**Leishmania infantum** and **Dirofilaria immitis** coinfection in dogs in Greece

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**SUMMARY**

Leishmaniasis and dirofilariasis are parasitic diseases of humans and dogs, worldwide, and they are often found as coinfections in endemic areas. Cases of human and canine dirofilariasis have been reported in Greece and leishmaniasis is endemic in most prefectures in humans and dogs. In most cases, dirofilariasis is established by parasitological (the modified Knott’s test) and/or immunological methods, whilst for leishmaniasis molecular techniques and culture are also used. During an epidemiological study in Greece, 22.1% of the 5772 dogs studied were found positive by serology for *Leishmania*. Blood cultures of 165 (12.94%) of these animals produced *Leishmania* promastigotes and 26 (2.03%) *Dirofilaria microfilariae* (L1), whilst only in two (0.16%) both *Leishmania* and *Dirofilaria* L1 appeared. The aim was to assess coinfections by the two parasites in dogs in Greece, the isolation and survival of *Dirofilaria* microfilariae and *Leishmania* promastigotes using clotted blood (a fast, simple and low-cost method) and the survival potential of the two parasites in coexistence, *in vitro*.

Key words: isolation, dog blood, *Dirofilaria microfilariae*, *Leishmania infantum*, coinfection.

**INTRODUCTION**

Leishmaniasis and dirofilariasis are both vector-borne diseases. *Leishmania* (Trypanosomatida, Trypanosomatidae) uses sand flies (*Phlebotomus* spp.) (WHO, 1997) and *Dirofilaria* (Spirurida, Onchocercidae) uses mosquitoes (species of *Aedes*, *Culex*, *Anopheles* and *Mansonia*) for their transmissions to mammalian hosts (Otto and Jachowski, 1981). During a blood meal, an infected sand fly introduces promastigotes to the warm-blooded host, whilst the infected mosquito introduces third-stage filarial larvae (L3) of *Dirofilaria*. In the mammalian host, promastigotes enter macrophages and inside the cell they replicate as amastigotes, whilst the L3 larvae undergo two moults to become adults who reside in pulmonary arteries and the heart. For 5–10 years the female worms produce microfilariae (L1), which enter into the peripheral blood (McCall et al. 2008). Infected macrophages and free L1 are then taken up by their vectors to complete their life cycles. In areas where hosts, infected by these parasites, and competent insect vectors exist, coinfections may arise (Aresu et al. 2007; Maia et al. 2015). It appears that changes in climate and human activities aid in the geographical spread of these zoonoses and human, as well as dog and feline; cases are increasing in numbers even in areas where the problems were not reported before (Orihel and Eberhard, 1998; Pampiglione and Rivasi, 2000; Dujardin, 2006; Maroli et al. 2008; Tanczos et al. 2012).

Five species of filarial nematodes: *Dirofilaria immitis*, *Dirofilaria repens*, *Dipetalonema reconditum*, *Dipetalonema grassi* and *Acanthocheilonema reconditum* have been recovered from dogs (Genchi et al. 2011); and *Leishmania infantum* (synonym: *Leishmania chagasi* in the New World) is the most common *Leishmania* species causing canine leishmaniasis (CanL), worldwide (Alvar et al. 2012). In Greece, *D. immitis* has been reported in dogs (Polizopoulou et al. 2000; Sinanis et al. 2012) and *D. repens* (*D. conjunctivae*) in humans (Vakalis and Himonas, 1997), whilst *L. infantum* comprises one of the most important parasitic diseases in dogs in most parts of Greece, and an important zoonosis for public health (Antoniou et al. 2009; Christodoulou et al. 2012; Ntais et al. 2013).

The aim of this work was to evaluate the geographical distribution of coinfected dogs in Greece, to isolate the two parasites from the blood of dogs, using a simple method, and to assess their survival and interaction *in vitro*.

**MATERIALS AND METHODS**

**Study area and animal sampling**

The study was conducted in all (54) prefectures of Greece where 63 veterinarians, collaborating with...
our laboratory, provided dog samples from 5772 animals visiting their clinic for any reason: vaccination, haircut, nail cut, deworming, general check-up, treatments and other purposes, without discrimination (Ntais et al. 2013). The animals were examined clinically and biological material was collected, after the written consent of the owner, and a questionnaire with personal, epidemiological and clinical data for each dog was completed. Biological samples, including peripheral blood without anticoagulants from all dogs taking part in the study, were collected and stored in sterile tubes at 4 °C until transfer to the laboratory for processing. The maximum number of samples reached 300 per prefecture and depended on the size of the prefecture and dog population (Ntais et al. 2013).

**Serology**

Dog sera were tested serologically using anti-dog, anti-immunoglobulin G antibodies by an indirect immunofluorescent antibody test (IFAT, Leishmania SPOT IF, BioMerieux, France). A series of 2-fold serum dilutions, starting from 1/40 were performed and a cut-off titre of ≥1/160 was regarded positive for dog sera since most animals lived in endemic areas (Ferroglio et al. 2002).

**Polymerase chain reaction (PCR) and PCR-restriction fragment length polymorphism (PCR-RFLP)**

The PCR for the detection of *Leishmania* DNA was carried out on whole blood, lymph node and/or spleen tissue from dogs, according to availability. The QI Amp DNA Blood Mini kit (QIAGEN, Hilden, Germany) and DNeasy Tissue kit (QIAGEN) were used for DNA extraction from blood and tissue, respectively. Primers T2 and B4 were used as described previously with few modifications (Minodier et al. 1997; Christodoulou et al. 2012). The *Leishmania* ITS1 region followed by a *Hae*III restriction endonuclease digestion of the positive PCR products (Schönian et al. 2003) was amplified in order to identify the parasite species infecting the dogs. All samples were tested in duplicates. Positive controls were used in all PCR assays, which consisted of DNA extracted from *L. infantum* (MCAN/GR/2009/GD70) and *Leishmania tropica* (MCAN/GR/2009/GD52), the two *Leishmania* spp. found in Greece, which had derived from Greek dogs and humans, respectively, and typed by enzyme electrophoresis (Ntais et al. 2013; Karayiannis et al. 2015). Negative controls included samples from healthy dogs born and lived in non-endemic areas in Crete (at >1000 m altitude), that had being tested for *Leishmania* serologically, by PCR and culture and proved negative.

**Parasite culture**

Parasite culture was performed using blood, without anticoagulants, of 1275 animals, which were positive both by IFAT and PCR. Sterile clotted blood (0.5 cm³ obtained from the core of the sample) was suspended in 3 mL RPMI 1640 culture medium containing 25 mM Hepes buffer, supplemented with 2 mM glutamine (GIBCO Invitrogen, Grand Island, NY), 10% heat inactivated fetal bovine serum (FBS – GIBCO Invitrogen), 100 IU mL⁻¹ penicillin, 100 mg mL⁻¹ streptomycin (Roche Diagnostics, Indianapolis, IN), and 5% filtered human urine, and incubated, in culture flasks, at 26 °C (±1 °C) (Howard et al. 1991; WHO, 1991).

**Parasite identification**

*Leishmania* was identified using PCR-RFLP, as described above, and *Dirofilaria* was identified after microscopic examination of the microfilariae using morphometric characters (body size and shape) (Magnis et al. 2013).

**RESULTS**

Of the 5772 randomly selected dogs examined for leishmaniasis by IFAT, and PCR, 1275, (22.09%) were positive both by serology and PCR (Ntais et al. 2013). PCR-RFLP showed the protozoan parasite to be *L. infantum* and morphometric characters of the microfilariae (length, 290–332 µm; width, 6–7 µm; straight body and tail; tapered anterior end) showed the Nematode parasite to be *D. immitis* (Magnis et al. 2013).

Culture, performed using clotted blood of only the 1275 PCR and IFAT-positive animals, resulted in 165 positive for *Leishmania* (12.94%) and 26 positive for *Dirofilaria* (2.04%) culture media. In two of these cultures (both from the island of Corfu, with animal IFAT titres against *Leishmania* 1/640 and 1/5120), both *Leishmania* and *Dirofilaria* emerged (0.16%).

The 26 dogs, from the blood of which *Dirofilaria* L1 was isolated, were 16 males and 10 females; none was a stray animal. Their age was: 6 > 3 years, 12 between 4 and 6 years, 4 between 7 and 9 years, and 4 > 10 years. Their weight was: 4 > 10 kg, 10 between 11 and 20 kg, 10 between 21 and 30 kg, and 2 > 30 kg. All 26 dogs presented antibodies against *Leishmania*: IFAT titres 1/320 to 1/5120 and had symptoms common to both diseases: unusual tiredness, ocular lesions, weight loss. Twenty-four of these cultures contained 5–6 L1 mL⁻¹ live and active *Dirofilaria*, which appeared on the first day and stayed alive until over 30 days at 26 ± 1 °C. From the other two L1 positive cultures, 1 and 2 L1 mL⁻¹ emerged on day 1. In the culture containing I microfilariae mL⁻¹, the microfilariae died on day 4 of the
culture and on day 7, *Leishmania* promastigotes appeared, grew in numbers and propagated successfully. In the second culture containing 2 L1 mL⁻¹, a small number of *Leishmania* promastigotes emerged on day 8, but disappeared after about 1 week, whilst the L1 survived for 30 days.

In the 165 cultures which yielded *Leishmania* promastigotes, the parasites propagated indefinitely in the RPMI medium at 26 ± 2 °C until the culture was terminated after the second passage to be frozen in liquid nitrogen for future work.

The *Dirofilaria* and *Leishmania* coinfected dogs originated from five prefectures (nine dogs from Corfu, seven from Drama, five from Fthiotida, two from Arta, two from Ioannina, one from Rodopi) in which CanL is endemic (Fig. 1).

**DISCUSSION**

Leishmaniasis is recognized as a very important disease for public health in the whole of the Mediterranean Basin, including Greece where it is notifiable. It appears that it has spread geographically nearly in the whole of the country (Antoniou et al. 2009; Ntais et al. 2013) implying that competent sand fly vectors are present and active. The problem of dirofilariasis in Greece, on the other hand, is not considered very important for public health.
health since a few sporadic human cases have being reported which, however, are increasing in number (Vakalis and Himonas, 1997). The presence of microfilaraemic dogs in an area, and possibly of other domestic and wild reservoirs, such as the red foxes (Karayiannis et al. 2015), allows the circulation of the parasite via the mosquito vectors, which explains the escalation of human cases and poses a threat for the future. The spread of both zoonoses is further favoured by the movement of people and pets and the spread of the vectors in space and time due to climatic changes, as well as human activities.

Measures for safeguarding unaffected areas from the introduction of these pathogens (through infected humans or animals) and their vectors must be undertaken. The introduction of Leishmania donovani MON-37 in Cyprus (Antoniou et al. 2008) and the arrival of the aggressive anthropophilic mosquito Aedes albopictus in Europe, which can play the role of the vector for D. immitis and could increase the risk of transmission of this parasite from animals to humans (Cancrini et al. 2003), is an indication of the enhanced movement of pathogens by globalization.

The culture method used is fast, simple and inexpensive and allows the isolation of both parasites from a big number of samples during an epidemiological study for monitoring risk areas. In this study, 12-94% of the dogs studied were positive for Leishmania and 2-04% were positive for Leishmania and Dirofilaria. Coinfections were found in the prefectures of Corfu, Drama, Fthiotida, Arta, Ioannina, Karayiannis, Campino, L. tropica (syn. L. infantum) in red foxes (Vulpes vulpes) in central Greece. (Karayiannis et al. 2013). Morphometric analyses of D. repens species in infections larial worms, including Dirofilaria, have being maintained in vitro (Silverman and Hansen, 1971), as far as we know, is the first record of isolation of live Dirofilaria L1 from the blood of a host and their maintenance for at least 1 month in the RPMI medium at 26°C. The method can also be used for the isolation of Leishmania promastigotes from biological samples (blood without anticoagulants, spleen tissue and lymph node), by placing a small segment of the sterile tissue in the RPMI culture medium. This method can be used to isolate L. infantum and L. tropica from biological samples without the need of a density gradient cell separation medium for the acquisition of lymphocytes to be used for the culture.

It is interesting to note that only in two out of the 26 IFAT and PCR positive for Leishmania samples in which Dirofilaria L1 emerged, Leishmania promastigotes appeared. In one of these two cases, promastigotes proliferated successfully after the three L1 (in the 3 mL culture medium) died, but the weak Leishmania culture which appeared on day 8 of the second culture died after 1 week in the presence of seven L1 in the culture medium. It would be interesting to investigate whether the presence of microfilariae do not favour promastigote proliferation and survival for long in the culture medium and possibly in the host as well. The microfilariae may feed on the promastigotes or produce metabolic substances that do not allow the protozoan parasite to exist. This could allow the development of potential new substances in combating leishmaniasis in dogs and humans.

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