a set number of parasites per ml of circulating media from 4 x 10^5 to 1.2 x 10^6 cells per ml caused the parasites to collect within the chambers resulting in over-infection and bursting of the macrophages at the 72 hour time point. This approach was also not pursued.

3. the addition of different numbers of promastigotes before the initiation of media perfusion at parasite:macrophage ratios from 0.5:1 to 25:1 in the medium for a 24 hour pre-infection
before media perfusion resulted in a controllable, reproducible infection after 72 hours. This approach was adopted.

4. There were decreased rates of macrophage infection with increasing flow rates from 50 to 360 to 1000 μl/min (Fig. 5). A flow rate of 360 μl/min was subsequently selected as it gave a more acceptable level of infection.

Infection of mouse peritoneal macrophages (PEMs) in the media perfusion system

The percentage of PEMs infected after 72 hours in each of the three flow conditions (i.e. static (0 μl/min), base of the chamber (1.45 x 10^-5 μl/min) and on the insert (1.23 x 10^-5 μl/min) using different parasite/macrophage ratios are shown in Fig. 5). The percentage infection after 24 hours before the transfer to the media perfusion system, at each of the starting infection ratios were reproducible across all of the infected cultures at that ratio. Mean initial percentage infection

---

Fig 5. Influence of flow rate on percentage infection at a L. major promastigote to macrophage ratio of 15:1. https://doi.org/10.1371/journal.pone.0219985.g005
Infection in PEMs under three conditions

Fig. 6. Box and whisker diagram showing the percentage of infected cells over a range of different infection ratios, parasite:Macrophage number, and different flow conditions. Significance tested using a two sided t-test p<0.01 * p<0.001 **** ns = not significant N = 6.

Parasite to Host Cell Ratio and Condition

levels ± 5D after 24 hours were 36 ± 1.51 ± 2.70 ± 4 and 87 ± 5% for the four different initial infection ratios of 0.5:1, 1:1, 3:1 and 6:1 parasite concentration to cell concentration. Media perfusion was maintained over the following 72 hours.

As the flow speed of the culture medium was increased from 0 m/s in the static condition to 1.45 x 10^-5 m/s to 1.23 x 10^-3 m/s (cells in the insert in chambers), the percentage infection of host cells decreased at all parasite to host ratios used (0.5:1, 1:1 and 3:1) (Fig. 6). However, the influence of medium flow speed on macrophage infection decreased as the parasite to host ratio increases, until at a parasite to host ratio of 6:1, increasing the flow speed of the culture medium had little effect on the percentage infection levels of the host. As expected increasing the initial parasite to host ratio increases the overall infection levels after 72 hours.

Comparison of data sets showed significant differences (at least p<0.01, by one-way ANOVA) from each other except when comparing the data at the 6:1 ratio (Fig. 6).

Incorporation of 5-ethynyl-2’-deoxyuridine (EdU)

The number of amastigotes per macrophage were counted microscopically after 24 hr under the three flow conditions, showing a similar parasite burden with approximately 2 amastigotes per infected cell (Fig. 7) across the infected cells that were imaged. Percentage infection rates were identical after the first 24 hr infection (63%) regardless of the speed of media perfusion per the cell would be maintained over the following 24 hr. The percentage of amastigotes that incorporated EdU into DNA was significantly lower in cultures maintained under perfusion conditions (Fig. 7), with a significant reduction observed (one way ANOVA, p<0.05) in cultures in flow systems compared to static cultures after 24 hr. On average, the mean percentage of amastigotes that incorporated EdU into DNA was 31 ± 7% in cells maintained in static
Fig 7. Left: Bar graph showing percentage of L. major amastigotes that incorporated the EdU marker into DNA at the three different conditions, static (0 min), low flow (1.45 x 10^{-3} m/s) and high flow (1.23 x 10^{-2} m/s). Right: Bar graph showing parasite burden in mouse peritoneal macrophages at the three conditions. *p<0.05 N=3.

Macrophage functions

Phagocytosis. Phagocytosis of latex beads by uninfected and infected PEMs showed a clear time dependent response (Fig 9) with phagocytosis increasing with duration of incubation. Phagocytosis was significantly higher (p<0.05 by t-test) in infected macrophages (infection rate of >80%) compared to uninfected ones (530± 30 x 10^3 versus 421± 30 x 10^2) beads/mg protein after 24 hours under static conditions.

Flow conditions caused a significant reduction in phagocytosis by infected macrophages as shown in Fig 9, such that after 24 h of incubation, phagocytosis had significantly decreased from 304± 30 x 10^2 beads/mg protein in static cultures to 304± 30 x 10^2 beads/mg protein at slow flow speed and 231± 30 x 10^2 beads/mg protein at fast flow speed (p<0.05 by one-way ANOVA).
Fig 8. Phagocytosis of fluorescent latex beads (2 μm) by uninfected and infected PEMs in static culture system. There is a significant increase in phagocytosis by infected PEMs compared to uninfected ones (p<0.05 by t-test). The data show mean ± standard deviations (SD), N = 3. Infection rate was > 80%.

https://doi.org/10.1371/journal.pone.021986-g008

Macropinocytosis

Macropinocytosis of pHrodo Red dextran by uninfected and infected PEMs showed a clear time dependent response (Fig 10) with macropinocytosis increasing with duration of incubation. Macropinocytosis was significantly increased in infected PEMs (p<0.05 by t-test) compared to uninfected ones (25±1.1 versus 19±1.0) μg/mg protein of pHrodo Red dextran after 24 hours of incubation.

Macropinocytosis was significantly reduced under flow conditions (Fig 11), with higher speed of culture medium flow causing greater reduction (p<0.05 by one way ANOVA) so that after 24 hours of incubation with pHrodo Red dextran, macropinocytosis was 25.3±1.1, 15.1±0.9 and 9.5±0.9 μg/mg protein under static, low flow and fast flow respectively.

Discussion

Media perfusion system and modelling

The importance of body fluid flow rates in physiology has been recognized for more than half a century [18]. Understanding the effects of fluid flow on solute transport in biological tissues and on cell-cell signalling and morphogenesis is now substantial. Media perfusion can provide more than just increased cell nourishment, it can also, for example, induce blood and lymphatic capillary morphogenesis in vitro [19–21], maintain the functional activity of
Fig 9. Phagocytosis of fluorescent latex beads (2 μm) by infected PEls in the three culture systems (static, low flow rate 1.45 x 10⁻⁸ m/s and fast flow rate 1.23 x 10⁻⁷ m/s). Phagocytosis is significantly higher in static than in flow system (p<0.05 by one-way ANOVA). The data are means ± standard deviations (SD), N = 3. Infection rate > 80%.

https://doi.org/10.1371/journal.pone.0219863.g009

...chondrocytes and osteocytes [22–25], drive fibroblast differentiation [26] and induce cytokine production by smooth muscle cells [27]. Static systems do not offer any form of dynamic chemical or physical stimulus to cells, such as concentration gradients, flow, pressure, or mechanical stress caused by movement of fluids around them. This is a major limitation in experiments investigating cellular responses in vitro since the complex interplay of mechanical and biochemical factors are absent. We used the QV900 system to introduce a fluid flow component to an in vitro L. major macrophage infection model. In addition, we adapted the QV900 system to enable comparison of the effect of different flow rates to static cultures on infection of macrophages. Experiments were performed with cells cultured at the base of the chamber (low flow) and cells cultured on top of a 3D printed insert (high flow). The 3D printed insert placed into the chambers enabled us to study media flow speed at the cell surface which is in line with values reported in the literature for interstitial flow in the skin [4,6,7].

Mathematical modeling also showed that cells cultured on the 9mm insert experienced flow speeds and shear stress that were two orders of magnitude higher than those affecting cells cultured at the base of the chamber. Oxygen concentrations at the base of the chambers with the insert were also determined to be higher when compared with the chambers without the insert.
The significance of our findings is that it is possible to expose cells to vastly different mechanical and chemical environments depending on where they are cultured in the chamber. This is consistent with previous studies where mathematical and computational models showed that changing the geometry of similar perfusion bioreactors has an impact on experimental conditions such as flow speed, shear stress and oxygen concentration [11].

Infection of macrophages in the media perfusion system

The macrophage infection level caused by parasite inocula over the three different conditions varied significantly. Macrophage infection by parasites was reduced by media flow, with significant reductions seen as the media flow speed increased, as shown in Fig 5. This pattern was also seen when using a larger range of initial infection ratios (See S2 File, where we used THP1 cells as the host cells). Possible reasons for the reduction in infection rate with increasing flow rate include: (a) reduced contact time between parasites and cells, (b) increase in the supply of nutrients to the host cells, (c) effect of higher shear stress on receptors, and (d) reduced proliferation of the parasite within the host cell. Promastigotes that are external to the cells but have remained on the glass coverslip may have reduced contact with the cells [28] after transfer to the perfusion system, as they will be pushed away from the cell by the media flow. Without sustained physical contact, the parasites will not be phagocytosed and will not establish an infection within the cell[25]. A lower probability of parasite invasion into macrophages could lead to
Fig 11. Macrophagocytosis of plasmodo Red dextran by infected PIMs at the three culture systems (static, slow flow rate $1.45 \times 10^{-9}$ m/s and fast flow rate $2.23 \times 10^{-7}$ m/s). Macrophagocytosis is significantly higher in static than in flow systems (p < 0.01 by one-way ANOVA). The data are mean ± standard deviations (SD), N = 3. Infection rate was > 80.

https://doi.org/10.1371/journal.pone.0219885.g011

...a lower infection rate, as fewer parasites would reach the phagolysosome, the site of parasite replication. The first step in the phagocytosis of the Leishmania promastigotes is binding to receptors on the cell surface: the Fc receptor (FcR), complement receptor type 3 (CR3), and mannose-fucose receptor have all been shown to be receptors for the parasite [29]. The flow of the media could cause a reduction in binding between the receptor and parasite. Another possible explanation for the effect of media perfusion on the final levels of infection is that the media flow provides more nutrients to the macrophages. Whilst we have considered only oxygen transport in our mathematical modelling, and demonstrated differential concentrations of oxygen at the cell surface with increasing media flow, it follows that the concentration of other important nutrients will be similarly affected. An increased supply of nutrients may provide more starting reagents for the production of anti-parasitic effector. In addition to this beneficial effect to the host cell, it is possible that the opposite occurs in the parasite, as the parasites could expend more energy [30] resisting the flow of the media reducing successful evasion of the macrophage cellular response and replication once they have been phagocytosed. Cells have been shown to respond to shear stress by changing shape [31], phenotype [32], and release of proteins/chemicals [33]. This stress will undoubtedly have an impact on the phagocytosis process [34].
Another possibility for the lower percentage infection in media perfusion system maintained cultures could be that the rate of parasite proliferation in the host cell is altered. The EdU incorporation assay demonstrated that fewer parasites are actively incorporating the labelled DNA base under media perfusion conditions. Although the average amastigote burden of the PEMs was the same under static, low flow and high flow, the lower EdU incorporation at higher flow speeds shows that parasites were replicating to a lower extent. This is a phenomenon also seen in plankton where increased flow reduces biomass build up [37].

Flow also affected phagocytosis and macroinocytosis of macrophages. Firstly, we established there was a significant increase in both cell functions in PEMs infected with L. major compared to uninected cells. These data are consistent with results described elsewhere, for example macrophages infected with either L. donovani or L. mexicana increased their phagocytic rates as measured by a fluorescent probe (fluorescein isothiocyanate dextran) [38]. Similar observations have been reported with RAW 264.7 macrophages infected with L. major showing increased uptake of fluorescently labelled liposomes [38]. This might be due to morphological changes of the infected cells or the parasitic infection may alter both the metabolic activity of the macrophages and their ability to ingest particulate material [39]. Our results demonstrated that phagocytosis and macroinocytosis were significantly decreased by media flow and that increasing the media flow speed caused a further reduction in the uptake. This is consistent with previous reports of decreased uptake of fluorescein isothiocyanate (FITC) -poly (ethylene glycol) diacylate particles (200 nm diameter) by human umbilical vein endothelial cells in a dynamic cell culture system exposed to shear stress of 10 dynes/cm² compared to the static culture [40]. Similar findings were also seen with a lower cellular uptake of solid silica particles (350 nm) by RAW 264.7 macrophages under dynamic condition compared with the static culture [41]. One explanation given was that the static system conditions might cause sedimentation of the beads on the cell surface or exposure to higher concentrations of pHfo/o Red dextran resulting in a local increase in their concentrations [42]. In contrast, medium flow prevents such localization of materials with subsequently reduced uptake [43].

In conclusion, in the media perfusion Leishmania-macrophage model flow speed was shown to affect infection rate even at interstitial fluid rates. This could have an impact on the development of Leishmania infection in skin especially when considering the possible higher flow rates in inflammatory sites. The role of mathematical modelling was essential to understanding different chemical and physical conditions resulting from the flow and, highlighting the need for mathematical modelling to be further integrated into this approach. The collateral effects of flow on pathogen replication rate and on host cell metabolism, as indicated by reduction in phagocytosis and macroinocytosis, further indicates research avenues and how these models might be used in studies on immune responses and drug and vaccine discovery. Additionally, our combined experimental and modelling approach has allowed us to generate hypotheses which we will test in future through the development of more advanced mathematical models and experiments.

Supporting information

S1 File. Supplementary material for “Development of a media perfusion model of macrophage infection by Leishmania major”.

S2 File. Supplementary material 2 for “Development of a media perfusion model of macrophage infection by Leishmania major”.

271
Acknowledgments

AO thanks Dr Vanessa Tardley for mentorship and guidance throughout the project and especially with the animal studies. SM and LH gratefully acknowledge a financial donation from Kirkstall Ltd this does not alter their adherence to PLOS ONE policies on sharing data and materials.

Author Contributions

Conceptualization: Alec O’Keeffe, Sean McGinty, Simon L. Croft.
Data curation: Lauren Hyndman.
Formal analysis: Alec O’Keeffe, Lauren Hyndman, Sean McGinty.
Funding acquisition: Sean McGinty, Simon L. Croft.
Methodology: Alec O’Keeffe, Lauren Hyndman, Sean McGinty, Alaiz Riezki.
Project administration: Simon L. Croft.
Software: Sean McGinty.
Validation: Alec O’Keeffe.
Visualization: Alec O’Keeffe.
Writing - original draft: Alec O’Keeffe, Lauren Hyndman, Sean McGinty, Alaiz Riezki.
Writing - review & editing: Sudarshana Murdan, Simon L. Croft.

References


8.3. Appendix 3: Paper 2

Copyright proof

Good morning, Alix,

Thank you for your message. This paper is being published under an open-access license (https://creativecommons.org/licenses/by/4.0/), so permission is not needed. Please see the link for the terms of the license (i.e., attribution).

Best,
Amanda

Amanda Donaldson
Assistant Production Editor
American Society for Microbiology
1730 N Street, NW
Washington, DC 20036
email: Amanda@asmusa.org

From: Alix Rock  
cSent: Wednesday, January 22, 2020 03:51 AM
To: Donaldson, Amanda
Subject: REQUEST TO USE COPYRIGHTED MATERIAL IN THESIS

Dear Amanda,

I am currently studying for a PhD at London School of Hygiene and Tropical Medicine. I am contacting you to ask permission to include the following paper within the electronic version of my PhD thesis:

Activity of chitosan and its derivatives against Listeria monocytogenes and L. monocytogenes in vitro

The thesis will be made available within LSE Library Online: http://eprints.lse.ac.uk/70738/our institutional repository. The repository is noncommercial and openly available to all.

Yours sincerely,

Alix
Results

In vitro activity of chloroquine and derivatives against L. major and L. donovani.

Antimonial activity against promastigotes and amastigotes of L. major and L. donovani was significantly higher compared to chloroquine.

The minimal inhibitory concentration (MIC) of chloroquine was 32 µM, while the MICs of L. major and L. donovani were 16 and 8 µM, respectively.

Derivatives of chloroquine showed increased activity against promastigotes and amastigotes of L. major and L. donovani.

The MICs of these derivatives were 16 and 8 µM, respectively, showing a significant increase in activity compared to chloroquine.

The results suggest that the increased activity of these derivatives may be due to structural modifications that enhance their affinity for the target parasite.

Conclusions

The in vitro activity of chloroquine and its derivatives against L. major and L. donovani is promising, with some derivatives showing MICs lower than 16 µM.

Further studies are needed to evaluate the in vivo efficacy of these derivatives and to determine their potential for clinical use in the treatment of leishmaniasis.

References

Cellular uptake of MIBI chromates and initiation of mitotic arrest

We found that the initial binding of MIBI chromates did not directly affect activity against intracellular amastigotes. Therefore, we investigated whether the anti-proliferative effects of MIBI chromates against intracellular amastigotes after 4 and 24 h were caused by the activity of the chromate against intra- cellular amastigotes at 4 h. To test whether chromates were effective against amastigotes at 4 h, we used a fumonisin B1-resistant clone of L. major. The chromates inhibited proliferation of the fumonisin B1-resistant clone (Fig. 7A). This result suggests that the chromates also inhibit proliferation of amastigotes at 4 h.

In a time-dependent manner, the chromates inhibited amastigotes by 4 h. This result suggests that the chromates are effective against intracellular amastigotes. However, the chromates did not affect the proliferation of amastigotes at 4 h.

In a time-dependent manner, the chromates inhibited amastigotes by 4 h. This result suggests that the chromates are effective against intracellular amastigotes. However, the chromates did not affect the proliferation of amastigotes at 4 h.

In a time-dependent manner, the chromates inhibited amastigotes by 4 h. This result suggests that the chromates are effective against intracellular amastigotes. However, the chromates did not affect the proliferation of amastigotes at 4 h.

In a time-dependent manner, the chromates inhibited amastigotes by 4 h. This result suggests that the chromates are effective against intracellular amastigotes. However, the chromates did not affect the proliferation of amastigotes at 4 h.

In a time-dependent manner, the chromates inhibited amastigotes by 4 h. This result suggests that the chromates are effective against intracellular amastigotes. However, the chromates did not affect the proliferation of amastigotes at 4 h.
The role of HMV on HIV-1 replication 

The role of HMV on HIV-1 replication was examined for the activation effects of the virus on the replication of HMV. 

1. **Virus and Cells** 
   - Virus: HIV-1 
   - Cells: Human lymphoblastoid cells 

2. **Methods** 
   - **Transduction Assay:** 
     - Cells were transduced with HIV-1 in the presence of HMV. 
     - Supernatants were collected at different time points. 

3. **Results** 
   - HMV significantly increased the number of HIV-1 copies per cell. 
   - The increase was dose-dependent. 

4. **Discussion** 
   - HMV enhances HIV-1 replication by activating specific cellular pathways. 
   - Further studies are needed to understand the molecular mechanisms involved. 

---

**References:** 
