Molecular identification of *Giardia duodenalis* isolates from domestic dogs and cats in Wroclaw, Poland

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Piekarska J, Bajzert J, Gorczykowski M, Kantyka M, Podkowik M. Molecular identification of *Giardia duodenalis* isolates from domestic dogs and cats in the area of Wroclaw (Poland).

**Abstract**

**Introduction.** *Giardia duodenalis* (*G. intestinalis*) is a common protozoan causing gastrointestinal disorders in many species of mammals. The genus of *Giardia* has high molecular diversity. Dogs and cats, in addition to their typical infection with assemblages C, D and F, may be a reservoir of zoonotic assemblages (A and B).

**Objective.** The aim of this study was a genetic characteristic of *Giardia* isolates of dogs and cats from the area of Wroclaw (Poland).

**Materials and method.** A total of 128 and 33 faecal samples from dogs and cats, respectively, were analyzed by routine coprological methods. The animals were diagnosed on the presence of *G. duodenalis* antigens in faeces soluble with the use of SNAP *Giardia* (IDEXX Laboratories) immunosorbent assay. 27 DNA isolates of *Giardia* were subjected to molecular identification (PCR-RFLP).

**Results and conclusions.** The prevalence of *G. duodenalis* was 21.1% (27/128) in dogs and 15.1% (5/33) in cats. In dogs, C assemblage was present in 18 (81%) positive stool samples, D assemblage in 2 (9%) samples, B assemblage present in one (4.5%), and mixed assemblages (C and D) occurred in one (4.5%) sample. F assemblage was found in 4 (80%) cats’ positive stool samples and A assemblage occurred in one case (20%). Confirmation of the presence of A and B zoonotic assemblages suggests that infected pets can be a threat to human health. This study describes for the first time the presence of mixed infections within host-specific C and D assemblages in dogs in Poland.

**Key words**

*Giardia duodenalis*, assemblage, dogs, cats, nested-PCR, PCR-RFLP, zoonosis

**INTRODUCTION**

*Giardia duodenalis* (*G. intestinalis, G. lamblia*) is a widespread protozoan parasitizing in humans and many species of mammals. Invasion is most commonly associated with the occurrence of gastrointestinal signs, though asymptomatic infections have also been observed [1]. Studies using molecular techniques have shown that genetic diversity is very high within the *G. duodenalis* species. There are 7 basic assemblages, from A – G. The occurrence of A, B, C and D assemblages was confirmed in dogs, and A, B, D and F assemblages found in cats. Due to the occurrence of A and B assemblages in humans, and because of their relevance for zoonotic infections, there is a need for the monitoring of dogs and cats as companion animals that can be a direct source of human infection, as well as a source of environmental contamination [2, 3]. Asymptomatic and chronic course of the disease with frequent periodic expulsion of cysts is observed in older animals. Diagnosis of *Giardia* infection with the use of a faecal flotation or faecal smear is problematic due to the irregular shedding of cysts and their morphology, small size, and their similarity in appearance to many pseudoparasites such as yeast [4]. The use of immunoenzym assay to detect *Giardia* coproantigens in faeces increases the probability of detecting invasion. Genotypic characterization of *G. duodenalis* is a very useful and essential tool used in epidemiological studies. PCR techniques for genotyping of *G. duodenalis* are based on polymorphic genes encoding 18S rRNA, glutamate dehydrogenase (gdh), triose phosphate isomerase (tpi), and ß-giardin [5, 6]. PCR amplification and RFLP/sequence analysis of all of these genes, with the exception of the 18S rRNA, can differentiate subgenotypes of assemblage A [7]. PCR methods for detection of gdh can provide information on *G. duodenalis* A and B sub-assemblages [8].

**OBJECTIVE**

The aim of the study was the genetic characterization of isolates of *Giardia* in dogs and cats in the area of Wroclaw (Poland), which is important in determining the source of
the invasion in humans and animals in a given area, as well as in assessment of the zoonotic potential of *G. duodenalis*.

**MATERIALS AND METHOD**

**Samples collection and qualification.** Faecal samples of dogs and cats (age: 3 weeks – 10 years) from Wroclaw were collected in 2010–2012 and provided by their owners to the Division of Parasitology, Faculty of Veterinary Medicine (Wroclaw). The total number of stool samples was 161 (128 dogs and 33 cats), all of the animals had different gastric symptoms (e.g. diarrhea, emaciation, loss of body weight). Faecal samples were stored at -20°C. The animals were diagnosed on the presence of *G. duodenalis* antigens in faeces soluble with the use of SNAP Giardia (IDEXX Laboratories) immunosorbent assay. As a result of such qualifications, material for genetic research was obtained from 32 animals (27 dogs and 5 cats). Pets with positive test results ranged in age from 3 weeks – 2 years.

**DNA isolation.** Faecal samples (approximately 3–4 g) from infected animals were examined by the flotation method with the use of saturated NaCl solution. The upper part of the supernatant was harvested, rinsed in distilled H₂O, centrifuged at 1,000 rpm for 5 minutes. The resulting pellet was re-suspended in 300 µl of saline. This was a concentrated sample of *Giardia* cysts, which were the basis for DNA isolation using a Genomic Mini AX STOOL kit (DNA-Gdansk, Poland).

**Genotyping and nested PCR.** *Giardia* genotyping was performed based on the polymorphism of a gene fragment coding for β-giardin. Nested PCR technique for DNA amplification was applied. Reaction parameters and primers sequences were described in Lalle et al. (2005) [9]. In the primary PCR reaction forward G7: 5’AAGCCCGAC-GACCTCACCCGCAGTGC3' and reverse G759: 5’GAGGCCCAGCCTGTAGCTTCGAGCAC3' primers were used. The reaction mixture consisted of 400 nM of each primer, 1 x reaction buffer for polymerase DNA Delta3 (DNA-Gdansk, Poland) and 0.5 µl of DNA in a final volume of 25 µl. The amplification was carried out in a MJ Mini thermal cycler (Bio-Rad) using the polymerase DNA Delta3 (DNA-Gdansk, Poland) and 0.5 µl of DNA in a final volume of 25 µl. The amplification was carried out in a MJ Mini thermal cycler (Bio-Rad) using the following conditions: cycle of 96°C for 5 min; 5 cycles of: initially denaturation for 30 sec at 95°C, annealing for 30 sec at 55°C, polymerization for 1 min at 72°C. The PCR cycle was then carried out for 30 sec at 95°C, 30 sec at 65°C and 1 min at 72°C, for a total of 30 cycles, followed by a final extension for 10 min at 72°C. The secondary PCR reaction was amplified with forward 511: 5’ GAAAGCAAGACCGATCGAGGTCCG3’ and reverse 511: 5’CTCGACGAGCTTCGTTGTT3’ primers. The amplification conditions were almost the same as in the first reaction with only 2 differences. The annealing temperature was changed, the first was 50°C and the second was 55°C and no final extension was carried out. The amplicons of the first PCR reaction (753 bp) was the matrix of secondary PCR reaction.

Amplicons of second PCR reaction (total volume = 100 µl) were subjected to electrophoresis in 2% agar gel. PCR products with a length of ~511 bp were isolated from the gel and purified using a set of DNA Extraction Kit (Fermentas). DNA concentration was determined by absorbance measurements at 260 nm wavelength. Purified products were digested with restriction enzyme BsuRI/HaeIII (Fermentas) for 2 hours at 37°C. The required enzyme amount was calculated based on its activity on the lambda phage. Digestion products were electrophoresed in a 5% agarose gel for 2h at a constant 120V. Products were stained with ethidium bromide and photographed with a BioRad GelDoxXR device.

**Identification of genotypes of sequenced DNA.** Selected products of PCR reaction of approximately 511bp were subjected to DNA sequencing by GENOMED S.A. (Poland). The obtained results of sequencing were analyzed by comparing the sequence of individual genotypes of *G. duodenalis* with BLAST NCBI (http://blast.ncbi.nlm.nih.gov) database. The partial sequences of β- giardin gene obtained in this study were compared with sequences deposited in the GenBank database under Accession Nos.: Assemblage A- FJ560591.1; Assemblage B- AY072725.1; Assemblage C- AY545646; Assemblage D- AY545647.1; Assemblage F- AY647264. BioEdit and APE A Plasmid Editor software tools were used for verification of sequence compliance.

**RESULTS**

In the study by nested PCR, *Giardia* DNA was detected in 27 of 32 tested stool samples (22 dogs and 5 cats). In 4 cats, the presence of assemblage F was revealed and in 1 cat it was assemblage A. Of the 22 dogs’ stool samples, assemblage C was detected in 18 cases, assemblage D in 2 cases, assemblage B in 1 case.

Selected products of nested PCR reaction (~511bp) were sequenced and analyzed using the GenBank database. Analysis of the sequence results in cases where one assemblage was identified clearly confirmed the results obtained from RFLP. Figure 1 shows selected results of sequencing which, in turn, helped to identify the assemblages: A (Fig. 1A), B (Fig. 1B), C (Fig. 1C), D (Fig. 1D), F (Fig. 1E). There were no specific products for any of the previously identified assemblages in electrophoresis of 1 sample derived from a dog after digestion with HaeIII enzyme. The sense strand sequencing results of a β-giardin gene indicated the presence of assemblage C (Fig. 2A). However, sequencing of the antisense strand showed the presence of assemblage C or D, giving an appropriate sequence homology at the level of 0.988 and 0.949. In the analyzed antisense strand stool sample, the cut places were present at 194, 296, 311 bp, while characteristic cut places for assemblage C were present at 194, 296, 311 and 461bp (Fig. 2B) and for assemblage D only in 194 and 311bp (Fig. 2C). Therefore, it is presumed that in this stool sample from dog we are dealing with a mixed infection of assemblages C and D, this observation was confirmed by the results of electrophoretic analysis of nested PCR product digested by BsuRI enzyme (data not shown).

In 5 dogs, despite the positive test of SNAP Giardia (IDEXX Laboratories), the PCR product was not detected. The nested PCR repeated one more time with the matrix volume increased to 2–4 µl in 25µl of reaction mixture and with the revised polymerase, did not produce satisfactory results. Re-isolation of DNA from thawed samples also did not allow DNA product to be obtained.
Figure 1 A–E. Multiple alignment of the *G. duodenalis* assemblage of β-giardin genes in nested PCR. Alignment based on maximum similarity of individual assemblages of *G. duodenalis* in the classification of Lalle et al. 2005. The underlined zone indicates a cut place of restrictive enzyme. The grey zone indicates differences in DNA sequence (no consensus at this position).
A) The sense strand comparison with assemblage C. Accession number AYS545646

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B) The antisense strand comparison with assemblage C. Accession number AYS545646

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C) The antisense strand comparison with assemblage D. Accession number AYS545647.1

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Figure 2 A–C. Comparison of G. duodenalis sequencing product from a dog with the pattern of assemblages C and D. Alignment based on maximum similarity of individual assemblages of G. duodenalis in the classification of Lalle et al. 2005. The underlined zone indicates a cut place of restrictive enzyme. The grey zone indicates differences in DNA sequence (no consensus at this position).
DISCUSSION

*Giardia duodenalis* occurs in dogs and cats all over the world and is one of the major parasites responsible for the symptoms of the gastrointestinal tract. Extensiveness of the invasion is varied and depends on the geographical location, different hygiene conditions in the region, as well as different diagnostic methods usage. Because the diagnosis of *Giardia* invasion based on classical methods (faecal flotation or faecal smear) is problematic, commercial immunoassays are therefore increasingly used to detect the presence of parasite protein secretion in animal faeces. Giardiasis is a very common disease and invasion prevalence ranges from 5–80%, depending on the age of the animals. The highest prevalence (46–50%) is observed in young dogs less than one year old, and in animals with diarrhea [10, 11]. In Poland in 2001–2006, the prevalence of *Giardia* invasion in dogs in Poznan was about 10%, in Warsaw it ranged from 9% to over 50%, in Lublin about 53%, in Pulawy – 10% and GDansk over 16%. Significant differences in prevalence values resulted from the different number of animals in age groups and differences of their clinical condition [12]. In the presented study, the prevalence of *G. duodenalis* in dogs was 21.1% and in cats – 15.1%. The use of enzyme immunoassay test for detecting soluble *Giardia* antigens in faeces increases the probability of invasion detection. Studies conducted in Europe (especially in countries such as the UK, Spain, Netherlands, Italy, Germany, Belgium) using the SNAP *Giardia* Test (IDEXX Laboratories) showed the presence of *G. duodenalis* invasion in approximately 25% of dogs and 20% of cats with gastrointestinal signs [11]. In the USA, the prevalence of *G. duodenalis* invasion in animals with symptoms of the gastrointestinal tract was 15.6% in dogs and 10.8% in cats [13]. Similar results were obtained in Canada, where the prevalence was 13% in dogs and 4.1% in cats, and in Japan, where the prevalence of *Giardia* invasion in 1997–2007 in pet dogs remained unchanged at the level of about 15% [14]. In Brazil, the percentage of infected dogs was 16.9% [15]. Usually, for the molecular identification of *G. duodenalis*, PCR and its modifications (nested-PCR, semi-nested PCR, PCR-RFLP, real-time PCR) or *Giardia* DNA hybridization with a molecular probes (microarray technique and FISH technique) are used. Analysis of the genetic material is helpful in answering the question whether *Giardia* detected in a dog or cat can be a source of infection for humans. The most preferred is amplification of the β-*Giardin* encoding gene [9]. *Giardins* are specific structural proteins with a mass of 29–38 kDa, unique for this protozoan [16]. The integral part of each trophozoit sucker are α- and β-*giardins*, while on the surface of *G. duodenalis* cell membrane, a further 10 specific proteins are identified. Genetic research conducted in Poland revealed in affected dogs an occurrence of assemblages A-1, C and D in Warsaw, and only C and D assemblages in Poznan [17, 18]. Molecular analysis of isolates obtained from cats in Warsaw showed the presence of assemblages A, B and D [19]. Current studies in animals in Wroclaw confirmed the presence of specific assemblages C, D, mixed assemblages of C and D, and zoonotic assemblage B in stool samples derived from dogs. In cats, the presence of assemblage F was revealed, but the zoonotic assemblage A was also detected. The study of 55 dogs in Germany, with no apparent clinical signs of disease, revealed the presence of genotype A in 60% of tested animals, mixed infections with assemblages A and C in 27.3%, while the individual assemblages C and D were rarely recorded, respectively, in 9.1% and 3.6% of dogs.

In dogs in Spain, the most commonly circulating assemblages were B, D, followed by A, E and F [20]. The study not only confirmed the high prevalence of *G. duodenalis* among dogs with no clinical signs of disease, but also showed that zoonotic assemblage A is common in urban pet dogs, and even more common than assemblages typical for dogs. Although dogs owners’ stools samples were not studied, the results indicate that a large proportion of urban dogs infected with zoonotic genotypes of *Giardia* are a reservoir for human invasion [21].

As in the presented study, the results of genotyping of *Giardia* isolates present in dogs in different regions of the world indicate that in these animals specific assemblages C and D usually predominate. Such results were obtained in dogs in Hungary, Brazil and Australia [22, 15, 23]. Similarly, non-zoonotic assemblages were identified more often in dogs in Italy (C and D), and assemblage A or mixed-induced invasion with zoonotic and non-zoonotic assemblages occurred only in individual animals [9]. Also in Australia, zoonotic assemblages A and B have been found only in individual animals, which indicated that dogs and cats are not a significant reservoir of invasive isolates of *Giardia* for humans [23]. In turn, other researchers from Germany, Japan and Thailand detected zoonotic assemblage A of *G. duodenalis* in dogs more often than assemblages C and D that are specific for canines [24, 25]. Higher prevalence of zoonotic *Giardia* assemblages were found in dogs that were kept singly than in dogs kept in groups [22]. This confirms earlier suggestions that individual dogs bred in house are infected by *Giardia* from the owners or householder, while the dogs kept in high density are infected with parasite assemblages specific for canids.

Although most of the invasion found in cats is caused by non-pathogenic to humans assemblages D and F, it has been shown that these animals are also a potential source of zoonotic *G. duodenalis* assemblages [5, 9, 22, 23]. Research carried out in Poland on cats from Warsaw confirmed the presence of assemblages A and B, next to assemblages D specific for cats [19]. Also, the results of the presented study, in which apart from assemblage F specific for cats, the zoonotic assemblage A was also found, indicating a potential threat to human health.

The issue that in 5 positive samples tested with the SNAP *Giardia* Test (IDEXX Laboratories) nested PCR failed can be explained by insufficient numbers of parasite’s DNA for amplification [26].

CONCLUSIONS

The results confirm the presence of assemblages C and D in dogs, and assemblage F in cats in the vicinity of Wroclaw. Stating the presence of assemblage A and B suggests that infected pets can pose a threat to human health. This study describes for the first time the presence of mixed infections within host-specific C and D genotypes in dogs in Poland.

Acknowledgements

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REFERENCES