Mini Review

Application of Nano Drugs in Treatment of Leishmaniasis

Abstract

Objective: Leishmaniasis is endemic in 88 countries with incidence rate of 1.5-2 million; the most common form of leishmaniasis is cutaneous leishmaniasis (CL) with 1.5 million new cases per year. Correct diagnosis and characterization of the particular parasite is important for evaluating prognosis and prescribing appropriate treatment. The current management of leishmaniasis is drug treatment of patients, to alleviate disease and vector control to reduce its transmission. Also, current treatments for visceral leishmaniasis are unsatisfactory because of their toxicity, resistance and high cost. The purpose of the present study was to review the application of nano drugs in treatment of leishmaniasis.

Materials and Methods: It was used as source of research the following databases: MEDLINE, Cochrane Library, Scielo, PubMed and the database of CAPES.

Results: It has been proved nanosilver particles with highly small sizes, have a good penetration into the cutaneous lesions and a good efficacy on leishmaniasis.

Introduction

Protozoan parasitic diseases stay an unsolved public health difficulty, especially in tropical regions [1]. The major death toll is due to malaria, leishmaniasis, and African and American trypanosomiasis, whose high mortality rates in developing countries are associated to poor sanitary conditions and lack of efficient prophylactic measures [1-4]. Despite the remarkable amount of research aimed at the production of protective vaccines, the success is still elusive measures [1-4]. Despite the remarkable amount of research aimed at the production of protective vaccines, the success is still elusive measures [1-4].

Leishmaniasis is an infection that is caused by an obligate intracellular protozoon of the genus Leishmania [6-8]. The natural transmission of Leishmania parasites is carried out by sandflies of the genus Phlebotomus or Lutzomyia [8]. These parasites cause three forms of leishmaniasis according to the localization of the parasites in mammalian tissues, notably visceral, cutaneous, and mucosal leishmaniasis [8].

Leishmaniasis is reported from 88 countries and estimated that 350 million world-wide are at risk of acquiring one form of the diseases, and 12 million are infected with annual occurrence rate of about 1.5 to 2 million [8-10]. According to findings 1-5 millions are affected annually in Iran. The real rate of incidence is 4 to 5 times higher than the reported prevalence [10-12].

The current management of leishmaniasis is drug treatment of patients, to alleviate disease and vector control to reduce its transmission [12-14]. Pentavalent antimonials (namely, sodium stibogluconate (SSG) and meglumine antimoniate) are the mainstay of anti-leishmanial therapy [14-17]. Although glucantime is commonly used for the treatment of leishmaniasis, it has some side effects including increased liver enzymes and electrocardiogram changes [17]. In addition, the drug is expensive, the injection is painful, and research shows that resistance of parasite to glucantime is growing in different parts of the world [17-20]. The use of nanotechnology is ever-expanding in today's technologically advanced world. This form of technology has many advantages over existing ones in many fields and medicine is one of them [20]. There are numerous devices and mechanisms developed with the aid of nanotechnology that can help cure diseases/disorders in a much better and more efficient manner [21]. The usefulness of nanotechnology can especially be seen in the treatment of infectious diseases [21,22]. This review has investigated the treatment of leishmaniasis by nano drugs.

Nano drugs against leishmaniasis

Nanotechnology must be applied for curing infectious disease like leishmaniasis [22]. Many drugs have been made and are still being discovered, but the protozoan re-emerges showing drug resistance to the effective medications [23]. Nanotechnology has just managed to show its promise in developing a liposomal formulation called amphotericin B for leishmaniasis but has serious side effects and is not cost effective in developing countries [23-25]. Initial, nanodelivery systems for delivering chemotherapeutics were nanodisks impregnated with amphotericin B, polymeric-nanoparticle loaded with pentamidine, primaquine, and niosomes, which still need validation at the clinical level [25-28]. There is an urgent need to take up assignments to use effective nanotechnology devices in combating this infectious disease [25-28].

Different mechanisms were proposed for antimicrobial property of nanoparticles [28]. One mechanism is binding of nanoparticle to sulfur/ phosphorus-containing biomolecules such as proteins and DNA, which leads to impairment of cell membrane, enzymes,
and DNA [28-31]. Some nanoparticles are deposited within mitochondria and impair oxidative stress pathway [28-31]. On the other hand, adenosine triphosphate (ATP) synthesis is inhibited when mitochondrial proteins are damaged [28-31]. Additionally, antimicrobial nanoparticles can impair glycoprotein and lipophosphoglycan molecules, which are responsible for infectivity of bacteria and parasites [31]. Another mechanism is ion release from nanoparticles [31]. These ions interact with cysteine-containing proteins, and inhibit protein functions [31-33].

Before penicillin, colloidal silver was a treatment of choice for many illnesses and infections. There are many reports that show colloidal silver is effective on about 650 different micro-organisms [27-29]. Therefore, it seemed Nanosilver particles with highly small sizes, have a good penetration into the cutaneous lesions and a good efficacy on Leishmania spp [27-33].

Silver has been used to treat different infections for many years; silver nano-particle form usage has opened new ways of treatment with development of nano-technology [1-3]. Nanosilver compounds show impact on a wide range of microorganisms including bacteria, viruses, fungi, and even protect against protozoa and influenza [1-3]. Nanosilver solution is a new drug with anti-bacterial, antifungal and antiviral properties [3]. Advantages of this drug compared to other drugs are efficacy at low concentration and long life duration [34-36]. Studies showed that nanosilver cement has high antibacterial activity and high effectiveness against multi resistant bacteria without cytotoxicity in vitro. Recently, findings have demonstrated that nanosilver has anti-inflammatory effects and increases wound healing and dressings of wounds [1,2,37]. If these results are confirmed in vivo, nanosilver may be appropriate for ulcer treatment [1]. Nanosilver DNA damage, denature proteins and enzymes and produce free radicals and some studies showed that nanosilver is cytotoxic to several different cell lines [1,38]. The teratogenicity of nanosilver in humans is indistinctive and is not reported in the literature but in vivo (animal studies) and in vitro studies showed nanosilver can exhibit a significant level of toxicity [39].

Nano-mediated drug delivery of plant-derived products is one of the most important strategies for the treatment of visceral leishmaniasis (VL) in the absence of adequate anti-leishmanial drugs. Artemisinin has been traditionally used for the treatment of malaria and has been reported to exhibit anti leishmanial and antitumor activities [2,38-40]. Earlier, it has been proved that formulation of artemisinin-loaded nanoparticles improved its activity against L. donovani amastigotes ex vivo [17].

Nanoparticles of amphotericin B have greater efficacy than conventional amphotericin B. This formulation may have a good safety profile, and if production costs are low, it may prove to be a feasible alternative to conventional amphotericin B in the treatment of VL [17,25-29].

If nanoparticles are systemically administered, they will be agglomerated after exposure to plasma, and their efficacy will be decreased [17]. Anti-leishmanial nanoparticles must be conjugated with biological compounds such as antibody or lectin, which bind to specific targets, in order to exert more toxicity for parasites and less toxicity for normal cells [21-32]. Some nanoparticles have high cytotoxicity on macrophages, which must be considered [17]. The use of nanoparticles for treatment of CL may have both positive and negative consequences. Some reports indicate that gold nanoparticles (Au NPs), titanium dioxide nanoparticles (TiO2 NPs), zinc oxide nanoparticles (ZnO NPs), magnesium oxide nanoparticles (MgO NPs), etc. have antibacterial properties [30-33].

On the other hand, some nanoparticles have photo thermal effect after exposure to near infra-red (NIR) light [31-33]. These nanoparticles absorb NIR energy, and alter it to heat [31]. Then, temperature is increased and cells will be damaged [31]. It is stated that leishmania parasites are sensitive to heat, and heat therapy has been used as a new way with nanoparticles [31-36].

Vaccine development in leishmaniasis is another major area where DNA nanotechnology may have an important contribution [26]. Till date, many efforts have been undertaken to develop a successful vaccine, but realizing them into the clinics has been a bottle neck [26].

Conclusions

Nanotechnology being applied to treat leishmaniasis is a relatively new area of research and a lot still needs to be done to set up merge the available datasets with new data that will arise when nanotechnology will be applied to various aspects of the leishmanial disease.

References

1. Mohelabi M, Rezayat MM, Giliani K, Sarkar S, Akhoundi B, et al. (2009) Nanosilver in the treatment of localized cutaneous leishmaniasis caused by Leishmania major (MRHO/IR/K5/ER): an in vitro and in vivo study. DARU 17: 285-289.
2. Nilloforouzhadeh MA, Shirani-Bidabadi LA, Zolfaghari-Baghbaderani A, Jafari R, Heidari-Beni M, et al. (2012) Topical effectiveness of different concentrations of nanosilver solution on Leishmania major lesions in Balb/c mice. J Vector Borne Dis 49: 249-253.
3. Rahi AA, Ali MA, Al-Charrakh AH (2013) Biosynthesis of silver nanoparticles by Leishmania tropica. Afr J Biotech 12: 6718-6722.
4. Mayeilit K, Taheri AR, Rajabi O, Sazgarnia A (2015) Ultraviolet B efficacy in improving antileishmanial effects of silver nanoparticles. Iran J Basic Med Sci 18: 677-683.
5. Kansal S, Tandon R, Verma A, Misra P, Choudhary AK, et al. (2014) Coating doxorubicin-loaded nanocapsules with alginate enhances therapeutic efficacy against Leishmania in hamsters by inducing Th1-type immune responses. Br J Pharmacol 171: 4038-4050.
6. Mario Zucca, Dianella Savoia (2011) Current Developments in the Therapy of Protozoan Infections. The Open Medicinal Chemistry Journal 5: 4-10.
7. Manandhar KD, Yadav TP, Prajapati VK, Kumar S, Rai M, et al. (2008) Antileishmanial activity of nano-amphotericin B deoxycholate. J Antimicrob Chemother 62: 376-380.
8. Collier MA, Peine KJ, Gautam S, Oghumu S, Varikuli S, et al. (2016) Host-mediated Leishmania donovani treatment using AR-12 encapsulated in acetlated dextran microcapsules. Int J Pharmacol 499: 186-194.
9. Mol M, Kosey D, Singh S (2015) Nano-Synthetic Devices in Leishmaniasis: A Bioinformatics Approach. Front Immunol 6: 323.
10. Carvalheiro M, Esteves MA, Santos-Mateus D, Lopes RM, Rodrigues MA, et al. (2015) Hemisynthetic triflurain analogues incorporated in liposomes for the treatment of leishmanial infections. Eur J Pharm Biopharm 93: 346-352.
11. Want MY, Islamuddin M, Chouhan G, Ozbak HA, Hemeq HA, et al. (2015) Therapeutic efficacy of artemisinin-loaded nanoparticles in experimental visceral leishmaniasis. Colloids Surf B Biointerfaces 130: 215-221.

12. Chaurasia M, Pawar VK, Jaiswal AK, Dude A, Pallikal SW, et al. (2015) Chondroitin nanocapsules enhanced doxorubicin induced apoptosis against leishmaniasis via TH1 immune response. Int J Biol Macromol 79: 27-36.

13. Ibrahim F, Thio TH, Faissal T, Neuman M (2015) The application of biomedical engineering techniques to the diagnosis and management of tropical diseases: a review. Sensors (Basel) 15: 6947-6995.

14. de Carvalho RF, Ribeiro IF, Miranda-Vilela AL, de Souza Filho J, Martins OP, et al. (2013) Leishmanicidal activity of amphotericin B encapsulated in PLGA-DMSA nanoparticles to treat cutaneous leishmaniasis in C57BL/6 mice. Exp Parasitol 135: 217-222.

15. Jebali A, Kazemi B (2013) Nano-based antileishmanial agents: a toxicological study on nanoparticles for future treatment of cutaneous leishmaniasis. Toxicol In Vitro 27: 1896-1904.

16. Zucca M, Savoia D (2011) Current developments in the therapy of protozoan infections. Open Med Chem J 5: 4-10.

17. Mohan S, Srivastava P, Maheshwari SN, Sundar S, Prakash R (2011) Nano-structured nickel oxide based DNA biosensor for detection of visceral leishmaniasis (Kalaj-azar). Analyst 136: 2845-2851.

18. Singodia D, Khare P, Dube A, Talegaonkar S, Khar RK, et al. (2011) Development and performance evaluation of alginate-capped amphotericin B lipid nanoconstrasts against visceral leishmaniasis. J Biomed Nanotechnol 7: 123-124.

19. Fattahi Baghi A, Shahcheraghi SH, Nematoitah S (2015) Comparison of hematological aspects: Visceral leishmaniasis and healthy children. Trop Parasitol 5: 133-135.

20. Ayatollahi J, Fattahi Baghi A, Shahcheraghi SH (2014) Rare variants of cutaneous leishmaniasis presenting as eczematous lesions. Med J Islam Repub Iran 28: 71.

21. Ayatollahi J, Ayatollahi A, Shahcheraghi SH (2013) Cutaneous leishmaniasis of the eyelid: a case report. Case Rep Infect Dis 214297.

22. DOLAT E, Rajabi O, Salarabadi SS, Yazegari-Dehkordi S, Sazgnaria A (2015) Silver nanoparticles and electroporation: Their combinational effect on Leishmania major. Bioelectromagnetics 36: 586-596.

23. Kalantari H, Hemmati AA, Bavarsad NS, Rezaie A, Amjadi S (2014) Effect of topical Nanoliposomes of Paromomycin on Rats Liver and Kidney. Jundishapur J Nat Pharm Prod 9: 173565.

24. Kumar R, Sahoo GC, Pandey K, Das V, Das P (2015) Study the effects of PLGA-PEG encapsulated amphotericin B nanoparticle drug delivery system against Leishmania donovani. Drug Deliv 22: 383-388.

25. Das S, Roy P, Mondal S, Bera T, Mukherjee A (2013) One pot synthesis of gold nanoparticles and application in chemotherapy of wild and resistant type visceral leishmaniasis. Colloids Surf B Biointerfaces 107: 27-34.

26. Badlee A, Heravi Shargh V, Khamesipour A, Jaafar MR (2013) Micro/nano particle adjuvants for antileishmanial vaccines: present and future trends. Vaccine 1: 735-749.

27. Santos DM, Carneiro MW, de Moura TR, Fukutani K, Clarencio J, et al. (2012) Towards development of novel immunization strategies against leishmaniasis using PLGA nanoparticles loaded with kinetoplastid membrane protein-11. Int J Nanomedicine 7: 2115-2127.

28. Van de Ven H, Vermeersch M, Matheusen A, Vandenoot J, Weyenberg W, et al. (2011) PLGA nanoparticles loaded with the antileishmanial saponin β-aescin: factor influence study and in vitro efficacy evaluation. Int J Pharm 420: 122-132.

29. Kunjachan S, Jose S, Thomas CA, Joseph E, Kiessling F, et al. (2012) Physicochemical and biological aspects of macrophage-mediated drug targeting in anti-microbial therapy. Fundam Clin Pharmacol 26: 63-71.

30. Danesh-Bahremini MA, Shokri J, Samiei A, Kamali-Sarvestani E, Barzegar-Jalali M, et al. (2011) Nanovaccine for leishmaniasis: preparation of chitosan nanoparticles containing Leishmania superoxide dismutase and evaluation of its immunogenicity in BALB/c mice. Int J Nanomedicine 6: 835-842.

31. Tafaghodi M, Eskandari M, Kharazizadeh M, Khamesipour A, Jaafari MR (2010) Immunization against leishmaniasis by PLGA nanospheres loaded with an experimental autoclaved Leishmania major (ALM) and Quillaja saponins. Trop Biomed 27: 639-650.

32. Tafaghodi M, Khamesipour A, Jaafari MR (2011) Immunization against leishmaniasis by PLGA nanospheres encapsulated with autoclaved Leishmania major (ALM) and CpG-ODN. Parasitol Res 108: 1265-1273.

33. Kota GF, de Sousa MR, Fereguetti TO, Saleme PS, Alvarisa TK, et al. (2016) The Cure Rate after Placebo or No Therapy in American Cutaneous Leishmaniasis: A Systematic Review and Meta-Analysis. PLoS One 11: 0149697.

34. Faliero RJ, Kumar R, Bunn PT, Singh N, Chauhan SB, et al. (2015) Combined Immune Therapy for the Treatment of Visceral Leishmaniasis. PLoS Negl Trop Dis 10: e0044145.

35. Wani GM, Ahmad SM, Khursheed B (2015) Clinical study of cutaneous leishmaniasis in the Kashmir Valley. Indian Dermatol Online J 6: 387-392.

36. Pineda-Reyes R, Llano-Cuentas A, Dancuart M (2015) Traditional treatments in an endemic area of american cutaneous Leishmaniasis in Peru. Rev Peru Med Exp Salud Publica 32: 761-765.

37. Shawler AJ, Boggild AK (2015) Cutaneous leishmaniasis in travellers: a focus on epidemiology and treatment in 2015. Curr Infect Dis Rep 17: 489.

38. Cardona-Arias JA, Vélez ID, López-Carvajal L (2015) Efficacy of thermotherapy to treat cutaneous leishmaniasis: a meta-analysis of controlled clinical trials. PLoS One 10: 0122569.

39. Yeşilova Y, Turan E, Sürüçi HA, Aksoy M, Özbilgin A (2015) Successful treatment of cutaneous leishmaniasis with amphotericin B, a case of unresponsive to pentavalent antimony therapy. Türkiye Parazitol Derg 39: 63-65.

40. Matin RN (2015) Cutaneous leishmaniasis--treatment options in children. Br J Dermatol 172: 844-845.