Vaccines for canine leishmaniasis

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Leishmaniasis is the third most important vector-borne disease worldwide. Visceral leishmaniasis (VL) is a severe and frequently lethal protozoan disease of increasing incidence and severity due to infected human and dog migration, new geographical distribution of the insect due to global warming, coinfection with immunosuppressive diseases, and poverty. The disease is an anthroponosis in India and Central Africa and a canid zoonosis (ZVL) in the Americas, the Middle East, Central Asia, China, and the Mediterranean. The ZVL epidemic has been controlled by one or more measures including the culling of infected dogs, treatment of human cases, and insecticidal treatment of homes and dogs. However, the use of vaccines is considered the most cost–effective control tool for human and canine disease. Since the severity of the disease is related to the generation of T-cell immunosuppression, effective vaccines should be capable of sustaining or enhancing the T-cell immunity. In this review we summarize the clinical and parasitological characteristics of ZVL with special focus on the cellular and humoral canine immune response and review state-of-the-art vaccine development against human and canine VL. Experimental vaccination against leishmaniasis has evolved from the practice of leishmanization with living parasites to vaccination with crude lysates, native parasite extracts to recombinant and DNA vaccination. Although more than 30 defined vaccines have been studied in laboratory models no human formulation has been licensed so far; however three second-generation canine vaccines have already been registered. As expected for a zoonotic disease, the recent preventive vaccination of dogs in Brazil has led to a reduction in the incidence of canine and human disease. The recent identification of several Leishmania proteins with T-cell epitopes anticipates development of a multiprotein vaccine that will be capable of protecting both humans and dogs against VL.

Keywords: vaccines, canine leishmaniasis, visceral leishmaniasis, zoonotic visceral leishmaniasis

Why is Leishmaniosis Important to Human and Veterinary Medicine?

Human leishmaniasis, caused by several species of Leishmania, comprises a group of diseases which are mostly zoonotic. These include visceral leishmaniasis (VL) and the cutaneous and mucocutaneous forms (CL). This group of infections is the third most important vector-borne disease after malaria and lymphatic filariasis (Solano-Gallego et al., 2009).

According to WHO, leishmaniasis is considered to be endemic in 88 countries, with more than 350 million people at risk. The estimated incidence is 2 million new cases per year, 0.5 million of VL, and 1.5 million of CL [World Heath Organization (WHO, 2011)]. Without prompt appropriate treatment as many as 95% of kala-azar patients die resulting in at least 50,000 deaths per year worldwide (Chappuis et al., 2007). Ninety percent of the human cases of VL are notified in Bangladesh, India, Nepal, Sudan, and Brazil, 95% of the cases of human CL are reported in Bolivia, Brazil, and Peru (mucocutaneous leishmaniasis), and 90% of cutaneous leishmaniasis cases occur in Afghanistan, Brazil, Iran, Peru, Saudi Arabia, and Syria (WHO, 2011). These statistics and epidemiological data however, are certainly an underestimation since they were first published by WHO in 1993 and have remained unaltered until now (WHO, 2011). Over the last two decades however many factors determined the increase in the incidence of human VL. Among them we can consider the geographical spread of VL in Europe, Africa, America, and Asia (Romero and Boelaert, 2010; Palatnik-de-Sousa and Day, 2011) caused by human migration carrying infected dogs to areas where the specific insect vector was already present (Maguill, 1995); the expansion of the phlebotomine habitats due global warming (Witt et al., 2009); the increase in dog cases with a travel history (Duprey et al., 2006); the urbanization of the disease after deforestation (Harhay et al., 2011); and the frequent coinfection in HIV-positive individuals (Evans and Kedzierski, 2012).

Visceral leishmaniasis is a chronic and frequently lethal disease caused by protozoan parasites of the Leishmania donovani complex, order Kinetoplastida. The etiological agents are: L. donovani in India and Central Africa and Leishmania infantum in the Americas, the Middle East, Central Asia, China, and the Mediterranean. The human disease is lethal if not treated soon after the onset of clinicopathological abnormalities that include: malaise, anemia, cachexia, hypergammaglobulinemia, hepato-splenomegaly, and progressive suppression of the cellular immune response. The L. donovani complex species are intracellular parasites of...
macrophages of lymphoid organs such as the spleen, lymph nodes, bone marrow, and liver. Their biological cycle alternates between the amastigote form in the vertebrate host and the promastigote form in the gut of the sand-fly vector (WHO, 2011).

Considering the presence or absence of animal reservoirs for *Leishmania*, two basic types of epidemiological cycles are noted: zoonotic (ZVL) or anthroponotic (AVL; reviewed by Palatnik-de-Sousa and Day, 2011).

AVL in India and Central Africa is caused by *L. donovani* and involves a severe parasitism of the blood and skin and an anthropophilic vector, making man the reservoir of the disease. In contrast, ZVL with dogs as the reservoir hosts is usually associated with *L. infantum* and is found in the Americas, the Middle East, Central Asia, China, and the Mediterranean. Sandflies become infected mostly by feeding on the skin of canids and humans are the final host of the parasites.

“One Health” proposes the unification of medical and veterinary sciences with the establishment of collaborative ventures in clinical care, surveillance and control of cross-species disease, education, and research into disease pathogenesis, diagnosis, therapy, and vaccination. The concept encompasses the human population, domestic animals and wildlife, and the impact that environmental changes (“environmental health”) such as global warming will have on these populations. “One Health” proposes taking a holistic view of the previously distinct disciplines of human medicine, veterinary medicine, environmental science, and wildlife conservation (Palatnik-de-Sousa and Day, 2011). ZVL is a disease that epitomizes perfectly the need for a One Health approach since the disease occurs in both, dogs and humans, with similar symptoms, clinical outcome, and commitment of the integrity of the T-cell related immunity (Reis et al., 2016; Palatnik-de-Sousa and Day, 2011). Tools for control of canine ZVL will have impact on the reduction of the human incidence of ZVL (Palatnik-de-Sousa et al., 2009).

The present review examines the current state-of-the-art of the development and use of vaccines for canine leishmaniasis as the most worthy cost–benefit tool for the control and prevention of ZVL.

**LEISHMANIASIS IN CANIDS**

A broad range of immune responses and clinical manifestations have been described in canine ZVL (Baneth et al., 2008; Solano-Gallego et al., 2011). These aspects have been thoroughly revised by the LeishVet group in order to establish guidelines for canine ZVL management (Solano-Gallego et al., 2011). Infection in dogs may be subclinical or manifested as a self-limiting disease, or a severe, and sometimes, even if treated, fatal illness (Solano-Gallego et al., 2011). Subclinical infection is not necessarily permanent and factors such as immunosuppression or concomitant diseases could break the equilibrium and lead to the progression of the clinical disease in dogs (Baneth et al., 2008; Solano-Gallego et al., 2009) as has been observed in humans coinfected with the human immunodeficiency virus and *Leishmania* (Alvar et al., 2008). Of note, the infection of cats by *L. infantum* was recently reported (Maia and Campino, 2011; Vides et al., 2011) and a correlation between cat infection by the feline immunodeficiency virus (FIV) and leishmaniasis was also described (Vides et al., 2011).

Since the proportion of clinical cases of canine ZVL is lower than that of the subclinical cases, and a positive correlation between clinical status and infectiosity to the sand-fly has been established (Travi et al., 2001), when high vector sand-fly and canine–host densities are present, the infection spreads quickly and extensively among the dog population (Quinnell et al., 1997, 2003; Oliva et al., 2006).

Several predisposing factors for the development of the disease have been described including breed, age, and genetic background. As for breed susceptibility, the German shepherd dog (Ranque et al., 1977; Abranches et al., 1991) together with the boxer (Ranque et al., 1977) and doberman (Abranches et al., 1991; Sideris et al., 1996) breeds appear to be predisposed to infection in France, Portugal, and Greece. In contrast, in Greece dogs of the collie breed are rarely infected (Sideris et al., 1996) and there is a well-known resistance in Ibizan hounds in Spain (Solano-Gallego et al., 2000). In Brazil however, the most affected breeds are the long-coated cocker spaniel (26.9%) and the short-coated boxer (24.6%) (França-Silva et al., 2003), while in Italy no breed-related predisposition has been reported (Pozio et al., 1981).

Age seems to be an important factor (Pozio et al., 1981; Sideris et al., 1996). The distribution of the disease is considered bimodal in Europe, with the highest prevalence reported in dogs younger than 3 years and older than 8 years (Abranches et al., 1991; Cardoso et al., 2004). This was not supported however by a Brazilian study of 33,937 dogs (França-Silva et al., 2003). No specific canine gender predisposition has been described for ZVL in various endemic countries (Pozio et al., 1981; Abranches et al., 1991; Sideris et al., 1996; Antônio et al., 2007); however, in France a greater prevalence of ZVL was found among male dogs (Lanotte et al., 1975) and in an endemic area of Brazil high rates of seropositivity were also found among male animals (Mouta-Confort et al., 2010).

ZVL in dogs is a systemic disease that may potentially involve any organ, tissue, or body fluid and is manifested by non-specific clinical signs. The most common clinical manifestations and clinicopathological abnormalities found in canine ZVL are thoroughly described by the LeishVet group (Solano-Gallego et al., 2011). Skin lesions are the most frequent manifestation among them and may be seen along with other clinical signs or clinicopathological abnormalities. Clinical manifestations include generalized lymphadenopathy, loss of weight, altered appetite, lethargy, mucous membrane pallor, splenomegaly, polyuria and polydipsia, fever, vomiting and diarrhea, dermatitis, onychogryphosis, blepharitis, conjunctivitis, keratoconjunctivitis, uveitis, oral, genital, and/or nasal mucocutaneous and mucosal lesions, epistaxis, vascular, and neurological disorders (Solano-Gallego et al., 2011). Among the main laboratory abnormalities, hypergammaglobulinemia, hypoalbuminemia, decreased albumin/globulin ratio, anemia, leukopenia, thrombocytopenia, proteinuria, renal azotemia, and elevated liver enzyme activities were described (Solano-Gallego et al., 2011).

Renal disease may be the sole clinical manifestation of canine ZVL and it can progress from mild proteinuria to the nephrotic syndrome or to an end stage renal disease. Chronic renal failure is a severe result of disease progression and the main cause of mortality due to ZVL. However, the variable and non-specific clinical
signs make the list of differential diagnoses wide and extensive (Solano-Gallego et al., 2011).

There have been numerous reports of the cutaneous pathology in the chronic stages of canine leishmaniosis and in some of these the presence of intracellular amastigotes within the macrophages forming the granulomatous dermatitis has been highlighted immunohistochemically (Koutinas et al., 1993; Maia and Campino, 2008; Day, 2011). Immunohistochemical studies have also shown that in the relatively milder clinical lesions of exfoliative dermatitis there is a low parasite burden, associated with enhanced expression of class II molecules of the major histocompatibility complex (MHC) by keratinocytes and a dermal T-cell infiltrate dominated by CD8+ cells over the CD4+ T helper (Th) subpopulation. In contrast, with the increasing severity of the cutaneous disease (the nodular form) there is reduced expression of class II molecules of the MHC by epidermal Langerhans cells and keratinocytes and fewer infiltrating T-lymphocytes (Papadogiannakis et al., 2005; Saridomichelakis, 2009). The lymph nodes of asymptomatic *Leishmania*-infected dogs are hyperplastic, but when the disease becomes systemic there is more often an atrophy of the lymph node cortex (Giunchetti et al., 2008a). The systemic pathological changes of canine ZVL have been reviewed recently (Day, 2011).

**CELLULAR IMMUNE RESPONSES IN CANINE ZVL**

Susceptibility and resistance to the development of canine leishmaniasis depends on the immune response elicited in the dog after natural infection. According to Reis et al. (2010), the evaluation of different factors like parasite burden, humoral response, cell-mediated immunity (CMI), or cytokine expression, have a predictive value for the progress of the infection and should, therefore, be taken into account as hallmarks of resistance and susceptibility to canine leishmaniasis (Reis et al., 2010). Increased levels of parameters like parasite load, IL-10, and TGF-β expression, *Leishmania*-specific cell immunodepression or *Leishmania*-specific IgG, IgM, IgA, and IgE serum antibodies are related to the clinical disease progression (Reis et al., 2010). By contrast, increased levels of parameters such as PBMCs proliferation after leishmanial antigen stimulation, IFN-γ and TNF-α expression, proliferation of CD4+, CD8+, and B-cell subsets, or a positive leishmanin skin test (LST) are related to resistance (Reis et al., 2010). The different balance established between the factors involved in the host immune response are reflected in the spectrum of clinical forms that can be observed in naturally infected dogs, ranging from symptomatic severe canine leishmaniasis to asymptomatic animals (Reis et al., 2010). In agreement with that, the increase of IFN-γ in cured cases was also described (Manna et al., 2008).

Other studies however, point out that the expression of IFN-γ was similar in symptomatic and asymptomatic dogs (Quinnell et al., 2001; Corrêa et al., 2007; Lage et al., 2007) or even higher and associated with the increase of parasite load and clinical status in susceptible dogs (Sanchez-Robert et al., 2008; Travi et al., 2009). Supporting these findings an initial asymptomatic phase, followed by a short-term production of Th1 type cytokines before the appearance of clinical signs (Santos-Gomes et al., 2002) or an initial elevation in IL-4 followed by an increase in the Th1-associated IFN-γ were described (Strauss-Ayali et al., 2007). Different from what Reis et al. (2010) stated, the high variability of the basal levels of these cytokines could not indicate the prognosis of the subsequent response against infection (Sanchez-Robert et al., 2008).

On the other hand, spleen cells from infected dogs showed a predominant expression of IL-10 that was positively correlated with parasitic load and clinical status severity (Pinelli et al., 1999; Chamizo et al., 2005; Lage et al., 2007; Alves et al., 2009).

A spectrum of innate and acquired immune responses to *L. infantum* infection is mounted by the canine–host (Baneth et al., 2006; Reis et al., 2010). Protective immunity to canine ZVL is T-cell-mediated and susceptibility to the overt disease is associated with a marked non-protective humoral immune response and a depressed CMI (Pinelli et al., 1994; Barbieri, 2006; Baneth et al., 2008).

When a cellular immune response is present, dogs appear asymptomatic and IDR (intradermal response to leishmanial antigen) positive, with higher levels of IL-2 and TNF-α and a mixed TH1/TH2 response involving TH1 evolution mediated primarily by IL-12, IL-18, and IFN-γ (Chamizo et al., 2005). In asymptomatic dogs, macrophages are capable of killing amastigotes via the nitric oxide route (Vouldoukis et al., 1996), and there is an increase in CD8+ T-cells in peripheral lymphocytes, and MHC class II molecules and the receptors CD45RA and CD45RB are also expressed (Reis et al., 2006a). In contrast, symptomatic dogs show a failed cellular response, with no IDR or lymphocyte proliferation and the following additional characteristics: decrease in CD14+ monocytes; decrease in CD8+ (Reis et al., 2006a), CD4+ T-lymphocytes, and CD21+ B lymphocytes, either *Leishmania*-specific or not; antibody increase; deficiency of the co-stimulatory response; decrease of IFN-γ; and reduced expression of MHC class II molecules, which renders the animal more susceptible (Bourdoiseau et al., 1997; Moreno et al., 1999; Pinelli et al., 1999; Guarga et al., 2000, 2002; Borja-Cabrera et al., 2004).

The majority of infected dogs are likely to develop positive specific CMI expressed as proliferation of lymphocytes stimulated in *vitro* by *Leishmania* antigen or in *vivo* by a positive skin test early in infection. However, as the disease progresses in susceptible dogs, these responses diminish. Blood parasite load and *Leishmania*-specific CMI were shown to be inversely correlated during a longitudinal follow-up of experimentally infected dogs (Rodriguez-Cortés et al., 2007a). CMI unresponsiveness in progressive disease has been postulated to be due to the decrease in peripheral CD4+ T-cell numbers or the decreasing expression of co-stimulatory molecules such as B7 (Pinelli et al., 1999; Guarga et al., 2000; Alvar et al., 2004; Barbieri, 2006).

It is now widely accepted that protective immunity against *Leishmania* parasites is mediated by CD4+ T helper1 (Th1) cellular responses (Strauss-Ayali et al., 2007) and that the decrease in CD4+ T-cell numbers is highly correlated with parasitism and with the infectious condition to the sand-fly vector (Guarga et al., 2000, 2002; Travi et al., 2001). Similar to the situation in human VL patients, symptomatic disease in dogs is correlated with a “suppressive” pattern of T-cell responses, with a dominant role of IL-10 in ongoing, non-protective immune responses (Nylen and Sacks, 2007; Alves et al., 2009; Carrillo and Moreno, 2009).
HUMORAL IMMUNE RESPONSE IN CANINE ZVL

Canine leishmaniosis is frequently associated with a marked humoral response, which is not protective and signifies failure to control the infection. The levels of *Leishmania*-specific immunoglobulins detected in symptomatic dogs are greater than those detected in infected but asymptomatic dogs, and a marked association was found between these levels, the clinical status, and the tissue parasite density (Reis et al., 2006b).

Canine ZVL shows very similar characteristics to the human disease (Palatnik-de-Sousa, 2008; Kaye and Aebischer, 2011). IgA, IgE, and IgM responses have been shown to be markers of the disease (Almeida et al., 2003; Iniesta et al., 2003; Rodríguez-Cortés et al., 2007b). An IgG antibody increase is also observed and correlated with symptomatology (Keenan et al., 1984; Courtenay et al., 2007b). An IgG antibody increase is also observed and correlated with symptomatology (Keenan et al., 1984; Courtenay et al., 2007b).

In dogs vaccinated with the CPa and CPb cysteine proteinases (Killick-Kendrick et al., 1997; Maroli et al., 2001) is referred to as the “culling control campaign” of infected dogs and of nets (with or without insecticides) in human homes and dog kennels is also recommended. A decrease in the prevalence of canine ZVL was observed after 65% permethrin spot-on treatment of dogs (Giffoni et al., 2002) and a significant reduction of anti-*Leishmania* antibody titers in dogs was observed in dogs using insecticide-impregnated collars (Killick-Kendrick et al., 1997; Maroli et al., 2001) and spot-on repellents (Miró et al., 2007; Ferroglio et al., 2008; Thomas et al., 2008). In southern Europe the use of deltamethrin-impregnated dog collars (Killick-Kendrick et al., 1997; Maroli et al., 2001) is preferred however they do not prevent relapses (Baneth and Shaw, 2002).

WHAT ARE THE SUITABLE ACTIONS FOR THE CONTROL OF CANINE ZVL

For the control of leishmaniasis, WHO recommends: (1) the treatment of human patients, (2) the culling of seropositive infected dogs, and (3) the insecticidal treatment of human homes (Tesh, 1995; WHO, 2011).

Human cases should be diagnosed and treated as swiftly as possible (Ministério da Saúde, 2006). However in the case of ZVL humans are not the main reservoir of the infection.

A second preventive tool for ZVL in Brazil is control of the canine reservoir by removal and humane destruction of the *Leishmania*-seropositive and/or infected dogs and elimination of their carcasses (Ministério da Saúde, 2006). Although many studies support the efficacy of the culling control campaign (Magalhães et al., 1980; Ashford et al., 1993; Braga et al., 1998; Jerónimo et al., 2000; de Oliveira and de Araújo, 2003; Costa et al., 2007; Nunes et al., 2010) a few investigations show opposite results (Dietze et al., 1997; Paranhos-Silva et al., 1998; Moreira et al., 2004; De Souza et al., 2008; Nunes et al., 2010) and this strategy is frequently debated (Quinnell and Courtenay, 2009; Romero and Boelaert, 2010). At present, the removal of infected dogs is undertaken systematically only in Brazil (Palatnik-de-Sousa et al., 2001; Romero and Boelaert, 2010) and eventually in other South American countries (Romero and Boelaert, 2010) and China (Wang et al., 2010) and not regularly performed in the Mediterranean basin where chemotherapy treatment is preferred however they do not prevent relapses (Baneth and Shaw, 2002).

A third tool for control is the monitoring of the sand-fly vector in human residences and peri-domestically by spraying with pyrethroids. The use of deltamethrin-impregnated collars in dogs and of nets (with or without insecticides) in human homes and dog kennels is also recommended. A decrease in the prevalence of canine ZVL was observed after 65% permethrin spot-on treatment of dogs (Giffoni et al., 2002) and a significant reduction of anti-*Leishmania* antibody titers in dogs was observed in dogs using insecticide-impregnated collars (Killick-Kendrick et al., 1997; Maroli et al., 2001) and spot-on repellents (Miró et al., 2007; Ferroglio et al., 2008; Thomas et al., 2008).

In southern Europe the use of deltamethrin-impregnated dog collars (Killick-Kendrick et al., 1997; Maroli et al., 2001) is preferred by many dog owners. Alternatively, there is substantial evidence for the efficacy of spot-on repellents containing imidacloprid, permethrin, pyriprole, metaflumizone, or amitraz (Miró et al., 2007; Ferroglio et al., 2008; Thomas et al., 2008). In the Mediterranean region, human and canine cases of ZVL are treated with antiparasitic drugs. In Europe, individual measures to protect dogs from sand-fly bites using insecticides are commonly practiced, but no public health surveillance and control interventions such as those applied in Brazil are in place (Palatnik-de-Sousa et al., 2001).

The mathematical model described by Dye condemned the epidemiological ZVL control campaign, considering it non-efficient and indeed, at low rates of canine seropositivity, no impact on the human incidence of the disease was observed (Dye, 1996; Palatnik-de-Sousa et al., 2004). However, at higher rates of canine seropositivity, corresponding to more sensitive diagnostic methods, the number of infectious dogs declined interrupting the transmission and the spread of epidemics (Palatnik-de-Sousa et al., 2004). The low acceptance of culling of companion animals by their owners, the ethical dilemmas of veterinarians and humane reasons demand the development of alternative preventive tools. Mathematical modeling suggests that vector control and vaccination of
dogs and/or humans would be more efficacious than dog culling (Dye, 1996).

**VACCINES FOR HUMAN LEISHMANIASIS**

In spite of an estimated 500,000 cases and more than 50,000 deaths annually (WHO, 2011), human VL has been considered an unattractive vaccine target for industry, being primarily a disease of the poor. Human VL however, ranks second only to malaria for mortality and fourth for morbidity amongst tropical parasitic diseases (Mathers et al., 2007).

Although the century-old practice of leishmanization, the deliberate infection of naive people with virulent *Leishmania major*, is defended by some authors for vaccination against tegumentary leishmaniasis (Khamesipour et al., 2006), the development of lesions resistant to treatment makes this practice unrecommended. Furthermore, while first generation vaccines of whole killed parasite vaccines have been developed and tested against CL and VL their average clinical efficacy was considered to be low (54%; reviewed in Palatnik-de-Sousa, 2008). Indeed, the only field trial against human VL was performed in Sudan (Khalil et al., 2000), with an autoclaved *L. major* vaccine with BCG and achieved 43.3% of vaccine efficacy (VE = incidence in non-vaccinated individuals – incidence in vaccinated individuals/incidence in non-vaccinated individuals × 100), only among LST converters, which is impressive considering the high mortality and virulence of kala-azar in that area. The safety of this preparation on children was also assayed (Khalil et al., 2006). These data however are contested by a recent metanalysis (Noazin et al., 2010).

Only a single product (Leish-111f), a fusion protein of three relatively conserved *L. major* proteins (thiul-specific antioxidant, stress inducible protein 1, and elongation initiation factor) formulated with MPL-SE is entering phase II clinical testing in humans, including human VL as a therapeutic vaccine (Coler and Reed, 2005; Nascimento et al., 2010) but there is no human vaccine licensed for prophylaxis against VL.

**FIELD ASSAYS OF VACCINES AND LICENSED VACCINES FOR CANINE LEISHMANIASIS**

Many *Leishmania* antigens have been identified as potential vaccine candidates against canine ZVL (reviewed by Palatnik-de-Sousa, 2008), but very few have been tested in field assays.

The first generation vaccines developed for humans were also tested against canine ZVL. In an initial comparison of vaccine formulations, with no challenge, Lasri et al. (1999) demonstrated that dogs vaccinated with autoclaved *L. major* promastigote lysate (ALM) and BCG showed an *in vitro* lymphocyte proliferative response, while dogs immunized with ALM and saponin expressed a main humoral antibody response against *L. infantum*. Mayrink and co-workers, using a *Leishmania braziliensis* lysate and BCG vaccine, in a formulation related to the Leishvacin, previously shown to be about 50% protective against tegumentary leishmaniasis in humans (Antunes et al., 1986), obtained 90% protection against experimental canine kala-azar in the kennel (Mayrink et al., 1996) but failed to detect any significant difference between vaccines and placebos in a well designed field Phase III assay (Genaro et al., 1996; Table 1). The 90% protection in the kennel assay was related to the absence of the parasite in bone marrow cultures of 9 of the 10 vaccinated dogs that were challenged with only 10⁶ cultured promastigotes of *Leishmania chagasi* (Mayrink et al., 1996). The difference between the artificial and the natural challenge in this case could account for the failure of the vaccine in the field assay. In Iran, on the other hand the use of Aluminum hydroxide precipitated *L. major* vaccine plus BCG resulted in 69.3% of vaccine efficacy against the natural infection by *L. infantum* in Iran (Mohebali et al., 2004; Table 1). The incidence rate was 3.7% (6/162) in vaccinated dogs and 12.0% (17/141) in control dogs.

In the first Phase III trial with a second-generation dog vaccine, Dunan et al., using a *L. infantum* semi-purified lyophilized protein preparation (94–67 kDa), paradoxically achieved, a significantly higher rate of infection in vaccinated than in control dogs. This vaccine then, while effective in murine models, developed no protection against canine kala-azar in the field (Dunan et al., 1998; Table 1).

Two dog vaccines achieved successful results in Phase III trials: the fucose–mannose ligand (FML)–saponin vaccine (Leishmune®; da Silva et al., 2001; Borja-Cabrera et al., 2002, 2010; Palatnik-de-Sousa et al., 2008) composed of the antigen of *L. donovani* promastigotes, named FML (Palatnik et al., 1989) and the QS21 and deacetylated saponins of *Quillaja saponaria* (Oliveira-Freitas et al., 2006; Table 1). The FML was antigenic for human (Palatnik-de-Sousa et al., 1995) and dogs (Borja-Cabrera et al., 1999) and Leishmune® was immunogenic, immunoprophylactic, and immunotherapeutic in mice and hamsters and field trials for dogs (Palatnik-de-Sousa et al., 1994; Santos et al., 2002, 2003, 2007; Borja-Cabrera et al., 2004, 2008, 2010). In the first Phase III assay, four obits and six symptomatic cases among 30 surviving placebo treated dogs (33%) were detected and confirmed by parasite analysis and PCR. No obits were detected among vaccines and infection was confirmed in 3/36 oligosymptomatic dogs (8.33%), making 92% protection, and 76% vaccine efficacy (da Silva et al., 2001). In the second assay, the infective pressure was higher and 2 years after vaccination, obits were detected in 8/33 (25%) of the placebo treated and 1/20 (5%) of the vaccinated dogs, making 95% protective effect and 80% vaccine efficacy. This protection lasted for at least 3.5 years and was concomitant with the reduction of the human incidence of the disease in the area (Borja-Cabrera et al., 2002). Noteworthy, the VE values for the FML–saponin vaccine revealed protection against severe diseases and obits due to canine ZVL (Palatnik-de-Sousa et al., 2008).

The FML-vaccine was considered a second-generation vaccine candidate and participated in the fourth Meeting on Second-Generation-*Leishmania* vaccines held in Mérida in May 2001 (Dumonteil et al., 2001). In 2003, the FML–saponin vaccine was licensed for prophylaxis against canine ZVL in Brazil under the name of Leishmune®, and has been used in Brazil since 2004 and it obtained its definitive license in October, 2011.

The vaccine was considered safe and was well tolerated (Parra et al., 2007; Table 1). After 2 years of vaccination of a cohort of 550 Leishmune® vaccinated exposed dogs, only 1% of the animals died of ZVL and 1.2% were symptomatic. Simultaneously, 39% of deaths and 20.6% of symptomatic cases were detected among untreated exposed control animals (p < 0.005; Borja-Cabrera et al., 2008). The Leishmune® vaccine is prophylactic
Table 1 | Canine vaccines for visceral leishmaniasis assayed in field trials.

| Vaccine         | Composition                              | Licensed* | VE (%)** | Results obtained                                                                 |
|-----------------|------------------------------------------|-----------|----------|----------------------------------------------------------------------------------|
| Leishvacin      | L. braziliensis lysate + BCG             | –         | –        | Protective in kennel assay but failed in Phase III assay in Brazil                |
| ALM             | Aluminum hydroxide-L. major              | –         | 69.3     | Protective against natural infection by L. infantum in Iran                       |
| Lyophilized protein preparation | L. infantum semi-purified proteins (94–67 kDa) | –         | –        | Protective in mice assays but not in field trial in France                        |
| Leishmune®      | L. donovani FML antigen and QS21 and deacylated saponins of Quillaja saponaria | +         | 76–80    | Immunogenic, immunoprophylactic, and immunotherapeutic in mice and hamsters       |
|                 |                                          |           |          | Long-lasting protection against infection, severe disease, and deaths of ZVL in dog field trials in Brazil |
|                 |                                          |           |          | Safe and well tolerated                                                          |
|                 |                                          |           |          | Vaccinated dogs do not expose parasites and are negative in xenodiagnosis         |
|                 |                                          |           |          | With double saponin concentration is therapeutic against naturally or experimentally acquired ZVL. In immunochemotherapy promotes the sterile cure |
|                 |                                          |           |          | Leishmune® enhances the levels of IFN-γ, NO, and anti-L. chagasi IgG2, the early and persistent activation of neutrophils and monocytes |
|                 |                                          |           |          | Leishmune® increases the CD8+ T-cells expressing IFN-γ and sustain or increases the proportions of CD4+ and CD21-B lymphocytes |
|                 |                                          |           |          | Leishmune® vaccination does not interfere in epidemiological serological control tests |
|                 |                                          |           |          | A decrease in human and canine incidence and in canine seroprevalence of ZVL was observed after dog vaccination in two Brazilian towns |
| LiESAP          | 54 kDa excreted protein of L. infantum + MDP | –         | 92%      | Long-lasting protection against, infection but not against deaths or severe disease by ZVL, in a field assay performed in France at a lower infective pressure endemic region |

*For prophylaxis against canine visceral leishmaniasis.
**VE, vaccine efficacy in Phase III trials.

against canine ZVL VL, protects 98% of vaccinated dogs (Borja-Cabrera et al., 2008), and reduces the parasite burden accessible for transmission back to sand flies, as disclosed by their negative results of PCR for Leishmania DNA in blood and lymph nodes and negative immunohistochemistry reactions in skin (Nogueira et al., 2005). The generated antibodies block the transmission of the disease by sand flies in the field (Saraiva et al., 2006; Palatnik-de-Sousa et al., 2008). Indeed, while sand flies fed upon pre-immune sera showed higher percent of infection and number of parasites/insect, giving a high infection index, sand flies fed upon Leishmune® sera (mainly IgG2 anti-FML antibodies) showed a 74.3% reduction in infection; (Saraiva et al., 2006). On the other hand, compared to the respective pre-immune fed controls, the sand flies fed on sera of infected animals (IgG1 predominant antibodies) showed, a pronounced enhancement of infection (331.9%; Palatnik-de-Sousa et al., 2008). Xenodiagnosis using Lutzomyia longipalpis disclose that 2/9 naturally infected dogs, infected phlebotomies, while 0/19 Leishmune® vaccinated dogs did not. The two infected dogs showing positive xenodiagnosis were the most symptomatic (more than three clinical signs) among the nine tested, while the other seven were asymptomatic or oligosymptomatic (Palatnik-de-Sousa et al., 2008; de Amorim et al., 2010). The anti-FML antibody response induced by Leishmune® is mainly of the IgG2 subtype (Mendes et al., 2003). A total of 150,000 healthy dogs were vaccinated in Brazil up to October 2011. We also observed that Leishmune® formulated with double saponin adjuvant concentration has a therapeutic effect against naturally (Borja-Cabrera et al., 2004) or experimentally acquired ZVL (Santos et al., 2007). While immunotherapy with the saponin-enriched Leishmune® reduces the symptomatology, the rate of obits and the parasite load in lymph nodes, immunochemotherapy with Leishmune®, Allopurinol, and Amphotericin B promotes the sterile cure, turning to negative the PCR reactions for Leishmania DNA (Borja-Cabrera et al., 2010).

Leishmune® induced an immunological pattern characterized by enhanced levels of IFN-γ, NO, and anti-L. chagasi IgG2 (Araújo
et al., 2009), the early and persistent activation of neutrophils and monocytes, and increased the CD8+ T-cells expressing IFN-γ (Araújo et al., 2008; Table 1). This increase of CD8+ T-cells is expected for the QS21 saponin adjuvant of Leishmune® (Oliveira-Freitas et al., 2006) and it was also described in the immunotherapy assays against naturally (Borja-Cabrera et al., 2004) and experimentally acquired ZVL (Santos et al., 2007). Leishmune® also induced early phenotypic changes in neutrophils (increase in MHCII+ and decrease in CD32+ and CD18+ activation markers) and monocytes, CD8+ T-cell activation, and a selective pro-inflammatory pattern (IFN-γ/NO; Araújo et al., 2011). Other studies also revealed the sustained or increased proportions of CD4+ and CD21-B lymphocytes and the increased proportions of CD8+ T-cells (Borja-Cabrera et al., 2004, 2010; Santos et al., 2007) and the diminished CD4+/CD25+ T-cell counts and increased IFN-γ levels in dogs vaccinated with Leishmune® (de Lima et al., 2010). Similar to the situation of human VL patients, the symptomatic disease in dogs is correlated with a “suppressive” pattern of T-cell responses, with a dominant role for IL-10 in ongoing, non-protective immune responses (Nylen and Sacks, 2007; Carrillo and Moreno, 2009). In view of this similarity between human and canine VL, the fact that Leishmune® has shown promise as a therapeutic vaccine in ZVL is encouraging. The therapeutic efficacy in ZVL of Leish-110f, which induces mainly B-cell and CD4+ T-cell responses, was indistinguishable from the effect of the MPL-SE adjuvants alone (Miret et al., 2008). This may suggest that a therapeutic vaccine against human VL should, akin to canine ZVL, aim to induce both CD4+ and CD8+ T-cells (Kaye and Aebischer, 2011).

Since canine ZVL in Brazil is epidemiologically controlled by serological enquire, and Leishmune® was designed to induce a strong humoral response against the FML antigen, to facilitate the monitoring during field assays, there was a concern that vaccinated dogs could not be differentiated serologically from infected dogs (Marcondes et al., 2011). The predominance of IgG2 antibodies to FML, on the other hand, was proposed to be a tool for differentiation (Mendes et al., 2003). The results obtained from the control campaign disclosed however that there was no reason for concern since in the field, dogs vaccinated with Leishmune® did not become seroreactive in the official test used by the enquire (Palatnik-de-Sousa et al., 2009; Table 1). In the epidemic area of Campo Grande, Brazil, the serological control campaign evaluated 110,000 dogs, 5,860 of which were Leishmune® uninfected vaccines. Only 1.3% of positivity (76 among 5,860) was detected. These seropositive dogs were seronegative for the L. chagasi HSP recombinant antigen and showed no parasite evidence in bone marrow and lymph nodes (Palatnik-de-Sousa et al., 2009).

A possible additive effect of Leishmune® vaccination over dog culling, on the decrease of the incidence of canine and human ZVL was studied in two Brazilian endemic areas, from 2004 to 2006 (Palatnik-de-Sousa et al., 2009; Table 1). In Araçatuba, a decline of 25% was seen in the incidence of ZVL in dogs with a 61% decline in human cases (36–14 cases), indicating the additive effect of Leishmune® vaccination over regular dog culling. In Belo Horizonte, where 8.1% (12,113/149,470) of the dogs were vaccinated up to 2006, the districts that had had greater vaccine coverage (85.7% of the doses) exhibited declined or sustained levels of canine and human cases of ZVL, while those with less vaccine coverage (14.3% of the doses), showed rising curves of canine and human cases of the disease. In the districts with higher vaccination levels, human cases declined by 36.5% falling from 2004 to 2006 outside of the 95% confidence interval (CI95%) of the less vaccinated districts (CI95% 2.23–21.11), which showed an average increase of 11.67%. From 1999 to 2006, the increase of canine seroprevalence and of human cases of the disease in all districts were significantly correlated (p = 0.001), confirming the importance of the dog as the infectious reservoir of the disease. The decrease in dog culling (−p = 0.007) and human incidence (−p = 0.043) were significantly correlated with the increase in the number of vaccinated animals, indicating the prophylactic impact of Leishmune® vaccination on the decrease of the proportions of infectious dog and human populations, and so indirectly indicating a decrease in the number of dogs sacrificed (Palatnik-de-Sousa et al., 2009).

The mathematical model for control of leishmaniasis developed by Dye compared the efficacy of various control methods in the reduction of human and canine incidence of the disease (Dye, 1996). According to this model, and regarding the canine incidence, killing infected dogs is the least successful strategy while an effective veterinary drug should perform better, even with the conservative assumption that only infectious dogs are treated. A dog vaccine has higher impact, because it is prophylactic and the insecticide treatment would be the best approach for dogs. The expected outcome of control on human incidence is that killing dogs is again the least successful strategy and that immunizing people is less effective than immunizing dogs because it has no impact on the dynamics of infection in the dog population. The insecticide control would be the most effective strategy.

We used the data of human and canine incidence obtained before and after Leishmune® vaccination in Araçatuba and Belo Horizonte (Palatnik-de-Sousa et al., 2009) in order to see if they are related to what expected according the control model of Dye (1996; Figures 1A,B). The expected efficacy for the vaccine in bringing down the canine incidence of the disease is disclosed by plotting along the y axis the incidence of VL in dogs after vaccination divided by its value before vaccination and expressed as a percentage, and in the x axis the fraction of the susceptible dogs converted to resistant by the vaccine treatment (Dye, 1996). In Araçatuba, Brazil, the VL incidence in dogs after vaccination was 21.77% (2006) and before vaccination 30.01% (2003) giving a percent ratio of 72.54% which corresponds to a transformation of 25% of the dog population from susceptible to resistant or protected (Palatnik-de-Sousa et al., 2009; Figure 1A). This was achieved vaccinating only 7.531% of the dog population. In Belo Horizonte, on the other hand, the decline in human incidence was observed after dog vaccination in several districts. In the Nordeste district, human cases declined from 24 to 18 giving a percent ratio of 72% which corresponds to 28% of the human population moving to a resistant class (Palatnik-de-Sousa et al., 2009; Figure 1B). This was achieved by vaccinating only 11.03% of the whole dog population.

The other second-generation vaccine, LiESAp, composed of the 54-kDa excreted protein of L. infantum plus MDP also reached Phase III trials after a kennel assay against L. infantum infection (Table 1). Parasites were detected in the bone-marrow of 3/3
FIGURE 1 | Impact of the use of Leishmune® vaccine on the decrease of dog and human incidence of ZVL in Brazil. The curves show the expected efficacy of different control methods in bringing down the (A) prevalence of infectiousness in dogs, and (B) the incidence of human visceral leishmaniasis according to the model of Dye (1996). The y-axes show the stable prevalence (A) or incidence (B) after control, divided by the respective values before control, and expressed as a percentage and the X axes record the percent change promoted by each control method of control moving a fraction of susceptible dogs or people to the resistant class. The arrows show the percent of dog (A) and human (B) incidence after/before Leishmune® vaccination in 2004–2006 and their respective plot of the percent of dog (A) and human (B) population that became protected and resistant to infection after vaccine treatment.

placebo treated controls, while they were absent in 3/3 vaccinated dogs (Lemesre et al., 2005). The double blind random trial further performed with LiESAp + MDP in naturally exposed dogs of South France revealed that after 2 years, an incidence of infection of 0.61% (1/165) in vaccines versus 6.86% (12/175) in control dogs making a 92% VE (Lemesre et al., 2007). In any dog showing clinical and/or serological evidence, infection was confirmed by the presence of parasites in bone marrow aspirates cultured in NNN media and also by PCR analysis (Lemesre et al., 2007). Differently from what was described for the FML–saponin vaccine (da Silva et al., 2001; Borja-Cabrera et al., 2002, 2008), the LiESAp vaccine induced protection against infection (Lemesre et al., 2007) but not against severe disease or death by VL. No obits at all, were described in the 2 years LiESAp assay (Lemesre et al., 2007), pointing to the lower infective pressure of the endemic region. The immune response generated in the LiESAp vaccinated dogs was confirmed by the finding of: a significant leishmanicidal effect of macrophages due to an IFN-γ dependent activation; a NO-mediated apoptosis of intracellular amastigotes (Holzmuller et al., 2005), a strong and long-lasting CMI revealed by positive IDR, an anti-leishmanial activity of monocytes, and by the in vitro activities of the anti-LiESAp antibodies (Bourdoiseau et al., 2009).

In Europe, a formulation related to the LiESAp vaccine was licensed for commercialization under the name of CaniLeish®, in
early 2011 and is currently being launched in Portugal, Spain, France, Greece, and Italy. It is as a second-generation defined peptidic antigen composed of Excreted–Secreted proteins of the supernatant of cultures of *L. infantum*. Differently from what was published before (Holzmuller et al., 2005; Lemesre et al., 2005, 2007; Bourdoiseau et al., 2009) this formulation does not include MDP but the QA21 adjuvant of *Q. saponaria* Molina instead (Virbac, 2011; Table 1).

As explained by WHO guidelines (WHO, 1997), confirmation of infection by very sensitive methods such as PCR or culture (Lemesre et al., 2005, 2007) represents a very early end-point of the development of the disease, while kala-azar obits and severe clinical cases (da Silva et al., 2001; Borja-Cabrera et al., 2002) are distant end-points which occur much later in the development of infection. A comparison between efficacies of Leishmune® and LiESAp should only be performed using the same infective pressure and the same end-point targets (WHO, 1997).

Finally, a vaccine called Leish-Tec® was also licensed in Brazil. It is composed of the recombinant A2-antigen of *Leishmania* amastigotes and is adjuvanted by saponin (Fernandes et al., 2008). While protection due to the Leishmune® vaccine has been extensively investigated in laboratory models (reviewed by Palatnik-de-Sousa et al., 2008), explained through an immunological approach (Araújo et al., 2008, 2009, 2011), and reported in: control versus experimental kennel assay with Leish-Tec® which was tested on seven dogs and compared with four untreated controls (Fernandes et al., 2008). There is no information about the infectivity of the strain used for challenge in that study and the lack of deaths in the control animals suggests that the challenge was mild. There are no reports of controlled-trial-field studies with Leish-Tec®, but despite the lack of peer-reviewed scientific publications, the vaccine was licensed in Brazil in 2008.

Since it was demonstrated that a canine vaccine promotes a decrease in the human and canine incidence of ZVL (Palatnik-de-Sousa et al., 2009), an increase in vaccine coverage for dogs, especially in the absence of a licensed human vaccine for leishmaniasis, would certainly bring about an interruption of epidemics.

**VACCINES FOR CANINE LEISHMANIASIS UNDER DEVELOPMENT**

The *L. braziliensis* lysate of a first generation vaccine against CL, which failed to show efficacy against canine ZVL when formulated with BCG, is now under research but adjuvanted with saponin (Giunchetti et al., 2007, 2008b). The vaccine increased the anti-*Leishmania* IgG isotypes, together with higher levels of lymphocytes, particularly circulating CD8(+) T-lymphocytes, and *L. chagasi* antigen-specific CD8(+) T-lymphocytes, as expected for a *Q. saponaria* saponin containing vaccine (Giunchetti et al., 2007). When combined to sand-fly saliva proteins it revealed an additional increase in circulating CD21+ B-cells, CD5+, and CD4+ T-cells (Giunchetti et al., 2008b).

There is a general consensus stating that the capacity to respond to multiple antigens may be an essential requisite of an efficacious vaccine (Working Group on Research Priorities for Development of Leishmaniasis Vaccines et al., 2011). Multiple antigens are included in a first generation or second-generation complex vaccines such as Leishmune® or LiESAp (Lemesre et al., 2007; Palatnik-de-Sousa et al., 2008) that already showed strong efficacy in field assays. However, from the industrial point of view, obtaining of protozoa vaccines has some limitations. Growing protozoa and extracting native antigens is laborious, expensive, and requires additional bovine products that are forbidden to be used since the emergence of bovine spongiform encephalopathy. On the other hand many subunits of *Leishmania* antigens have already been described, used alone, or in combination and are even available in the recombinant form (reviewed by Palatnik-de-Sousa et al., 2008; Working Group on Research Priorities for Development of Leishmaniasis Vaccines et al., 2011). These defined vaccines could be used in future multiple recombinant vaccines against canine ZVL.

The most promising candidates seem to be the LACK, LeIF, TSA, LmSTI1, H1, Cpa + CPb, KMP11, and NH36 (reviewed by Palatnik-de-Sousa, 2008). The last approach in second-generation vaccines is the use of recombinant proteins that were tested in kennel assays in a dog model against ZVL (Molano et al., 2003; Fujiwara et al., 2005; Gradoni et al., 2005; Poot et al., 2006; Moreno et al., 2007). None have advanced to Phase III dog trials. The multicomponent Leish-111f fusion protein containing the antigens TSA, LmSTI1, and LeIF; in formulation with MPL-SE or AdjuPrime, was immunogenic in dogs challenged with *L. chagasi* (Fujiwara et al., 2005) and *L. infantum* (MML; Moreno et al., 2007), but failed to prevent *L. infantum* natural infection, or disease progression in dogs in an open kennel trial (Gradoni et al., 2005). However it was useful as an adjunct therapy together with Glucantime in treating a field population of dogs suffering from ZVL due to *L. chagasi* infection (Miret et al., 2008). The vaccination was safe and induced a twofold to threefold increase in antigen-specific proliferative response in vitro after cure, but was lacking a clear clinical benefit (though the trial was not powered to reveal small effects; Kaye and Aebischer, 2011). Leish-110f in the form of an experimental vaccine designated MML was also tested alongside recombinant *L. infantum* histone H1 and hydrophilic acylated protein B1 (HASPB1; Moreno et al., 2007) as prophylactic vaccines against experimental ZVL. Dogs were vaccinated with either MML adjuvanted with MPL-SE or H1 or HASPB1 adjuvanted with Montanide-ISA 720 and subsequently challenged with 10³ *L. infantum* promastigotes. All vaccines were immunogenic, however they developed different boost responses after infection. For example, MML antibody responses were strongly boosted, HASPB1 responses weakly so and H1 response unaffected by infection (Moreno et al., 2007). Fewer H1 and HASPB1 immunized dogs developed CVL symptoms (37 and 50%, respectively) compared with control or MML-vaccinated dogs (71 and 75%, respectively), but larger studies would be needed to confirm the efficacy of these vaccines (Palatnik-de-Sousa et al., 2008; Kaye and Aebischer, 2011; Working Group on Research Priorities for Development of Leishmaniasis Vaccines et al., 2011).

Compared to recombinant protein vaccines, DNA vaccines are much more stable and have the advantage of low production costs,
no need for cold chain distribution, and the flexibility of combing
multiple genes in a simple construct. A lot of interest has
been generated for the development of a vaccine against leish-
mania in recent years, with studies going on in the labs on
experimental models. LACK, LeIF, TSA, LmST11, H1, CpaA + Cpb,
KMP11, and NH36 are the most promising candidates that may
find a place in the forthcoming years, since they have already been
tested in various animal models (reviewed by Palatnik-de-Sousa,
2008).

Immunization with HPB-LACK protected dogs against ZVL
(Ramiro et al., 2003) by increasing the IFN-γ and IL-12 expres-
sion, lymphocyte proliferative response, and the IgG2 to IgG1 ratio
and it led to decreasing clinical symptoms, number of parasites
in target tissues, and IL-4 expression. A prime boost vaccina-
tion with CpaA + Cpb, Montanide 720, and CPG protected dogs
against ZVL, as evidenced by the increase in IgG2 specific anti-
body synthesis, lymphocyte proliferation, IFNγ/IL-10 secretion,
and DTH response (Rafati et al., 2005). No death or clinical signs
were reported, probably due to the very low infective challenge
(5 × 10⁶ promastigotes). Eight out of 10 vaccinated dogs were
considered protected based on their PCR negative results; however,
no significant conclusion could be drawn from the study, because
it used only two untreated control dogs (Rafati et al., 2005). A
cocktail of plasmid DNA encoding KMPII, TRYP, LACK, and
GP63 did not protect dogs against L. infantum virulent challenge
(Rodríguez-Cortés et al., 2007b).

Recently, a dominant antigen in the FML complex, a nucleo-
side hydrolase of 36 kDa, has been shown in the recombinant
form or as a DNA vaccine to reproduce some of the Leishmune®
effects in mice (Aguilar-Be et al., 2005). The NH36 DNA vac-
cine protected mice against infection by L. chagasi, Leishmania
mexicana (Aguilar-Be et al., 2005), and Leishmania amazonen-
sis (Souza and Palatnik-de-Sousa, 2009), indicating its potential
usefulness in a bivalent immunoprophylactic vaccine for the
control of both endemics. Protection by the DNA vaccine was higher
than that induced by the recombinant NH36 or the FML anti-
gen plus saponin, and it is related to IFN-γ-producing CD4+ T-cells, which are characteristic of a TH1 type immune response
(Aguilar-Be et al., 2005). The DNA vaccine showed prophylac-
tic and immunotherapeutic effects against ZVL in a kennel assay
(Borja-Cabrera et al., 2009, 2012). The immunotherapy treatment
increased survival and reduced clinical status of L. chagasi-highly
infected dogs through the enhancement of a NH36-specific CD4+ T-
cell response (Borja-Cabrera et al., 2012).

The protection induced by the NH36 recombinant vaccine is
related to its C-terminal domain bearing the required T-cell epitopes that are responsible for a main CD4+ T-cell-mediated
immune response that has the enhancement of the IDR reac-
tion and the increases in ratios of TNFa/IL-10 CD4+ producing
cells as strong correlates of protection (Nico et al., 2010). The less
defined but nevertheless effective Leishmune® vaccine may there-
fore become replaced by a synthetic product comprising its active
ingredients (Kaye and Aebsicher, 2011).

The control of the companion dog population for VL is funda-
mental in order to avoid the spread of the disease between humans
and dogs. A high level of infection in dogs, particularly in impover-
ished areas, just before the start of the human epidemic, or during
the epidemic, or a rising prevalence of canine infection before the
epidemic all strongly predicted a high incidence of human VL
(Werneck et al., 2007). Recent research demonstrates that the use
of insecticides both in residences and in impregnated dog neck-
laces and/or the use of preventive canine vaccine could potentially
substitute the dog culling contributing to the eradication of the
disease. In any case, the research on human vaccines and the large
use of canine vaccines against VL should be stimulated in order to
reduce the incidence of both the canine and human disease.

ACKNOWLEDGMENTS
The author thanks David Straker for the English language editing
of this manuscript.

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Conflict of Interest Statement: The author declares that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 06 December 2011; accepted: 20 March 2012; published online: 17 April 2012.

Citation: Palatnik-de-Sousa CB (2012) Vaccines for canine leishmaniasis. Front. Immun. 3:69. doi: 10.3389/fimmu.2012.00069

This article was submitted to Frontiers in Microbial Immunology, a specialty of Frontiers in Immunology.

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