High prevalence and diversity of zoonotic and other intestinal parasites in dogs from Eastern Spain

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Abstract

The diversity and frequency of enteric parasites in dog populations in the Castellón province (Eastern Spain) was assessed by means of a prospective cross-sectional epidemiological survey. A total of 263 canine faecal samples were collected between July 2014 and July 2016. Detection of intestinal parasites was conducted by routine coprological methods. In addition, identification of *Giardia duodenalis* and *Cryptosporidium* spp. was carried out by direct immunofluorescence microscopy, whereas the presence of *Strongyloides* spp. was assessed by real-time PCR in a selected number of specimens. Based on conventional and/or immunofluorescence microscopy examination, 65.8% (95% CI: 59.7%–71.5%) of the investigated dogs were found infected by at least one gastrointestinal parasite. *Giardia duodenalis* (35.4%) and members of the family Ancylostomatidae (27.0%) were the most prevalent protozoan and helminth parasites found, respectively. Other pathogens potentially infective to human included *Toxocara canis* (8.0%), *Cryptosporidium* spp. (6.8%), and *Strongyloides* spp. (1.1%). Frequency of occurrence of helminthic, but not protozoan, enteroparasites was geographical origin-dependent (*P* = 0.02), with dogs living in coastal areas presenting higher infection rates than those living in inland regions. Similarly, rural dogs were significantly more infected than urban dogs (*P* < 0.001). Our results revealed that zoonotic agents were common in dogs from the Castellón province. Animals from rural areas and sheltered dogs were particularly at risk of these infections.

*Keywords:* Intestinal parasites; Dogs; Prevalence; Zoonoses; Spain.
**Introduction**

Intestinal parasites including a wide range of helminth and protozoan species are common infectious agents of dogs. Some of them can be transmitted to humans causing diseases such as hydatidosis by *Echinococcus granulosus* sensu lato (s.l.), giardiosis by *Giardia duodenalis*, and cryptosporidiosis by *Cryptosporidium* spp. (Soriano et al. 2010; Deplazes et al. 2011; Overgaauw and Knapen 2013; Otranto et al. 2017).

Many studies on canine intestinal parasites have been conducted worldwide with heterogeneous results. Dogs, primarily stray and semi-domesticated animals, living in poor-resource settings with favourable environmental features for pathogen transmission harbour greater diversity and higher prevalences (above 80%) of parasitic infections (Dantas-Torres and Otranto 2014). In developed countries, where domestic dogs are generally well-cared and under adequate sanitary conditions, several surveys have revealed rates of intestinal parasitic infections in pet dogs typically ranging from 20% to 30%, although large variations may occur among different dog populations and geographical areas (Dubná et al. 2007; Claerebout et al. 2009; Zanzani et al. 2014). In Spain, the presence of intestinal parasites in canine populations has been investigated in a limited number of epidemiological surveys conducted in the autonomous regions (ARs) of Andalusia (Martínez-Moreno et al. 2007), Aragon (Causapé et al., 1996), the Basque Country (Benito et al. 2003), Cataluña (Gracenea et al., 2009; Ortuño et al. 2014), Madrid (Miró et al., 2007), and Murcia (Martínez-Carrasco et al. 2007) during the period 1992–2014. No relevant reports in the field have been published in the country since then.

The aim of this work was to assess the diversity and frequency of intestinal parasites in both urban and rural dogs from Castellón Province in the AR of Valencia,
an area where the intestinal parasite fauna in the canine host has not been investigated yet.

Material and methods

Study area and design

Castellón is a province of the AR of Valencia in Eastern Spain. It has a surface area of over 6,632 km² and has a total population of 582,327 inhabitants. Most of the population lives in the coastal strip (a third of them in the capital city Castellón de la Plana), whereas the mountainous interior is practically uninhabited. The province is divided in eight administrative regions called “comarcas” (FIG. 1). Agriculture and livestock raising constitute the principal economic activities of the province. The climate of the region is typically Mediterranean, characterized by mild, rainy winters and warm, dry summers.

There were 151,311 domestic dogs officially censed in Castellón in 2015, 21,936 of them belonging to hound-type breeds commonly used in hunting, whereas an undetermined number of guard or shepherd dogs were used in agricultural exploitations in rural areas (Registro Informático Valenciano de Identificación Animal 2015). Overall. Stray, abandoned, or surrendered animals in the province were managed by five private animal shelters, one of them also acting as a licensed breeding kennel. Surveyed animals were categorised as pet dogs, breeding dogs, sheltered dogs, shepherd dogs and hunting dogs. To achieve the objectives of the present project a cross-sectional study was carried out in this Spanish province between July 2014 and July 2016.

Faecal sample collection
Estimated sample size \((n = 217)\) was calculated using Open Source Epidemiological Statistics for Public Health OpenEpi 3.01 software (Dean 2013). Power was set considering an expected prevalence of 50%, a marginal error of 7% with a 95% confidence interval (CI), and a loss rate of 10%. A total of 263 faecal dropping samples from individual dogs were regularly collected during the study period. Faecal specimens belonged to dogs attended at four of the five animal shelters located at the province \((n = 139)\), breeding dogs for sale \((n = 18)\), hunting \((n = 68)\), shepherd \((n = 24)\) and pet \((n = 14)\) dogs. Faecal specimens were placed in screw-topped specimen containers and uniquely labelled indicating identification number and date of collection. Data on sex, age, status, and geographical origin of the dog and consistency of the faecal material were also recorded.

**Parasitological procedures**

Faeces were stored at 4 °C in 5% (v/v) formaldehyde until further treatment. Macroscopic examination was firstly performed for detection of proglottids and adult worms. After homogenization, each faecal sample was divided into two aliquots. In order to detect parasite eggs, cysts and oocysts, one aliquot was analysed using routine coprological procedures based on the modified Ritchie formalin-ether and Sheather's sugar flotation concentration methods (Thienpont et al. 1979). Each sample was microscopically examined at 10× in triplicate and suspected parasite structures confirmed at 40× magnification in a Leica DM500 microscope (Wetzlar, Germany). Parasite eggs, cysts and oocysts were identified according to their morphometric characteristics. The second aliquot was used to assess the presence of *Giardia duodenalis* cysts and *Cryptosporidium* spp. oocysts by direct fluorescent antibody test (DFAT) using a commercially available kit (MERIFLUOR® Cryptosporidium/Giardia,
Meridian Bioscience, EE.UU). A sample was recorded as positive if at least one parasite form was observed by any given method.

**DNA extraction and purification**

The presence of *Strongyloides* spp. was assessed by a PCR-based method (see below) in a limited number of faecal samples from dogs considered at higher risk of being in contact with the nematode, including shepherd and hunting dogs. Aliquots of selected faecal samples were stored in 70% ethanol. Total DNA was extracted from ~200 mg of faecal material using the QIAamp® DNA Stool Mini Kit (Qiagen, Hilden, Germany) following the manufacturer’s instructions. Purified DNA samples (200 μL) were stored at -20 °C and shipped to the Parasitology Reference and Research Laboratory, Spanish National Centre for Microbiology (Majadahonda) for further PCR testing.

**Molecular detection of *Strongyloides* spp.**

Genus-specific (F: 5´‒GAATTCCAAGTAAACGTAAGTCATTAGC‒3´ and R: 5´‒TGCCTCTGGATATTGCTCAGTTC‒3´) primers were used to amplify a partial sequence of the small subunit ribosomal RNA (*ssu* rDNA) gene of *Strongyloides* spp. (Verweij et al. 2009) by a qualitative real-time PCR (qPCR) assay as described elsewhere (Saugar et al. 2015). qPCR reactions (25 μL) contained 1× Quantimix EasyMaster Mix (Biotools B&M Laboratories, Madrid, Spain), 0.2 μM of each specific primer, 0.5 μL of 50× SybrGreen (Invitrogen, San Diego, CA, USA), and 10 μL of total DNA extracted from faecal specimens. Purified genomic DNA from *S. venezuelensis* L3 was used as positive control. All DNA isolates were assayed in duplicate. An internal inhibition control including 10 ng of *S. venezuelensis* DNA was used for each sample. Negative and no template controls were included in each run.
Cycling conditions were 15 min at 95 ºC followed by 50 cycles of 10 s at 95 ºC, 10 s at 60 ºC and 30 s at 72 ºC. DNA amplification and detection of fluorescence at the end of each amplification cycle were performed on a Corbett Rotor Gene™ 6000 real-time PCR system (Qiagen). Data were analysed with Rotor Gene 6000 Series software version 1.7.

**Data analyses**

Infection rates of total and individual intestinal parasites were determined. The Chi-square ($\chi^2$) test was used to compare total and individual parasite infection rates in the canine population under study by sex, status, and geographical origin of the animals. Prevalence risk ratios (PRR) with 95% confidence intervals (CI) were also calculated to assess the association between the above mentioned variables and the occurrence of enteric parasite infections. A probability ($P$) value < 0.05 was considered evidence of statistical significance. Data were analysed with the free software RStudio Version 1.0.44 (https://www.rstudio.com/) using the Epitools library.

**Results**

The study included dogs from six of the eight administrative regions of the Castellón Province (FIG. 1). Intestinal parasites were found in 65.8% (173/263; 95% CI: 59.7%–71.5%) of the faecal samples analysed. Frequencies of appearance of each individual parasite species are shown in Table 1.

Helminth (38.4%; 101/263) and protozoa (43.0%; 113/263) infections were observed in the examined samples with no significant differences ($P > 0.05$). In addition, co-infections involving protozoan and helminth species were detected in 15.6% (41/263) of the samples, whereas no parasite infections were found in 34.2%
of the faecal specimens examined. Overall, seven species of protozoa, three of cestodes, and seven of nematodes were identified. Intestinal protozoa were the most frequent type of enteric parasites identified in all dog groups excepting shepherd and hunting dogs, which were primarily infected by nematodes (Table 1).

The most frequently observed parasite species were *Giardia duodenalis* (35.4%), followed by hookworms (27.0%), *Toxocara canis* (8.0%), and *Trichuris vulpis* (6.8%). However, these figures varied largely depending on the dog's status considered (Table 1). Out of the 93 *Giardia*-positive samples, 90 were detected only by DFAT, whereas the remaining three tested positive both by microscopy examination and DFAT. Additionally, two shepherd dogs carried sporulated *Eimeria* spp. oocysts. Because this coccidia does not naturally infect dogs this finding very likely reflects events of coprophagy or predation of other species. In samples with a positive result to any given intestinal parasite, single and multiple (two or more) infections were identified in 36.5% (96/263) and 29.3% (77/263) of the cases, respectively. Double (*n* = 48), triple (*n* = 21), quadruple (*n* = 4), and quintuple (*n* = 4) infections were recorded. Co-infection by *G. duodenalis* and *Ancylostoma caninum* was the most frequent association identified (9.4%; 33/173). The distribution of single and multiple parasite infections according to the status of the investigated dogs is shown in Table 2. Interestingly, urban (pet and breeding) dogs were significantly less likely (*χ²* = 24.8; *P* < 0.001) to harbour parasitic infections than rural (shepherd and hunting) dogs.

Table 3 shows the assessment of risk factors that may influence the occurrence and transmission patterns of canine intestinal parasites in the present study. In order to increase statistical power, shepherd and hunting dogs were grouped together and categorised as rural animals, whereas pet and breeding dogs were regarded as urban animals. Similarly, and based on their region of origin, dogs were allocated between
two categories: inland (municipalities of Alcalatén, Alto Palancia, and Alto Maestrazgo) and coastal (Bajo Maestrazgo, Plana Alta, and Plana Baja) areas (see also FIG. 1). The male/female ratio was 1.5. Although intestinal parasites were more prevalent in female (69.1%; 65/94) than in male (58.1%; 79/136) dogs, sex was not a risk factor for infection ($P = 0.09$). As expected, rural dogs were at higher risk of harbouring enteroparasites than urban dogs (PRR: 1.51; $P < 0.001$). However, no statistically significant differences in the occurrence of enteric pathogens were demonstrated between dogs living in coastal areas and those living in inner regions of the Castellón province (PRR: 1.24; $P = 0.12$).

Importantly, surveyed dogs were demonstrated to frequently harbour intestinal protozoan and helminth species potentially infective to humans including *G. duodenalis* (35.4%), hookworms (27.0%), *T. canis* (8.0%), *Cryptosporidium* spp. (6.8%), *Dipylidium caninum* (1.9%), members of the family Taeniidae (1.5%), and *Blastocystis* spp. (1.1%). Of note, three dogs (1.1%) were initially found positive for *Strongyloides* spp. at microscopy. Considering that Castellón has been long regarded as an endemic area for *S. stercoralis*, and that human strongyloidiosis cases are still sporadically detected in the province (Martinez-Perez and Lopez-Velez 2015), we further expanded this preliminary finding. Thus, the presence of *Strongyloides* spp. was assessed by molecular methods in selected faecal samples ($n = 87$) from dogs considered at higher risk of being infected by the parasite, including shepherd ($n = 16$) and hunting ($n = 19$) dogs. A number of sheltered dogs ($n = 52$) were also tested based on recently published literature (Paradies et al. 2017). Interestingly, PCR-positive results for *Strongyloides* were obtained in 13.8% (12/87) of the dogs investigated. The infection was more prevalent in shepherd (25.0%, 4/16) than in sheltered (15.4%, 8/52) dogs, but was not
detected in hunting dogs. Neither sex nor origins of the animals were significantly
associated to higher Strongyloides spp. infection rates.

Discussion
This study provides the first description of the diversity and frequency of intestinal
parasites in dogs from the Castellón province. The overall canine infection rate (66%)
recorded here is one of the highest reported in Spain to date, only behind of those
(∼70%) previously identified primarily by microscopy in southern (Martínez-Moreno et
al. 2007) and north-eastern Spain (Ortuño et al. 2014). Additionally, a high diversity of
intestinal parasites was also identified, including seven protozoa, three cestode, and
seven nematode species. Taken together, these data depict an epidemiological scenario
characterised by elevated prevalences leading to high infection and (very likely) re-
infection rates.

In the European context, our results are in agreement with those documented in
Belgium (Claerebout et al. 2009), Czech Republic (Dubná et al. 2007), France (Osman
et al. 2015), Germany (Barutzki and Schaper 2003), Greece (Kostopoulou et al. 2017),
Italy (Zanzani et al. 2014), and Portugal (Mateus et al. 2014). In these surveys G.
duodenalis, A. caninum, and T. canis were demonstrated to be the most common
endoparasite species infecting dogs, although variations in parasite diversity and
frequency rates were often reported among different dog populations and geographical
areas. Of note, the G. duodenalis infection rates observed in the present study (up to
43.2% in sheltered dogs), together with that (43.9%) previously reported in Belgium
also by DFAT (Claerebout et al. 2009) are among the highest documented in Europe to
date. This fact is probably associated to the superior diagnostic sensitivity of DFAT
compared to conventional microscopy, and the high infection pressures and crowded
conditions commonly seen in kennelled dogs (Gil et al. 2017; Adell-Aledón et al. 2018).
Indeed, sheltered dogs harbour the highest parasite diversity (15 species) detected in
the present survey.
Interestingly, shepherd and hunting dogs (both categories linked to rural
activities) were significantly more infected by helminth species than dogs from urban
areas such as pet and breeding dogs. Thus, infections by hookworms (50–76%) and
*T. canis* (7–17%) were particularly abundant among the former dog categories. Similar
prevalence rates have been previously reported in farm and hunting dogs for *A. caninum*
(70%) in neighbour Portugal (Mateus et al. 2014), and for *T. canis* (13%) in Greece
(Papazahariadou et al. 2007). These findings are indicative of failure of dog owners to
comply with prescribed deworming protocols.
Data presented here are also relevant from a public veterinary health perspective.
Among the recovered protozoa, *G. duodenalis* was the most prevalent species.
Importantly, *G. duodenalis* was present in 36% and 28% of the pet and breeding dogs
analysed, respectively. Because of their close contact with their owners, these animals
may act as potential sources of human giardiosis. In this regard, it should be noted that
zoonotic sub-assemblages AII, BIII, and BIV of the parasite have been previously
described in sheltered dogs in northern Spain, although the genotypes found seemed
primarily transmitted within canine cycles and posed therefore limited risk to humans
(Gil et al. 2017). Furthermore, no evidence of zoonotic (or anthroponotic) transmission
of *G. duodenalis* was demonstrated between humans and pet dogs sharing households in
the geographical area (de Lucio et al. 2017). Similar results and conclusions were
reached for the molecular characterization of the *G. duodenalis* samples generated in the
present survey, as described elsewhere (Adell-Aledón et al. 2018). Taken together, all
these molecular data indicate that domestic dogs do not play a relevant role as natural
source of human giardiosis in Spain. Other zoonotic protozoan parasites including
*Blastocystis* spp. and *Cryptosporidium* spp., were found at lower rates.

*Toxocara canis* represents an important public health concern not only in
developing countries but also in industrialized settings with adequate sanitary facilities
(Stolk et al. 2016; Salas-Coronas et al. 2018). Over the last few years, toxocariasis has
gained an increasing international attention and was listed among the five most
neglected parasitic infections according to the US Centers for Disease Control and
Prevention (Chen et al. 2018). Human toxocariasis has been described in more than 100
countries, with Spain ranking first among the European countries reporting cases of the
visceral form of the disease. Humans acquire the infection via contact with soil
contaminated with *Toxocara* eggs. *Toxocara* worms have a tendency to cause extra-
testinal pathologies including four clinical (visceral larva migrans, ocular toxocariasis,
covert toxocariasis, neurotoxocariasis) forms which can lead to serious health
consequences. Due to the non-specific symptoms of this disease, its medical and public
health impact might be underestimated (Chen et al. 2018). The high prevalence of
*Toxocara* in dogs poses also a considerable public health risk as the eggs are
environmentally resistant. Considering that the latest available treatment protocols have
improved the control of the disease (Rehbein et al. 2017), it should be emphasized that
regular pet deworming would be a useful tool to reduce this problem.

Also noteworthy was the finding of taenid eggs in faecal specimens belonging to
sheltered and hunting dogs. The family Taeniidae comprises cestodes of the genus
*Taenia* and *Echinococcus*, important (and neglected) zoonotic helminths of dogs whose
eggs are morphologically indistinguishable at microscopy examination. Although we
did not conduct any molecular test for the specific detection of *E. granulosus* s.l. (the
causal agent of human CE or hydatid disease), the possibility that some of the
investigated dogs were naturally infected by this cestode cannot be completely ruled out. Indeed, an *E. granulosus* infection rate of 0.5% (5/1,040) by necropsy has been previously described in sheltered dogs in Northern Spain (Benito et al. 2003). Therefore, more studies are required to investigate the current epidemiological situation of canine equinococciosis in this geographical area. One of the most intriguing contributions of this paper was the detection of *Strongyloides* spp. in a significant number of shepherd and sheltered (but not hunting) dogs. Members of the family Canidae and Felidae are considered suitable hosts for a number of *Strongyloides* species including *S. stercoralis*, the etiological agent of human strongyloidiosis (Thamsborg et al. 2017). Whether domestic dogs can act as suitable reservoirs of human infections remains a matter of intense debate, but a recent molecular survey conducted in rural Cambodia has demonstrated that humans and their dogs can be infected by the same genetic variant of *S. stercoralis* (Jaleta et al. 2017). Arguing in favour of the occurrence of zoonotic transmission, the authors suggested that in order to reduce the exposure of humans to infective *S. stercoralis* larvae, dogs should be treated against the infection along with their owner. In Europe there are few studies on the prevalence of this parasite in dogs. The infected animals were usually asymptomatic and when signs and symptoms appeared they were unspecific. However, the increase of human strongyloidiasis cases diagnosed globally has lead the scientific community to reconsider the role of domestic dogs as potential natural reservoirs of human infections (Paradies et al. 2017). Imported human strongyloidiasis associated to immigrant populations and returning travellers from endemic areas is increasingly reported in Spain (Martinez-Perez et al. 2018, Belhassen-García et al. 2017), although in Castellón Province sporadic autochthonous cases of the disease are still recorded. The fact that these cases correspond to individuals of older age has been interpreted as evidence of
successful interruption of the transmission cycle of the parasite (Martinez-Perez and Lopez-Velez, 2015). Still, it would be very interesting to isolate *Strongyloides* larvae from fresh faecal material of canine origin in order to identify the species involved and assess the associated zoonotic risk.

**Conclusions**

This is the first coprological, microscopy-based study targeting different dog populations conducted to date in the Castellón Province. Investigated dogs were infected at high rates by a wide range of protozoa and helminth species, some of them with zoonotic potential. Dogs from rural areas (mainly shepherd and hunting dogs) were more exposed. Simple measures, such as periodic deworming, prompt removal of faeces from kennels, and improving owner's education on zoonotic transmission are all cost-effective methods to limit the risk of animal and human infections by enteric parasites. People at higher risk of infection (e.g. veterinarians, slaughterhouse workers, animal husbandry workers, kennel personnel, and hunters) should be provided with accurate information on the potential risks associated to dog handling and management. Finally, data provided here are expected to be of interest for public veterinary health authorities and decision makers in order to design and implement effective control measures against these infections.

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Author Disclosure Statement

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and cats from metropolitan and micropolitan areas: prevalence, zoonotic risks, and

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**Figure legends**

**FIG.1.** Map of the administrative divisions of the Castellón province. The
municipalities where sampling was conducted and the status of the dog sub-populations
are indicated. The location of Castellón in Spain is highlighted in red in the upper left corner. Image reproduced with permission of BioMed Central.