Molecular Characterization of New Haplotype of Genus *Sarcocystis* in Seabirds from Magdalena Island, Southern Chile

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**Simple Summary:** Sarcocystidae is a family of apicomplexan protozoa highly prevalent in vertebrates. The definitive hosts of sarcocystids eliminate oocysts or sporocysts that infect intermediate hosts. After infection, mature tissue cysts (sarcocysts) develop in intermediate hosts, mostly in muscle and neurological tissues. *Sarcocystis* are infectious for definitive hosts, which acquire them through carnivorous or scavenging habits. Intermediate hosts and definitive hosts are the natural hosts of sarcocystids in which infections are usually mildly or not asymptomatic. In 2017, muscular and neurological tissues of 22 birds from Magdalena Islands, southern coast of Chile, were screened for the presence of DNA of sarcocystids. DNA of organisms of the genus *Sarcocystis* was identified in two Chilean skuas (*Stercorarius chilensis*). The genetic makeup of the parasite detected in skuas was unprecedented and probably represent a new species in the genus. It is well known that *Sarcocystis* may cause severe infections in aberrant hosts, which are susceptible animals that do not behave as natural hosts for the parasite and have low resistance to the infection, thus more studies are needed to characterize this parasitosis in skuas and other hosts to understand the epidemiology of the infection and its impact on the health of marine fauna.

**Abstract:** Evidence of sarcocystid infection was investigated in samples of 16 penguins (*Spheniscus magellanicus*), four Dominican gulls (*Larus dominicanus*) and two Chilean skuas (*Stercorarius chilensis*) found in Madalenas Islands, Chile, in 2017. Samples of skeletal muscle, cardiac muscle and brain from all birds were screened by a pan-sarcocystid nested-PCR targeting a short fragment of the gene encoding the small ribosomal unit (nPCR-18Sa). The only two positive samples by nPCR-18Sa, both from skuas, were tested by a nested-PCR directed to the internal transcribed spacer 1 (nPCR-ITS1), also a pan-sarcocystidae nested-PCR, and to a nested-PCR directed to the B1 gene (nPCR-B1), for the exclusive detection of *Toxoplasma gondii*. The two nPCR-18Sa-positive samples were nPCR-ITS1-positive and nPCR-B1-negative. The nPCR-ITS1 nucleotide sequences from the two skuas, which were identical to each other, were revealed closely related to homologous sequences of *Sarcocystis halieti*, species found in seabirds of northern hemisphere. Larger fragments of genes encoding 18S and partial sequences of genes coding for cytochrome oxidase subunit 1 were also analyzed, corroborating ITS1 data. The haplotypes found in the skuas are unprecedented and closely related to species that use birds as the definitive host. Further studies need to be carried out to detect, identify and isolate this parasite to understand the epidemiology of the infection and its impact on the health of marine fauna.

**Keywords:** wild birds; coccidian; molecular; apicomplexa; marine
1. Introduction

The phylum Apicomplexa is composed of obligate intracellular parasites that are characterized by having a specialized structure called the apical complex, which is used to invade vertebrate host cells [1]. Within this phylum, the Sarcocystidae family comprises more than 196 species of coccidia that form cysts in tissues of intermediate hosts. Although taxonomic controversies still exist, this family has been divided into three subfamilies: Sarcocystinae, represented by the genera Frenkelia and Sarcocystis; Cystoisosporinae, containing the genus Cystoisospora; and Toxoplasmatinae, a subfamily with a few species grouped in the genera Besnoitia, Hammondia, Neospora and Toxoplasma [2–5].

Toxoplasma gondii is a coccidian parasite with worldwide distribution. It infects virtually all warm-blooded animals, including humans, but only cats (domestic and wild) act as definitive hosts. Toxoplasmosis has been reported in many avian species; however, little information is available in relation to populations of Spheniscus magellanicus, Stercorarius chilensis and Larus dominicanus [6]. Recently, T. gondii antibodies were detected in 57 (43.18%) out of 132 serum samples collected from free-living Magellanic penguins (Spheniscus magellanicus) on Magdalena Island, Chile, with titers that ranged from 20 to 320 [7].

The genus Sarcocystis has an obligate two-host life cycle. Asexual stages develop in intermediate hosts, usually omnivores, through forming cysts in the musculature and central nervous system. Infection of intermediate hosts occurs through their ingestion of food or water contaminated with sporocysts. Sexual stages only develop in the definitive host, which is typically a carnivore or an omnivore, and infection in this case occurs through ingestion of meat contaminated with cysts [8]. Sarcocystids of the genus Sarcocystis may cause severe infections in aberrant hosts, which are susceptible animals that do not behave as natural hosts of the parasite and have low resistance to the infection. Thus, Sarcocystis potentially pose risk to human and animal health, depending on the susceptible host behaving as aberrant host or not [8].

A few studies have documented the presence of Sarcocystis spp. in wild animals in Chile. Presence of cysts of this parasite has been confirmed in muscle tissues of pudu deer (Pudu puda), guanacos (Lama guanicoe) and sea lions (Otaria byronia) [9–11]. However, Sarcocystis has not yet been described in Chilean wild birds.

The Chilean skua Stercorarius chilensis is a large predatory seabird that inhabits shore ecosystems along the southern cone of South America from central Peru to northern Argentina, with occasional occurrence on the coasts of Ecuador, Brazil, Uruguay and Antarctica [12]. Skuas belong to the order Charadriiformes and are considered to be opportunistic feeders, preying on a wide diversity of animals such as small seabirds, fish, scraps and carrion [13,14]. Populations of skuas may be small, but they do not approach the thresholds for vulnerable classification following a population-size criterion (<10,000 mature individuals) [15].

Considering other coastal birds’ species, the Kelp Gull (Larus dominicanus) is an opportunistic feeder like numerous Laridae and consumes a wide variety of fishes, invertebrates and fisheries waste [16,17]. A high diversity in the use of habitat types has been recorded throughout its distributional range in the Southern Hemisphere, including Argentina, Brazil, Chile, Peru and Uruguay, and the breeding population has been estimated at least 160,000 pairs [17]. In contrast, the Magellanic penguin (S. magellanicus) has approximately 1.1 to 1.6 million breeding pairs that nest along the eastern and western coasts of South America, in Argentina Chile and the Malvinas/Falkland Islands [18]. Spheniscus magellanicus has a primarily piscivorous diet with the presence of some cephalopods and crustaceans [19].

To date, more than 25 species of Sarcocystis have been found to use birds as intermediate hosts [8,20]. Sarcocystis falcataula, Sarcocystis calchasi and the recently described unnamed species Sarcocystis sp. Chicken-2016-DF-BR, which can possibly be interpreted as Sarcocystis wenzeli [21] are species that may be pathogenic for intermediate hosts [22–24].

Focusing on T. gondii and the genus Sarcocystis, the aim of this study was to screen for the evidence of new species or species genotypes of Sarcocystidae in seabird carcasses from southern Chile, a region with scarce data on the occurrence of this group of parasites.
Molecular evidence of a unique haplotype of genus *Sarcocystis* was found in two Chilean skuas (*S. chilensis*).

2. Materials and Methods

2.1. Ethical Considerations

Sample collections on Magdalena Island were conducted under license no. 039/2016 issued by the National Forestry Corporation (Corporación Nacional Forestal; CONAF), and permit no. 2799 issued by the National Fisheries Service (Servicio Nacional de Pesca; SERNAPESCA), Chile. This study was approved by the Ethics Committee on Animal Use (CEUA-no. 9701041113) of the School of Veterinary Medicine, University of São Paulo (FMVZ-USP).

2.2. Collection of Samples

In January 2017, fragments from the pectoral muscle, heart and brain, comprising approximately 5–10 g each, were collected from fresh seabird carcasses on Magdalena Island. This island is located in the Strait of Magellan, near the city of Punta Arenas, in southern Chile (52°55′10.0″ S; 70°34′37.7″ W), and constitutes a natural reserve named “Monumento Natural Los Pinguinos”. Necropsies were performed in situ and samples were stored in sterile microtubes at −20 °C until the time of analysis. Samples were collected from 22 birds: 16 penguins (*S. magellanicus*), four Dominican gulls (*L. dominicanus*) and two Chilean skuas (*S. chilensis*), totaling 66 samples (22 from pectoral muscles, 22 from hearts and 22 from brains).

2.3. Molecular Identification

Initial screening targeting the Sarcocystidae family was performed using a pan-sarcocystid nested PCR based on primers [28] directed to a short fragment of 18S rDNA gene (nPCR-18Sa). The nPCR-18Sa positive samples were further investigated for the presence of DNA of *T. gondii* to amplify partial fragments of gene B1 (nPCR-B1) using the primers described by [29]. The nPCR-18Sa positive samples were also tested by a second pan-sarcocystid nested PCR directed to internal transcribed spacer 1 (nPCR-ITS1) [30,31]. The nPCR-ITS1 were used in order to obtain genetic sequences capable of differentiating the species of the Sarcocystid screened with nPCR-18Sa. The nPCR-ITS1 positive samples were further tested with a third pan-sarcocystid nested PCR, now targeting a larger fragment of 18S rDNA gene (nPCR-18Sb) using primers described by [32], as well with a *Sarcocystis* specific nested PCR [30] directed to cytochrome oxidase subunit I (nPCR-CO1). The primers are depicted in Table 1.

The first round of nPCR-18Sa were performed with 3.0 μL of extracted DNA, 1.8 μL of 10× PCR Buffer (KCl 50 mM; Tris HCl 10 mM; pH 9.0) (Life Technologies Corporation, Carlsbad, CA 92008 USA), 0.7 μL of MgCl₂ (50 mM), 1.4 μL of mixed dNTPs (10 mM), 0.1 μL of each primer (25 μM), 0.14 μL of Platinum™ Taq DNA Polymerase (5 U/μL) (Life Technologies Corporation, Carlsbad, CA 92008 USA) and ultrapure autoclaved water to a volume of 18 μL per reaction. The PCR thermal cycling consisted of an initial incubation at 94 °C for 30 sec, followed by 30 cycles (94 °C for 25 sec, 55 °C for 1 min, 72 °C for 1.5 min) and a final extension at 72 °C for 10 min. For the second rounds: 1 μL of template derived from the first reactions, 2.5 μL of 10× PCR Buffer (KCl 50 mM; Tris HCl 10 mM; pH 9.0) (Life Technologies Corporation, Carlsbad, CA 92008 USA), 2.5 μL of MgCl₂ (50 mM), 4.0 μL of mixed dNTPs (10 mM), 1.25 μL of each primer (10μM), 0.15 μL of Platinum™ Taq DNA Polymerase (5 U/μL) (Life Technologies Corporation, Carlsbad, CA 92008 USA) and
ultrapure autoclaved water to a volume of 25 µL per reaction. The PCR thermal cycling consisted of an initial incubation at 94 °C for 4 min, followed by 30 cycles (94 °C for 30 s, 55 °C for 1 min, 72 °C for 2.0 min) and a final extension at 72 °C for 10 min. For the second rounds the same quantities of the reagent mixture with primers at 50 µM, using 2 µL of the product of the PCR diluted in ultra-pure water (1:2). The nPCR thermal cycling consisted of an initial incubation at 94 °C for 4 min, followed by 35 cycles (94 °C for 30 s, 55 °C for 1 min, 72 °C for 2.0 min) and a final extension at 72 °C for 10 min.

Table 1. Primers for the detection of Sarcocystidae in tissues of seabirds from Magdalena Island, Chile.

| PCR     | Primers       | Sequences                        | PCR Step a | Reference |
|---------|---------------|----------------------------------|------------|-----------|
| nPCR-18S| Tg18s48F      | CCATGCATGTCTAAAGTATAAGC          | 1          | [28]      |
|         | Tg18s359R     | GTTACCCGTAACGGCCCAC              | 1          | [28]      |
|         | Tg18s58F      | CTAAGTATAAGCTTTTTATACCCGC       | 2          | [28]      |
|         | Tg18s348R     | TGGCCAGGTTAATCCATAC             | 2          | [28]      |
| nPCR-B1 | T1            | AGCCCTCTCTCTCAAGAAGCCGTAA       | 1          | [29]      |
|         | T2            | TCCGGAGGACTCTTTATCTAGTG         | 1          | [29]      |
|         | T3            | TGGGAATGAAAGAGAGGCTAATAGG       | 2          | [29]      |
|         | T4            | TTAAGCGCTCTTTGCTGAAAACTATCG     | 2          | [29]      |
| nPCR-18Sb| 18s9L        | GGATAACCTGGTAAATCTATG           | 1 + 2      | [32]      |
|         | 18s1H         | GGCAAATGCTTTCGCGACTAG           | 1 + 2      | [32]      |
| nPCR-CO1| COX1-227F25   | GTTTTGGTAAACTACTTGTACGGAT       | 1          | [31]      |
|         | COX1-885R25   | GAAATATGACGGATATCTCTCTCTT       | 1          | [31]      |
|         | COX1-275F22   | GTACCGAGAATTAAGGTAG            | 2          | [31]      |
|         | COX1-844R21   | GTGGTCCCCATACAGAAGGA            | 2          | [31]      |
| nPCR-ITS1| JS4           | CGAAATGGGAAGTTTGGAAC            | 1          | [33]      |
|         | CT2c          | CTGCAATTCACATTCGCC             | 1          | [30]      |
|         | JS4b          | AGTCGTAAACAGGTTCCGTTAGG        | 2          | [30]      |
|         | CT2b          | TTGGCGAGGCAAAGACATC            | 2          | [30]      |

*Primers used in the first round of amplification (1); primers used in the second round of amplification (2); primers used in both first and second round of amplification (1 + 2).

The first round of nPCR-B1 were performed with 1.0 µL of extracted DNA, 2.5 µL of 10× PCR Buffer (KCL 50 mM; Tris HCl 10 mM; pH 9.0), (Life Technologies Corporation, Carlsbad, CA 92008 USA), 0.75 µL of MgCl₂ (50 mM), 4.0 µL of mixed dNTPs (10 mM), 1.25 µL of each primer (10 µM), 0.15 µL of Platinum™ Taq DNA Polymerase (5 U/µL) (Life Technologies Corporation, Carlsbad, CA 92008 USA) and ultrapure autoclaved water to a volume of 25 µL per reaction. The PCR thermal cycling consisted of an initial incubation at 94 °C for 3 min, followed by 25 cycles (94 °C for 25 s, 55 °C for 1 min, 72 °C for 1.5 min) and a final extension at 72 °C for 10 min. For the second rounds: 1 µL of template derived from the first reactions, 2.5 µL of 10× PCR Buffer (KCL 50 mM; Tris HCl 10 mM; pH 9.0) (Life Technologies Corporation, Carlsbad, CA 92008 USA), 2.5 µL of MgCl₂ (50 mM), 4.0 µL of mixed dNTPs (10 mM), 1.25 µL of each primer (10 µM), 0.15 µL of Platinum™ Taq DNA Polymerase (Life Technologies Corporation, Carlsbad, CA 92008 USA) and ultrapure autoclaved water to a volume of 25 µL per reaction. The PCR thermal cycling consisted of an initial incubation at 94 °C for 3 min, followed by 35 cycles (94 °C for 30 s, 55 °C for 1 min, 72 °C for 1.5 min) and a final extension at 72 °C for 10 min.

The first round of nPCR-18Sb, nPCR-ITS1 and nPCR-CO1 were performed with 4 µL of extracted DNA, 2.5 µL of 10× PCR Buffer (KCL 50 mM; Tris HCl 10 mM; pH 9.0) (Life Technologies Corporation, Carlsbad, CA 92008 USA), 1.0 µL of MgCl₂ (50 mM), 0.5 µL of mixed dNTPs (10 mM), 1.0 µL of each primer (10 µM), 0.2 µL of Platinum™ Taq DNA Polymerase (Life Technologies Corporation, Carlsbad, CA 92008 USA) (5 U/µL) (Termoﬁschier Scientiﬁc) and ultrapure autoclaved water to a volume of 25 µL per reaction. The nPCR thermal cycling consisted of an initial incubation at 94 °C for 3 min, followed by 35 cycles (94 °C for 30 s, 55 °C for 1 min, 72 °C for 1.5 min) and a final extension at 72 °C for 10 min.

The first round of nPCR-18Sb, nPCR-ITS1 and nPCR-CO1 were performed with 4 µL of extracted DNA, 2.5 µL of 10× PCR Buffer (KCL 50 mM; Tris HCl 10 mM; pH 9.0) (Life Technologies Corporation, Carlsbad, CA 92008 USA), 1.0 µL of MgCl₂ (50 mM), 0.5 µL of mixed dNTPs (10 mM), 1.0 µL of each primer (10 µM), 0.2 µL of Platinum™ Taq DNA Polymerase (Life Technologies Corporation, Carlsbad, CA 92008 USA) (5 U/µL) (Termoﬁschier Scientiﬁc) and ultrapure autoclaved water to a volume of 25 µL per reaction. The PCR thermal cycling consisted of an initial incubation at 94 °C for 3 min, followed by 35 cycles (94 °C for 30 s, 56 °C for 30 s, 72 °C for 50 s) and a final extension at 72 °C for 5 min. For the second rounds: 2 µL of template derived from the first reactions, 2.5 µL of 10× PCR Buffer (KCL 50 mM; Tris HCl 10 mM; pH 9.0) (Life Technologies Corporation, Carlsbad,
CA 92008 USA), 1.0 µL of MgCl₂ (50 mM), 0.5 µL of mixed dNTPs (10 mM), 2.5 µL of each primer (10 µM), 0.2 µL of Platinum® Taq DNA (Life Technologies Corporation, Carlsbad, CA 92008 USA) (5 U/µL) and ultrapure autoclaved water to a volume of 25 µL per reaction. The thermal cycling was the same used in the first round.

DNA of Sarcocystis neurona, Neospora caninum and Hammondia hammondi was used as positive controls and ultrapure DNase-free water as the negative control for all reactions.

PCR products were resolved on 2.0% agarose gels and viewed through UV transillumination. Amplicons of the expected sizes were treated with ExoSAP-IT (Affymetrix/Thermo Fisher Scientific, Santa Clara, CA, USA), prepared for sequencing using the Big Dye Terminator cycle sequencing kit (Applied Biosystems, Foster City, CA, USA) and sequenced in an ABI automated sequencer (ABI 3500 Genetic Analyzer, Applied Biosystems). Sequencing was performed using the same primers as in the nPCR consensus. Sequence edition and contig assemblies were done by using the software Codoncode aligner, Codoncode Corporation. Final sequences were compared with homologous available in GenBank, using the BLASTn algorithm (Table S1) (https://blast.ncbi.nlm.nih.gov/Blast.cgi).

For the phylogenies, sequences were aligned using the Clustal W program, as implemented in the BioEdit Sequence Alignment Editor [34]. The phylogenetic tree based on ITS1 was inferred using MEGA X [35], through the maximum likelihood method and T92 model of evolutionary distances. Branch supports were tested through 1000 bootstrap replications.

3. Results
3.1. Molecular Identification

Sixty-six tissue samples from 22 seabirds were screened by nPCR-18Sa and only two samples of pectoral muscle from two Chilean skuas were positive. None of these two samples were positive for the T. gondii-specific nested-PCR (nPCR-B1). The two nPCR-18Sa-positive samples were also positive by nPCR-ITS1, nPCR-18Sb and nPCR-CO1 (Figure 1, left panel). After sequencing nPCR-ITS1, nPCR-18Sb and nPCR-CO1 amplicons and removal of primer-derived sequences, 861, 783 and 547 base pairs were obtained, respectively. Fragments of the sequences obtained are shown in Figure 1, right panel. The homologous sequences of the two samples were identical to each other, thus only one set was submitted to the GenBank, under the accession numbers MW160469, MW161469, MW157378. Through BLAST searches, ITS1, CO1 and 18S genetic sequences were compared with sequences producing the most significant alignments, with query coverage ≥ 99% and percentage similarities ≥ 99.00% in the cases of CO1 and 18S. All ITS1 sequences with query cover ≥ 96% were used for analyses of the genetic sequence of the skuas.

Figure 1. Left panel: Agarose gel electrophoresis of nPCR-ITS1 (1), nPCR-CO1 (2), and nPCR-18Sb (3) amplicons from Sarcocystis sp. ex Stercorarius chilensis, nPCR-ITS1 amplicons (4) from Sarcocystis neurona and Ladder Scada 100 bp, Sinapse, Inc. (L). Right panel: segments of the electropherogram obtained after sequencing nPCR-18Sb (top), nPCR-CO1 (middle), and nPCR-ITS1 (bottom) amplicons from Sarcocystis sp. ex Stercorarius chilensis.

The ITS1 fragment from the skuas showed 96.14–96.28% identity to sequences of Sarcocystis halieti from herring gulls (Larus argentatus) (MN450340–MN450356), great cormorants (Phalacrocorax carbo) (MH130209, JQ733513) and white-tailed sea-eagles (Haliaeetus albicilla) (MF946589–MF946596). Sarcocystis sp. from Cooper’s hawk (Accipiter cooperii)
eral species descri...

to find at least one positive animal, as the prevalence of the infection might not have been adeq...

generated in branches represent bootstrap values after 1000 replicates. The red dot identifies the sequence of Stercorarius chilensis.

3.2. Phylogeny

The ITS1-based phylogeny demonstrated that the species that shares the most recent ancestral commonality with Sarcocystis sp. ex Stercorarius chilensis was S. halieti (Figure 2). These sequences were separated with high support from a major clade comprising the species Sarcocystis sp. ex Columba livia (FJ232948), Sarcocystis calchasi (KC733715–KC733718) and Sarcocystis wobeseri (MN450365–MN450373, HM159421, JN256121), which exploit Anseriformes, Charadriiformes, Columbiformes, Psittaciformes and other birds as intermediate hosts.

Altogether, the Sarcocystis species that were most similar to Sarcocystis sp. ex Stercorarius chilensis used birds as intermediate hosts.

Figure 2. Phylogenetic tree of Sarcocystis species based on ITS1 sequences. The tree was constructed using the maximum likelihood method and Tamura 3 parameters nucleotide substitution model. The final alignment contained 49 sequences and 814 aligned nucleotide positions. All positions containing gaps and missing data were eliminated (complete deletion option). Numbers in branches represent bootstrap values after 1000 replicates. The red dot identifies the sequence of Sarcocystis sp. ex Stercorarius chilensis.

4. Discussion

Toxoplasma gondii has high genotypic diversity and several new genotypes have been described in wildlife [36–38], which has aided to understand the shape of molecular evolution and the epidemiology of the infection [38]. Similarly, several species descriptions have been made for the genus Sarcocystis, most of them with the aid of molecular methods [39–43]. This study presents the results of a molecular screening of Sarcocystidae focusing on animal species and geographical areas where these parasites have rarely or not
yet been identified. DNA of organisms of the genus *Sarcocystis* was identified in two Chilean skuas, whereas DNA of *T. gondii* were not found in any sample. The *Sarcocystis* haplotype detected in skuas was named *Sarcocystis* sp. ex *Stercorarius chilensis*.

Although all the samples tested negative for the presence of *T. gondii*, antibodies against this parasite were detected previously in 57 (43.18%) of the 132 serum samples from free-living Magellanic penguins from the same region, with titers that ranged from 20 to 320 [7]. Herein, infected animals were not encountered possibly because the sampling was insufficient to find at least one positive animal, as the prevalence of the infection in seabirds in the sampled area are not known and the sampling might have not been representative of the population surveyed. In addition, the mass of tissue that was tested might have been insufficient because of the very sparse and focal distribution of *T. gondii* cysts in the tissues of the HI, thus, digestion of samples previously to the DNA extraction and subsequent DNA detection would be more appropriate than direct DNA extraction, as used in this study [44].

Oocysts of *T. gondii* can sporulate and survive in seawater for months [45,46]. Marine mammals in different groups (cetaceans, pinnipeds and sirenians) and seabirds might become infected through consumption of water containing the oocysts. Thus, *T. gondii* oocysts from felidae feces might enter the marine environment and contaminate both the water and several invertebrate species, which could act as vectors of infection for mammals and seabirds [47]. Mice can be experimentally infected when fed with *T. gondii*-contaminated oysters (*Crassostrea virginica*) [45] proving that *T. gondii* was able to survive for several months in these mollusks [48]. Anchovies and Pacific sardines can be experimentally contaminated with *T. gondii* oocysts, which indicates that migratory filter feeders may serve as biotic vectors for this parasite [49]. Another study proved that freshwater crustaceans were able to bioaccumulate *T. gondii* oocysts. It should be noted that crustaceans are part of penguins’ and many seabirds’ food chain [50,51]. Thus, although the birds screened here were found not infected by *T. gondii*, marine fauna are at risk of acquiring the infection, by ingesting oocysts carried by transport hosts (oysters, fish and other) or through predation of intermediate hosts in the marine or in the coastal environment.

Based on molecular data, *Sarcocystis* sp. ex *Stercorarius chilensis* is an undescribed *Sarcocystis* species, closely related to *S. halieti*. The molecular identification based on ITS1, CO1 and 18S rRNA gene sequences showed a closed relationship between *Sarcocystis* sp. from Chilean skuas and other *Sarcocystis* spp. that use birds as intermediate hosts and predatory birds as definitive hosts. As expected, the most variable locus was ITS1, and phylogenies based on 18S rRNA and CO1 genes showed insufficient discrimination power to differentiate between species within the genus [39,41].

The most similar sequences to ITS1 of *Sarcocystis* sp. ex *Stercorarius chilensis* are those from *Sarcocystis* spp. that use hawks as definitive hosts. *Sarcocystis* sp. ex *Stercorarius chilensis* grouped together with *S. halieti*, a species that uses white-tailed sea-eagles (*Haliaeetus albicilla*) and Eurasian sparrowhawks (*Accipiter nisus*) as definitive hosts [39]. Other taxa found through ITS1-based BLAST searches encompass *Sarcocystis* spp. that also use hawks as definitive hosts (*Accipiter cooperii, Accipiter nisus*), except for *S. corvus*, for which this information remains unknown [52,53]. Accipiter hawks (*Accipiter gentilis, Accipiter nisus*) are definitive hosts for *S. calchasi* [54–56].

Several studies have expanded the knowledge on the host specificity of *Sarcocystis*, as unequivocal identification of the parasite can be achieved after identifying sarcocysts and oocysts to species level using molecular methods. *Sarcocystis halieti* and *Sarcocystis lari* were found to have formed oocysts in the intestine of white-tailed sea eagle (*Haliaeetus albicilla*), showing for the first time the potential role of sea eagle as definitive host of those species of *Sarcocystis* [53]. Likewise, European seabirds were found to harbor several species of *Sarcocystis* after DNA of *Sarcocystis lari*, *S. wobeseri*, *S. columbae* and *S. halieti* were detected in sarcocysts infecting muscle of herring gulls (*Larus argentatus*), great black-backed gulls (*Larus marinus*) and great cormorants (*Phalacrocorax carbo*) in Lithuania [40–42].
The four morphologically indistinguishable Sarcocystis species, *Sarcocystis lari*, *S. wobeseri*, *S. columbae* and *S. halieti*, could only be differentiated in *L. argentatus* by means of ITS1 sequence analysis [42]. Likewise, only ITS1 clearly discriminated *Sarcocystis* sp. ex *Stercorarius chilensis* from *S. halieti*, which reinforces that molecular characterization using this marker is of paramount importance to distinguish closely related species within the genus.

It is well known that a single animal can host more than one *Sarcocystis* species [40]. Here, sarcocysts were not individually excised and subjected to molecular examination, notwithstanding, the possibility of mixed infected samples of skuas was discarded because single peaks and clean sequence throughout the chromatograms were obtained for each sequence. Thus, a haplotype could be confidently assigned to the samples.

Although screening *Sarcocystis* by using molecular methods without morphological characterization of parasitic structures is obviously not enough to name a new species, this procedure may provide subsides to future studies on the epidemiology of the infection and its impact on the health of marine fauna. To our knowledge, *Sarcocystis* in south American seabirds were identified only once [43], which suggests that a wide field of research on diversity of sarcocystidae can be explored on this continent.

5. Conclusions

Although few animals have been screened in this study and morphological characterization of the parasites was not carried out, evidence of an unprecedented haplotype of *Sarcocystis* was found in skuas from Chile, which demonstrate that molecular screening of *Sarcocystis* can be a valuable tool to prospect for new species, contributing to knowledge on the epidemiology of sarcocystosis and life cycle of *Sarcocystis*. Sporocysts shed with feces, sarcocysts in tissues or rapid dividing structures in acute sarcocystosis (schyzonts and merozoites) can be more easily and accurately identified as data on *Sarcocystis* genetic sequences increases. Nevertheless, a complete study encompassing aspects of life cycle and morphological data is necessary to fully describe *Sarcocystis* sp. ex *Stercorarius chilensis* and additional studies are needed to better understand the epidemiology of the infection and its impact on the health of marine fauna.

**Supplementary Materials:** The following are available online at [https://www.mdpi.com/2076-2615/11/2/245/s1](https://www.mdpi.com/2076-2615/11/2/245/s1), Table S1. Results from BLAST search using sequences of ITS1, COX1 and 18S of *Sarcocystis* sp. ex *Stercorarius chilensis*.

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