

Research Article

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
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New insights into the coexistence of *Contracaecum rudolphii* A and *Contracaecum rudolphii* B (Nematoda: Anisakidae) in *Phalacrocorax carbo sinensis* from Sardinia: genetic variability and phylogenetic analysis

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Abstract

Contracaecum sp. nematodes are important parasites of fish eating birds that can cause animal health problems. In the present study, specimens of *Contracaecum rudolphii* sensu lato, from the great cormorant *Phalacrocorax carbo sinensis* from Sardinia, were characterized based on morphological and molecular data. The morphological analysis allowed to identify all the fourth stage larvae ($n = 1918$) as *Contracaecum* sp., and adults, male ($n = 5845$) and female ($n = 8312$), as *C. rudolphii* sensu lato. Population genetics and phylogenetic relationships were inferred based on mitochondrial and nuclear markers. Multiple sequence alignment of the ribosomal internal transcribed spacer showed the coexistence of *C. rudolphii* A ($n = 157$), *C. rudolphii* B ($n = 22$) and a rare heterozygote of these species. Moreover, mitochondrial markers, namely NADH dehydrogenase subunits I (*nad1*), cytochrome c oxidase subunit (*cox1* and *cox2*) and small subunit of rRNA (*rrnS*), showed that the studied *C. rudolphii* A populations had undergone bottleneck, or founder effect event, subsequent to a rapid population growth and expansion. The observed heterozygote is with a mitochondrial pattern of *C. rudolphii* B. Although, both *Contracaecum* species showed high genetic diversity, no genetic structure between localities was detected. Phylogenetic reconstructions supported the paraphyly of the avian *Contracaecum* species including *C. ogmorhini* (parasite of otariids).

Introduction

Contracaecum sp. nematodes are parasites of aquatic and terrestrial animals with a global distribution (Hartwich, 1964; Shamsi, 2019). This genus is considered the most diverse within the family Anisakidae (Hartwich, 1964), with over one hundred described species (Shamsi, 2019), and has a significant pathogenic impact on hosts (Shamsi, 2019). Molecular studies are providing a continuous update of *Contracaecum* nematode species composition. Mattiucci *et al.* (2010) reported two new species *C. gibsoni* and *C. overstreeti* from Greece, Garbin *et al.* (2011) described a new taxa *C. australe* from Chile and D'Amelio *et al.* (2012) reported two new taxa *C. fagerholmi* and *C. rudolphii* F from Mexico.

Contracaecum rudolphii Hartwich, 1964 is worldwide distributed and can be found as an adult in diverse seabirds, as cormorants (*Phalacrocorax* spp.) and pelicans (*Pelecanus* spp.) (Hartwich, 1964; Nadler *et al.*, 2000, 2005; Szostakowska *et al.*, 2002; Barson and Marshall, 2004; Li *et al.*, 2005; Mattiucci *et al.*, 2008, 2020; D'Amelio *et al.*, 2012; Cole and Viney, 2018; Shamsi, 2019). Epidemiological and molecular studies can help to explain the population genetics and the dispersal pattern of the different *Contracaecum* and host species, and provide useful data for the conservation of host seabird populations (Cole and Viney, 2018). Hosts can play an important role in increasing the population genetic variability of parasites, by mixing parasites from multiple localities. Indeed, host migration may lead to important gene flow among parasite populations, and so to a poor genetic structuring (Cole and Viney, 2018).

The variety of hosts and the wide range for spicule length of *C. rudolphii* suggested, since its first description (Hartwich, 1964), that this taxon included several sibling species. The application of allozyme analysis of *C. rudolphii* collected from *Phalacrocorax aristotelis* and *P. carbo sinensis* in Europe revealed the co-existence of two sibling species, *C. rudolphii* A and *C. rudolphii* B (Bullini *et al.*, 1986; D'Amelio *et al.*, 1990; Mattiucci *et al.*, 2003, 2020; Li *et al.*, 2005). Subsequently, based on molecular markers, several new species were characterized from different regions and hosts: D'Amelio *et al.* (2007) described *C. rudolphii* C from *P. auritus* in west-central Florida; Shamsi *et al.* (2009) described *C. rudolphii* D from *P. carbo* and *P. varius* and *C. rudolphii* E from *P. varius* in Australia; D'Amelio *et al.* (2012) described *C. rudolphii* F from *Pelecanus occidentalis* in Mexico.

In Europe, several studies confirmed *P. carbo* as the main host of *C. rudolphii* (Szostakowska *et al.*, 2002; Szostakowska and Fagerholm, 2012). Moreover, investigations on the parasites of brackish and freshwater fish showed that *C. rudolphii* B exists in both ecosystem, while *C. rudolphii* A occurs mainly in marine and brackish environments (Szostakowska *et al.*, 2002; Szostakowska and Fagerholm, 2007, 2012; Culurgioni *et al.*, 2014; Mattiucci *et al.*, 2020). However, Szostakowska and Fagerholm (2012) reported mixed infections in *P. carbo*, and a rare 'hybrid' of *C. rudolphii* A and B in the Baltic Sea.

Two sub-species of *P. carbo* are recognized in Europe, differing mainly in their geographical distribution. The sub-species *P. carbo carbo* (L.) lives in Europe, North Africa and on the Atlantic coast of North America, whereas *P. carbo sinensis* ranges from western Europe to China, India and North Africa. In Italy, several studies have shown that the breeding colonies of the great cormorant have disappeared in the past centuries, but not in Sardinia (Carpegna *et al.*, 1997), where between 2006 and 2012, an increase of about 82% in the reproductive number of cormorants (*P. carbo sinensis*) has been reported, corresponding to a total of 48 colonies.

The aim of the present study is to contribute to extending the knowledge on the *Contraecaecum* taxa that infect the great cormorant in the Mediterranean Sea (Sardinia), using both morphological and molecular data (four mitochondrial and one nuclear genes). This information will allow to provide a phylogenetic analysis of the studied taxa and their congeners, to elucidate the pattern of dispersion of *Contraecaecum* spp. populations, useful to evaluate the possible impact on the host.

Materials and methods

Sampling

The stomach contents of 60 specimens of *P. carbo sinensis* from six localities of Sardinia were examined for *Contraecaecum* sp. The hosts came from the controlled abatement campaign carried out from 5 to 27 February 2009 in the Oristano coastal brackish water ponds, in order to protect fish production (Decreto Regione Autonoma della Sardegna, No 2225/DecA/3, 30/01/2009). The first sorting of the stomach contents was carried out by Merops s.r.l. and the Oristano Department of the Istituto Zooprofilattico Sperimentale della Sardegna. The localities ($n = 6$) and the number of hosts examined are shown in Fig. 1. Collected specimens of *Contraecaecum* sp. were sorted by locality, host, stage and sex, then fixed in ethanol 70% prior to morphological and molecular analyses.

Morphological identification

Nematodes were identified based on the main characters with taxonomic significance, i.e. shape of interlabia (both sexes), number and pattern of pre- and post-cloacal papillae, length and shape of the extremity of spicula (males) (Hartwich, 1974; Barus *et al.*, 1978; Abollo *et al.*, 2001). Specimens were deposited in the collection of the Department of Zoology, King Saud University.

DNA extraction, amplification and sequencing

From each locality, five hosts were randomly selected, and from each host, six specimens of *C. rudolphii* (two fourth-stage larvae, two adult males and two females) were stored in 70% ethanol for molecular studies (N total = 180). Genomic DNA was extracted using SDS/proteinase K protocol (Gasser *et al.*, 1993).

Polymerase chain reactions (PCR) were performed in a 50 μ L reaction mix containing 100 ng DNA, ddH₂O, 1U Taq polymerase (Bioline, USA), 5X buffer and 5 pmol from each primer (further information on the primer sets are listed in the Supplementary Table S1). One nuclear locus, the ribosomal internal transcribed spacer (ITS) including ITS1 and 5.8S ribosomal RNA gene and ITS2, and four mitochondrial loci, the cytochrome c oxidase 1 (*cox1*), cytochrome c oxidase 2 (*cox2*), the NADH dehydrogenase subunit I (*nad1*) and the small subunit of rRNA (*rrnS*), were PCR-amplified. Amplifications comprised a denaturation at 95°C (4 min), 35 cycles at 95°C for 60 s, 55°C (ITS), 50°C (*cox1*, *cox2*, *nad1* and *rrnS*) for 90 s and 72°C for 60 s with a final extension at 72°C for 8 min. Negative controls were added to check for eventual contamination. PCR products were sequenced at Macrogen sequencing facility (Macrogen Inc, Seoul, Korea).

Data analysis

The obtained sequences were checked and aligned using Unipro UGENE 1.3 (Okonechnikov *et al.*, 2012). Representative sequences of *Contraecaecum* sp. (Supplementary Table S2) were included in the datasets from Genbank using BLAST algorithm (Altschul *et al.*, 1990). Mixed Sequencer Reader (MSR program) was used to examine mixed chromatograms (Chang *et al.*, 2012), useful to identify and compare heterozygous ITS sequences. For each base position, MSR calculates the log ratio of the chromatographic intensities (LRi). The LRi value varies from 0, for heterogeneous chromatographic traces with equivalent intensities, to infinite, for position containing only one main band.

DnaSP v 5.10 (Librado and Rozas, 2009) was used to estimate the number of haplotypes (h), nucleotide (π) and haplotype (Hd) diversities. Network 5 (Fluxus Technology Ltd., United Kingdom) was used in order to generate Median-Joining network (Bandelt *et al.*, 1999) depicting evolutionary relationships between haplotypes obtained from each single marker and from the concatenated mitochondrial set. Analysis of molecular variance (AMOVA) was estimated between populations with sufficient sample size, using Arlequin 3.5 (Excoffier and Lischer, 2010). *Contraecaecum* sp. populations were clustered according to locality into six groups (Fig. 1).

Fu's F_s (Fu, 1997), Tajima's D (Tajima, 1989) and Mismatch distributions were performed for all mitochondrial markers to evaluate changes in population size using DnaSP v 5.10 (Librado and Rozas, 2009).

Models of evolution and the best partitioning scheme were inferred, for each marker and for the concatenated sequences, using PartitionFinder 2.1 (Lanfear *et al.*, 2016). The appropriate models were selected under the best value for the Bayesian Information Criterion (BIC) (Lanfear *et al.*, 2016). Sequences of *Toxocara cati* (KY003086, AM411622) and *Hysterothylacium auctum* (AF115571) were used as outgroup (Supplementary Table S2). Bayesian (BI) and maximum likelihood (ML) phylogenetic trees were constructed using MrBayes v3.0 (Huelsenbeck and Ronquist, 2001) and RAXML (Stamatakis, 2006), respectively. BI analyses were performed with two independent runs of 1×10^8 generations. Trace plots were used to test the number of burn-in fraction (20%). The chains were sampled every 1000 generations. For ML analyses, bootstrap was assessed by 2000 pseudoreplicates.

Results

Morphological identification

The morphological analysis of 15375 nematodes, based on the size and shape of interlabia (both sexes), and the number and pattern of the pre- and post-cloacal papillae, length and shape

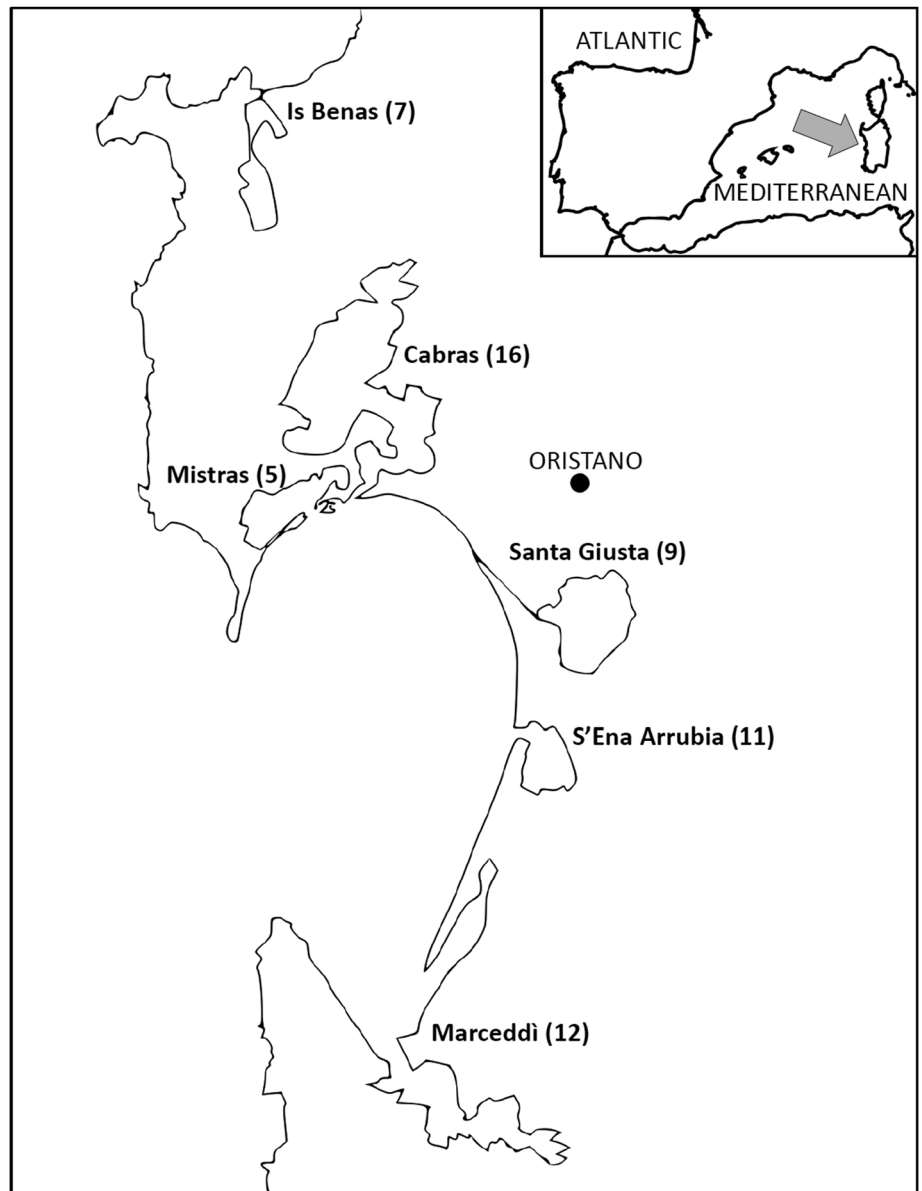


Fig. 1. Sampling localities of *Phalacrocorax carbo sinensis* in the Oristano coastal brackish water ponds. In brackets number of hosts examined.

of spicula (males), allowed to identify adult males ($n = 5845$) and females ($n = 8312$) as *C. rudolphii* sensu lato, and the fourth stage larvae as *Contracaecum* sp. ($n = 1218$). The main morphological characters were the following. Adults, cuticle brownish-yellowish and striated transversely, lips and interlabia well developed, interlabia with bilobed distal end. Males, smaller than females, 20 mm (14–25 mm) long, tail conical, curved at the tip, with 27–40 pairs of pre-cloacal papillae and seven pairs of post-cloacal papillae, spicules sub-equal with longitudinal alae and rounded tips. Females, 30 mm (18–48 mm) long, vulva located on the second quarter of body length, tail conical, rounded at the tip. Fourth stage larvae, 7.5 mm (5–12 mm) long, cuticle whitish, with striations at the anterior extremity, oral opening with three lips, interlabia absent, tail conical without papillae.

Table 1 shows the levels of infection of fourth stage larvae and adult males and females according to sex, weight and locality of the host. All the examined specimens of *P. carbo sinensis* were infected with adults, and 88% with fourth-stage larvae. Total intensity of infection ranged from 15 (Santa Giusta) to 1075 (Cabras). Hosts from Cabras and Mistras showed the highest total mean intensity of infection (334.9 and 332.2, respectively), those from Is Benas and Santa Giusta the lowest (143.6 and 179.7, respectively). In all localities, the intensity of infection of

fourth stage larvae was the lowest, and that of females the highest, showing an unbalanced sex ratio (0.7:1.0).

Molecular characterization

Sequences of the ribosomal ITS [ITS1 (434 bp) and ITS2 (286 bp)] revealed the presence of *C. rudolphii* A ($n = 157$, accession numbers: MT341236, MT341180), *C. rudolphii* B ($n = 22$, accession numbers: MT341234, MT341178) (Table 2, Supplementary Fig. S1A). Pairwise comparison revealed seven variable positions within ITS1 and 4 substitutions and an indel (9 bp) within ITS2. One sample from Is Benas showed a mixed chromatogram (accession numbers: MT341235, MT341179). The analysis of the mixed chromatograms with MSR (Supplementary Fig. S1A–C) exhibited a shift of signal intensity at 635 bp, corresponding to an ‘indel’-type heterozygosity. Eleven LRi line drops were observed, corresponding to the heterogeneous fluorescence traces within the chromatogram (Supplementary Fig. S1B, C).

All the fourth-stage larvae were identified as *C. rudolphii* A. Fifteen hosts from all the six localities showed a mixed infection. Of the five *P. carbo* from Mistras, only one was infected with both *Contracaecum* species, whereas hosts from Is Benas showed the highest rate of coinfection (4/5).

Table 1. Levels of infection of *Contracaecum rudolphii* in *Phalacrocorax carbo sinensis* from the Oristano ponds according to sex, weight and locality.

	P%	Mean I	Mean I (P = 100%)		
			L4	Adult M	Adult F
Tot	88%	20.3	97.4	138.5	256.3
Host sex					
M	86%	18.6	108.9	161.3	288.8
F	89%	21.8	91.5	126.8	240.1
Host weight (kg)					
1.5–2.0	92%	25.9	100.2	140.4	266.5
2.0–2.5	92%	23.8	101.0	146.2	271.1
2.5–3.0	80%	13.1	97.1	140.3	250.5
3.0–3.5	80%	6.0	66.6	88.6	161.2
Locality					
Cabras	69%	19.0	125.6	190.3	334.9
Is benas	100%	24.7	46.9	72.0	143.6
Marceddi	100%	16.4	105.3	145.5	267.2
Mistras	100%	37.8	129.4	165.0	332.2
Santa Giusta	89%	17.4	70.0	92.2	179.7
S'ena Arrubia	91%	18.0	88.0	123.8	229.8

P%, prevalence; Mean I, mean intensity of infection.

The length of the PCR products of the mitochondrial markers *cox1*, *cox2*, *nad1* and *rrnS* was, approximately, 450, 560, 400 and 530 bp, respectively. For all mtDNA region, no size variation was detected on agarose gel, but among *nad1* amplicons, two different patterns were observed, 200 and 400 bp (Fig. 2). No pseudo-genes were identified after the examination of mitochondrial genes translation. The nucleotide variation of the *cox1*, *cox2*, *nad1* and *rrnS* was mainly related to changes at the third codon position, while few changes were detected at the first or second position. The G + C contents were 36.8% (*cox1*), 36.2% (*cox2*), 30.0% (*nad1*) and 30.5% (*rrnS*), respectively (Table 3).

Phylogenetic relationships

For all mitochondrial markers, both *C. rudolphii* A and *C. rudolphii* B were characterized by a very high haplotype diversity

whereas their respective nucleotide diversity was very low (Table 3). Most haplotypes were unique and represented by a single individual (Table 4).

Alignment of the partial *cox1* sequences (440 bp) of *C. rudolphii* A and *C. rudolphii* B generated 64 and 14 haplotypes, respectively (Table 4, Fig. 3a) (accession numbers: MT338400-77). *Contracaecum rudolphii* A and *C. rudolphii* B differed by 33 mutations. For *C. rudolphii* A, only five haplotypes (H_{A1} , H_{A2} , H_{A4} , H_{A21} and H_{A36}) were shared among localities. *Contracaecum rudolphii* A network consisted of two star-like haplogroups, the first included 53 haplotypes with three major haplotypes (two from Is Benas, one from Cabras) and the second one was more aggregated, with one haplotype from S'Ena Arrubia grouping 11 haplotypes. Both network patterns were not dependent on the samples' origin. The heterozygote sample showed the *C. rudolphii* B pattern with the following haplotype combination: $H6_{cox1}$ - $H3_{cox2}$ - $H1_{nad1}$ - $H7_{rrnS}$.

Alignments of the mtDNA *cox-2* sequences (540 bp) of *C. rudolphii* A and *C. rudolphii* B included 61 and 40 variable sites, respectively (accession numbers: MT338999-038). Network analysis revealed a star-like haplogroup for each studied species. *Contracaecum rudolphii* A haplogroup consisted of 33 haplotypes aggregated around three major haplotypes (two from Is Benas, one from Cabras) and differing from *C. rudolphii* B haplogroup by 33 mutations (Table 4, Fig. 3b).

Mitochondrial *nad1* sequencing generated variable sequence sizes, 400 bp for all *C. rudolphii* A haplotype ($n = 56$) and for one *C. rudolphii* B haplotype, and 200 bp for the four remaining *C. rudolphii* B haplotypes (accession numbers: MT338938-99). Multiple alignment of *nad1* dataset revealed a homologous region to the forward primer starting from the position 166. The network analysis yielded one star-like haplogroup for each studied species differing by 32 mutations. *Contracaecum rudolphii* A haplotypes were aggregated around three haplotypes, whereas *C. rudolphii* B haplotypes were derived from a major haplotype (Table 4, Fig. 3c).

Analysis of *rrnS* sequences (530 bp) yielded two star-like haplogroups differing by 12 mutations, including *C. rudolphii* A ($n = 39$) and *C. rudolphii* B ($n = 14$) haplotypes (accession numbers: MT341181-233). *Contracaecum rudolphii* A samples were divided into two main groups differing in one site from major haplotypes H_{A1} and H_{A22} (Table 4, Fig. 3d). The second haplogroup included *C. rudolphii* B samples, differing from the major haplotype H_{B1} by one to three mutations.

The obtained matrix of the four concatenated mitochondrial genes consists of 1940 bp after alignment (450 bp for *cox1*, 560 bp for *cox2*, 400 bp *nad1* and 530 bp for *rrnS*). In the total matrix, 292 positions were variable of which 231 were parsimony informative. Multiple alignment of the concatenated set generated

Table 2. Number of identified individuals of *Contracaecum rudolphii* A, and *C. rudolphii* B, and heterozygote of these (*C. rudolphii* A/B), in cormorants from the different localities of the Oristano ponds

Locality	Hosts		Parasites			Total
	Tot	Mixed infection	<i>C. rudolphii</i> A	<i>C. rudolphii</i> B	<i>C. rudolphii</i> A/B	
Cabras	5	3	24	6	0	30
Marceddi	5	2	25	5	0	30
Is benas	5	4	26	3	1	30
Mistras	5	1	28	2	0	30
Santa Giusta	5	3	26	4	0	30
S'Ena Arrubia	5	2	28	2	0	30

Fig. 2. Gel showing the size variation in PCR products of the mitochondrial nad1, the two different patterns, 200 bp and 400 bp. lanes 1, 3–6, 10 and 11 representative samples of *C. rudolphii* A and lanes 2, 7–9 of *C. rudolphii* B.

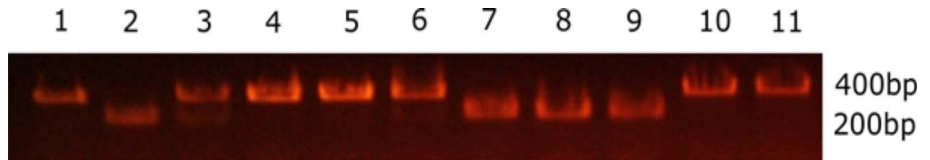


Table 3. Genetic variability and G+C content of four mitochondrial markers, cox1 (450 bp), cox2 (560 bp), nad1 (400 bp) and rrnS(530 bp), within *C. rudolphii* A (n=158) and *C. rudolphii* B (n=22), from the Oristano ponds

		H	Number of variable sites	G + C	Hd	π
cox1	B	14	38	0.362	0.970	0.022
	A	64	59	0.374	0.994	0.016
cox2	B	7	40	0.359	0.952	0.021
	A	33	61	0.365	0.985	0.013
nad1	B	5	13	0.299	0.731	0.0071
	A	56	50	0.306	0.986	0.0166
rrnS	B	14	17	0.305	0.972	0.0101
	A	39	64	0.306	0.894	0.0166

H, haplotype number; Hd, haplotype diversities; π , nucleotide diversity.

75 and 17 mitochondrial haplotypes for *C. rudolphii* A and *C. rudolphii* B, respectively (Fig. 4a, b). Network analysis generated two star-like haplogroups, supporting the results of single markers analysis. One major haplotype was shared by three populations, Cabras, Mistras and S'Ena Arrubia, in the case of *C. rudolphii* A and one was observed in two populations, Cabras and Marceddi, for *C. rudolphii* B (Fig. 4a, b).

AMOVA analysis revealed that the highest amounts of total genetic variation originated from differentiation within populations. Within populations proportions varied from 95% ($P < 0.0001$) for rrnS to 100% ($P < 0.0001$) for cox1, cox2 and nad1 (Table 5).

Neutrality tests Tajima's *D* and Fu's *F_s* revealed for *C. rudolphii* A significant negative values (Supplementary Table S3) indicating an excess of rare polymorphisms in the studied population suggesting a recent expansion and/or positive selection (Tajima, 1989). Overall Tajima's *D* and Fu's *F_s* statistics in *C. rudolphii* B datasets were not statistically significant and consistent with a population at drift-mutation equilibrium (Pichler, 2002). Mismatch distributions for all markers in *C. rudolphii* A were unimodal (Supplementary Fig. S2), patterns in concordance with the above neutrality test values supporting the assumption of a sudden expansion model. Mismatch distribution plot *C. rudolphii* B showed a multimodal shape, revealing demographic equilibrium or a stable population.

The best evolution models and the optimal partition schemes selected for each single marker and for the concatenated mitochondrial set are shown in Table 6. BI analysis and ML yielded trees with similar topology for the nuclear ITS, the four mitochondrial loci (cox1, cox2, nad1 and rrnS) and the concatenated dataset (Fig. 5). All the obtained trees showed two highly supported clades including *C. rudolphii* A and *C. rudolphii* B samples, which only differed from the resolution within both clades. *Contraecacum ogmorhini* (parasite of the otariid *Arctocephalus pusillus*) appeared within the highly-supported clade of *C. rudolphii* (avian hosts), as sister group to *C. rudolphii* B in all mitochondrial trees, or sister group to [*C. rudolphii* D + (*C. rudolphii* E + *C. rudolphii* F)] in ITS tree. *Contraecacum eudyptulae* and *C. septentrionale* are sister group to the clade containing *C. rudolphii* complex (A, B, C, D, E and F) + *C. ogmorhini*. Only in cox1 tree, *C. septentrionale* was more divergent with a basal position. Cox2 topology revealed a highly supported clade including *C. ogmorhini*, to the avian *C. margolisi* (host: California sea lion). In ITS and cox2 trees, *C. osculatum* complex, *Phocascaris cystophorae* *C. radiatum* and *C. miroungae* (all from phocid hosts) appeared as sister

Table 4. Observed mitochondrial haplotypes of two *Contraecacum* spp. used in this study, with their numbers, codes and locality

<i>Contraecacum</i> spp.	Locality	cox1		cox2		nad1		rrnS	
		Number	Codes	Number	Codes	Number	Codes	Number	Codes
B	Cabras	5	H1-H5	2	H1-H2	2	H1-H2	3	H1-H3
	Marceddi	3	H3, H8-H9	1	H1	1	H1	4	H3-H6
	Is benas	2	H6-H7	1	H3	2	H1, H3	2	H7-H8
	Mistras	1	H6	1	H4	1	H1	2	H9-H10
	Santa Giusta	3	H12-H14	1	H5	1	H4	3	H3, H11-H12
	S'Ena Arrubia	2	H10-H11	2	H6-H7	2	H3, H5	2	H13-H14
A	Cabras	8	H1-H8	9	H1-H9	10	H1-H9, H21	8	H1-H8
	Marceddi	12	H2, H4, H22-H31	7	H9-H15	10	H10, H13, H15, H19-H20, H23-H27	6	H1-H2, H9-12
	Is benas	13	H9-H21	8	H1, H11-H17	9	H10-H18	10	H1, H12-H20
	Mistras	11	H1, H31-H42	10	H1, H18-H26	12	H3, H19, H22, H28-H36	11	H1, H2, H21-H29
	Santa Giusta	13	H11, H18, H21, H55-H64	4	H27-H30	3	H3, H19, H56	9	H1, H2, H30-H36
	S'Ena Arrubia	14	H1, H36, H43-H54	4	H1, H31-H33	10	H3, H37-H45	6	H1-3, H37-H39

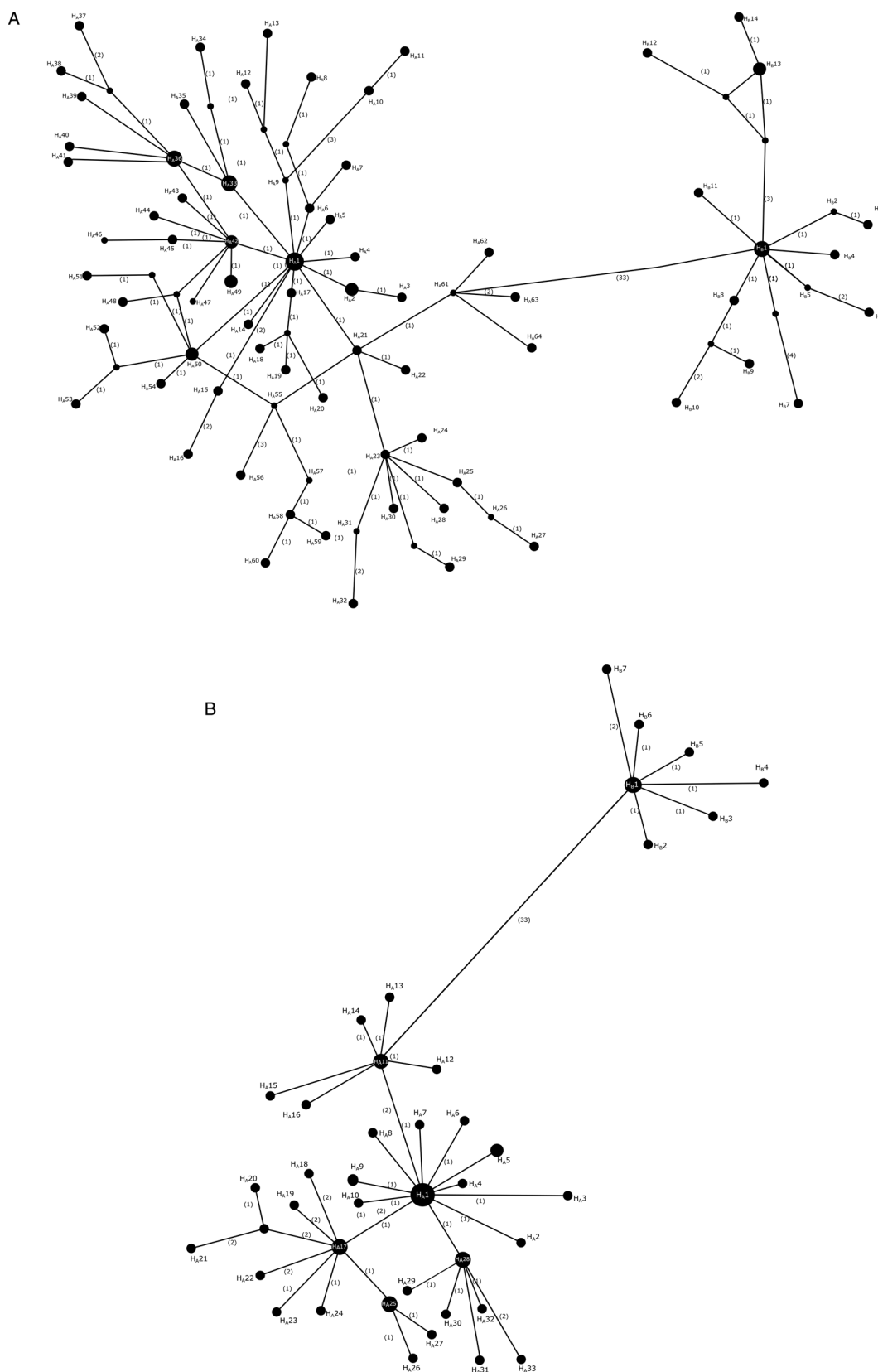


Fig. 3. Haplotype network obtained from (a) *cox1* (450 bp), (b) *cox2* (560 bp), (c) *nad1* (400 bp) and (d) *rnsS* (530 bp). (Ha) *C. rudolphii* A haplotype, (Hb) *C. rudolphii* B haplotype. The area of each circle is proportional to the haplotype frequency and mutation numbers are indicated in each branch. The heterozygote sample showed the *C. rudolphii* B pattern with the following haplotype combination: H6_{cox1}-H3_{cox2}-H1_{nad1}-H7_{rnsS}.

group to *C. microcephalum*, *C. micropapillatum* and *C. multipapillatum* species that mature in birds. Trees of both BI and ML analyses are shown in Fig 5, with node support including BI posterior probability and ML bootstrap values.

Discussion

Combination of morphological and molecular data allowed the identification of *Contracaecum* specimens from *P. carbo sinensis*

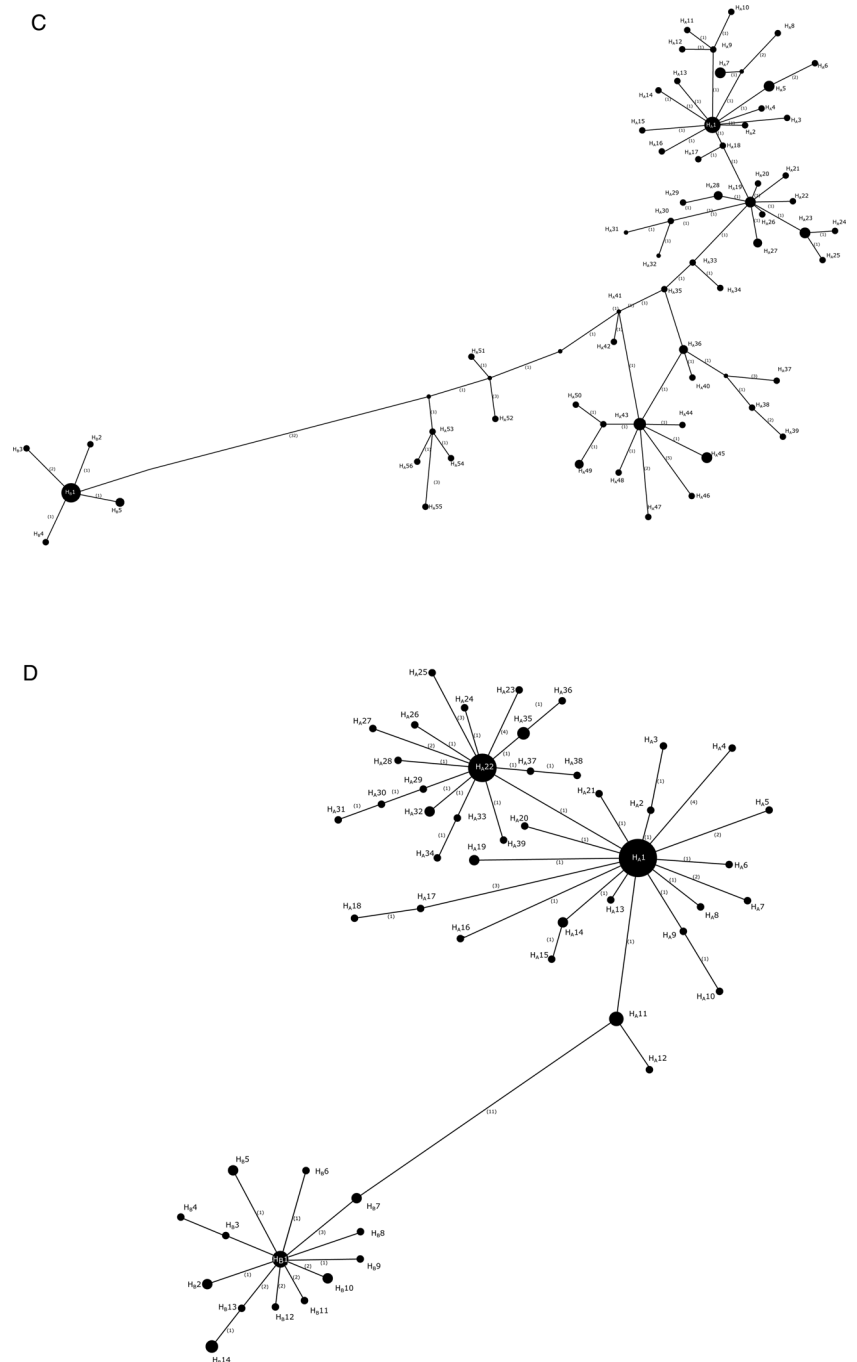


Fig. 3. Continued.

of the Oristano ponds as *C. rudolphii* A and *C. rudolphii* B. It is confirmed that sequence analysis of both nuclear and mitochondrial markers effectively discriminates both *Contraecaecum* spp. The present study confirms that the two *Contraecaecum* spp. can coexist in a single host (bird), as previously suggested by Szostakowska and Fagerholm (2007, 2012). *Contraecaecum rudolphii* A ranges mainly in marine and brackish waters of Europe, whereas *C. rudolphii* B in the freshwater habitats of the central Europe (Abollo *et al.*, 2001; Szostakowska *et al.*, 2002; Li *et al.*, 2005; Szostakowska and Fagerholm, 2007, 2012; Farjallah *et al.*, 2008a; Mattiucci *et al.*, 2020). The identification of all the fourth stage larvae as *C. rudolphii* A and none as *C. rudolphii* B suggests that only specimens of *C. rudolphii* A had recently infected the cormorants in the same sampling locality, while the infections of *C. rudolphii* B probably occurred sometime before, and not in the sites of sampling. Indeed, *P. carbo sinensis* is a migratory species,

widespread in the inland and coastal wetlands of central Europe and the Mediterranean Sea. Thus, it is probable that *C. rudolphii* B could have infected the host in the central European region, where this species is common in brackish and freshwater habitats (Szostakowska *et al.*, 2002; Szostakowska and Fagerholm, 2007, 2012; Mattiucci *et al.*, 2020). In the same Island of Sardinia (Archipelago di La Maddalena), Farjallah *et al.* (2008a) reported only *C. rudolphii* A from the European shag *P. aristotelis*. Unlike its congener, *P. aristotelis* is a typical marine species, feeding mainly on marine fish, and with a more restricted geographical distribution (western Palearctic).

Several studies confirmed the important role of *Phalacrocorax* spp., and several other bird species, for the dispersion of *Contraecaecum* spp., and in modelling their genetic variability within the distribution areas (Szostakowska and Fagerholm, 2007, 2012; Farjallah *et al.*, 2008a; Mattiucci *et al.*,

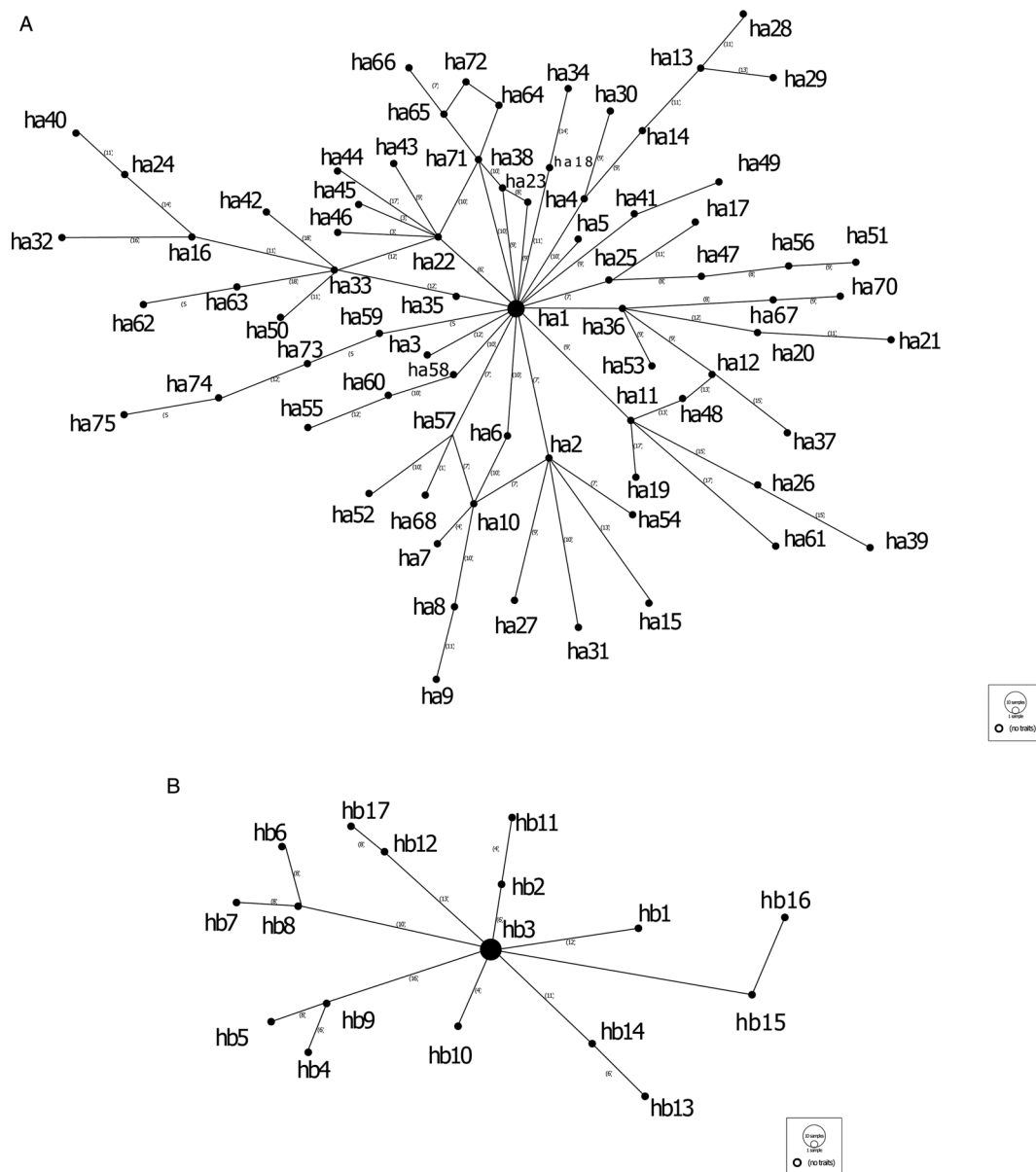


Fig. 4. Haplotype network obtained from concatenated mitochondrial loci [*cox1* (450 bp), *cox2* (560 bp), *nad1* (400 bp) and *rns*(530 bp)]. (a) *C. rudolphii* A haplotype network, (b) *C. rudolphii* B haplotype network. The area of each circle is proportional to the haplotype frequency and mutation numbers are indicated in each branch.

2008, 2020). The pattern of distribution of *C. rudolphii* A and *C. rudolphii* B has long been debated, and although several studies have demonstrated that both species may occur in fresh, brackish and marine water systems (Nadler, 2000; Szostakowska and Fagerholm, 2007, 2012; Farjallah *et al.*, 2008a; Mattiucci *et al.*, 2008), present results show that *C. rudolphii* A appears to be more seaward and *C. rudolphii* B more freshwaterward, as recently suggested by Mattiucci *et al.* (2020). In fact, the out-numbering of *C. rudolphii* A compared to *C. rudolphii* B (157 specimens *vs* 22), as well as the identification of all fourth stage larvae such as *C. rudolphii* A suggest that *C. rudolphii* A was acquired by the host in the studied brackish water ponds; on the other hand, the absence of larval stages of *C. rudolphii* B suggests that this species has infected the host in environments other than those of the studied lagoons. Considering the migratory habits of *P. carbo sinensis*, which in Sardinia is mainly present as wintering, it is likely that *C. rudolphii* B could have infected the host in the breeding grounds of central Europe,

where the third-stage larvae of this species have been identified in fish (Szostakowska and Fagerholm, 2007; Mattiucci *et al.*, 2008; Molnár *et al.*, 2019).

Among all *C. rudolphii* examined, multiple alignment of ITS sequences combined to chromatogram analysis distinguished one heterozygote *C. rudolphii* A/B. Previously, only Szostakowska and Fagerholm (2012) reported a similar heterozygote in the Baltic Sea. Few other studies reported 'hybrids' within anisakid species, i.e. *A. simplex*/*A. pegreffii* in the Atlantic, Mediterranean Sea and in Western Pacific Ocean (Abollo *et al.*, 2003; Marques *et al.*, 2006; Farjallah *et al.*, 2008b; Cavallero *et al.*, 2012, 2014); however, such observations are still very rare. Even when parental species are reproductively isolated, the coexistence of different anisakid congeneric species, with high intensity in single hosts, increases the probability of observing such a rare heterozygote. It should be emphasized that the heterozygote was found in Is Benas, where the hosts showed the highest coinfection rate (4/5). The present results confirm that in the brackish-marine waters

Table 5. Analysis of molecular variance (AMOVA)

	Source of variation	D.F.	Sum of squares	Variance components	% variation	P value
Cox1	Among groups	3	13.288	-0.359 Va	-4.495	0.000
	Among populations Within groups	5	24.192	0.325 Vb	4.48	0.000
	Within populations	179	605.924	7.305 Vc	100.47	0.000
	Total	187	643.404	7.266		
Cox2	Among groups	3	11.004	-0.41063 Va	-5.11799	0.000
	Among populations Within groups	5	26.776	0.34785 Vb	4.33550	0.000
	Within populations	179	697.405	8.08611 Vc	100.78249	0.000
	Total	187	735.185	8.02333		
Nad1	Among groups	3	7.513	-0.03732Va	-1.37	0.000
	Among populations Within groups	5	6.357	0.03514 Vb	1.29	0.000
	Within populations	179	186.008	2.73541 Vc	100.08	0.000
	Total	187	199.878	2.73322		
rrnS	Among groups	3	11.631	0.08149Va	3.49	0.000
	Among populations Within groups	5	4.922	0.01679 Vb	0.71.3	0.000
	Within populations	179	151.118	2.25549 Vc	95.82	0.000
	Total	187	167.671	2.35378		

Percentage of variation explained by different hierarchical levels for cox1, cox2, nad1 and rrnS loci. Degrees of freedom (D.F.).

Table 6. Evolutionary models by gene and codon position found in PartitionFinder.

	Subset	Best model	Partition names
ITS	1	HKY + G	ITS
cox1	1	GTR + G	Gene1_pos1
	2	F81	Gene1_pos2
	3	GTR + G	Gene1_pos3
cox2	1	HKY + I	Gene1_pos1, Gene1_pos2
	2	HKY + G	Gene1_pos3
nad1	1	HKY + I + G	Gene1_pos1, Gene1_pos2
	2	GTR + G	Gene1_pos3
rrnS	1	HKY + G	rrnS
Concatenated (cox1, cox2, nad1, rrnS)	1	GTR + I + G	Gene3_pos2, Gene1_pos1, Gene2_pos1, Gene2_pos2, Gene1_pos2
	2	GTR + G	Gene1_pos3, Gene2_pos3, Gene3_pos3
	3	HKY + I + G	rrnS, Gene3_pos1

of the southwestern Atlantic Ocean and of the southern Mediterranean Europe *C. rudolphii* A is the most abundant species in *Phalacrocorax* spp., whereas in the same regions *C. rudolphii* B is dominant in freshwater habitats (Abollo *et al.*, 2001; Li *et al.*, 2005; Farjallah *et al.*, 2008a, 2008b; Mattiucci *et al.*, 2020). On the other hand, in the cormorants of northeastern Europe, *C. rudolphii* B is always more abundant than *C. rudolphii* A, although the second species is significantly more abundant in the brackish water habitats compared to freshwater ones, where it is practically absent (Szostakowska and Fagerholm, 2012). The coinfection rate of *C. rudolphii* A and *C. rudolphii* B ranged from 20 to 80% (mean 50%), a similar value (42%) was reported by Szostakowska and Fagerholm (2012) in brackish water habitats of northern Europe, whereas the same authors found a much lower coinfection in the freshwater habitats of the same region (17%).

For mitochondrial markers, the AT content was higher than 65%, which agrees with the previous studies of *Contracaecum* species (Nadler *et al.*, 2000; Dzido *et al.*, 2012; Lin *et al.*, 2012; Delgado and Garcia, 2015). Polymorphism analyses of *C. rudolphii* A and *C. rudolphii* B based on several markers revealed high Hd and low Pi, which are evidence of an important genetic diversity (Ferreri *et al.*, 2011). Similar values have also been reported for other parasite species (Szostakowska and Fagerholm, 2007, 2012; Dzido *et al.*, 2012; Lin *et al.*, 2012; Szudarek *et al.*, 2017). Such pattern of diversity, coupled to the 'star-like' haplotype networks with a high ratio of singletons, indicates that the studied *C. rudolphii* A populations had undergone a bottleneck or founder effect event subsequent to a rapid growth and expansion of the population (Ferreri *et al.*, 2011; De Jong *et al.*, 2011; Levin and Parker, 2013; Dabert

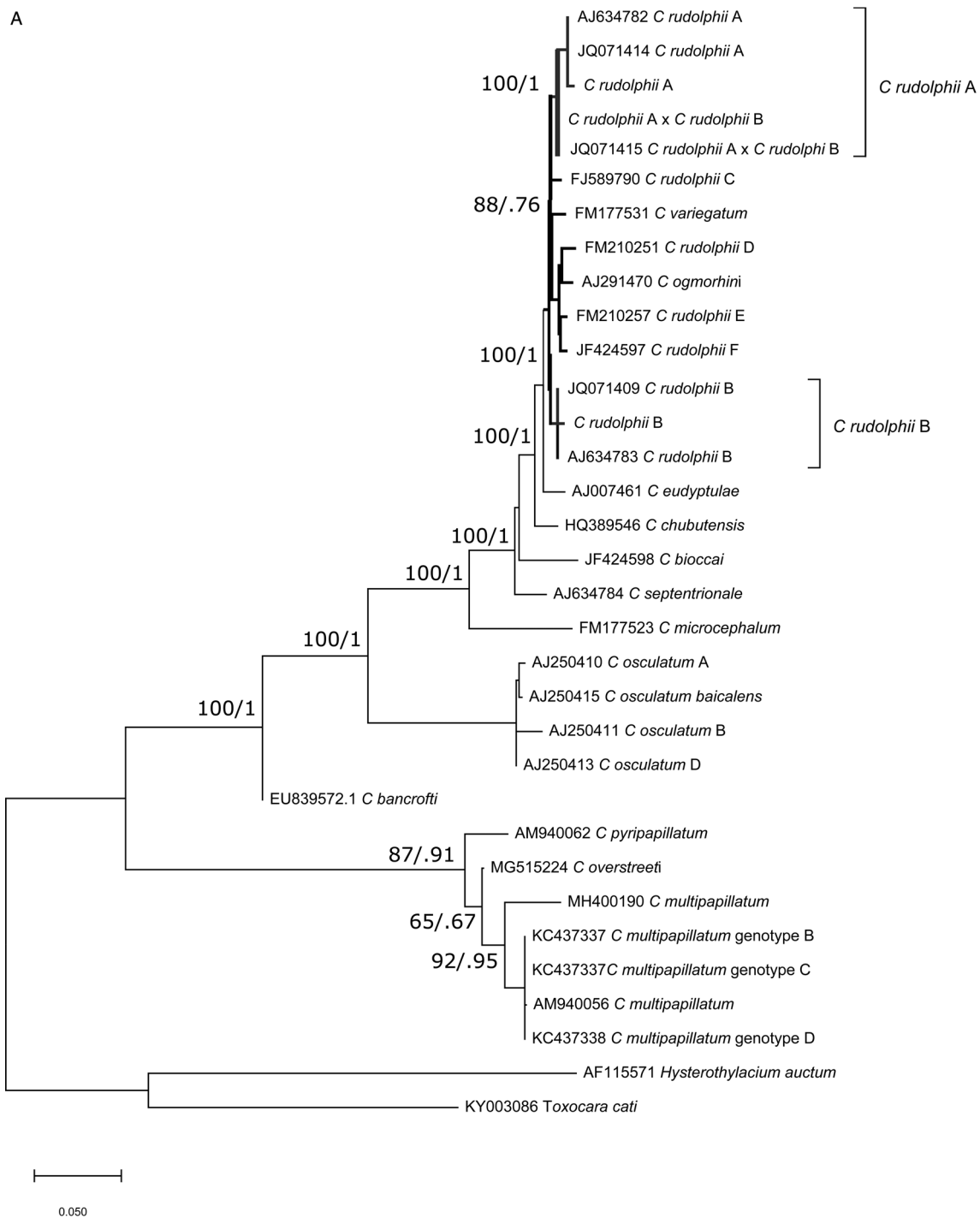


Fig. 5. Inferred phylogenetic relationships among *Contracecum* spp. based on sequences of (a) ITS, (b) *cox1*, (c) *cox2*, (d) *nad1*, (e) *rrnS* and (f) concatenated mitochondrial loci. The numbers along branches indicate bootstrap probability (BP) values resulting from ML analysis and posterior probability (PP) from the MrBayes analysis.

et al., 2015). Neutrality test values and Mismatch distributions observed in *C. rudolphii* A samples provide strong evidence for past population expansion. These parameters pointed to different demographic history in the case of *C. rudolphii* B. However, it is important to point out that in the case of *C. rudolphii* B, small numbers of samples originated from different localities and hosts were studied. This may conduct to misevaluate the polymorphism level for these samples (Rand *et al.*, 1994). Thus, it will decrease the ability to detect evolutionary forces acting at the population level (Schmid *et al.*, 1999).

Population genetic analysis of *Contracecum* samples revealed high genetic diversity, and no genetic structure among different localities. The lack of genetic structure is common within parasitic nematodes and several hypotheses have been suggested to clarify such pattern. One hypothesis is the high intensity of infection of these nematodes in the definitive host (mammals and birds) leading to a high effective population size (N_e), which slows down the genetic drift (Cole and Viney, 2018). A second hypothesis is the parasite dispersal model, actually the long distance gene flow mediated by host dispersal dynamic was suggested as the cause of lack of differentiation between parasite populations (Slatkin



Fig. 5. Continued.

and Hudson, 1991; Li *et al.*, 2005; Levin and Parker, 2013; Dabert *et al.*, 2015; Cole and Viney, 2018).

Phylogenetic reconstructions yielded two well-supported monophyletic clades corresponding to *C. rudolphii* A and *C. rudolphii* B, and thus confirm the status of the studied taxa. This topology was concordant with the haplotype network results. In all trees, *C. eudyptulae* and *C. septentrionale* are sister group to

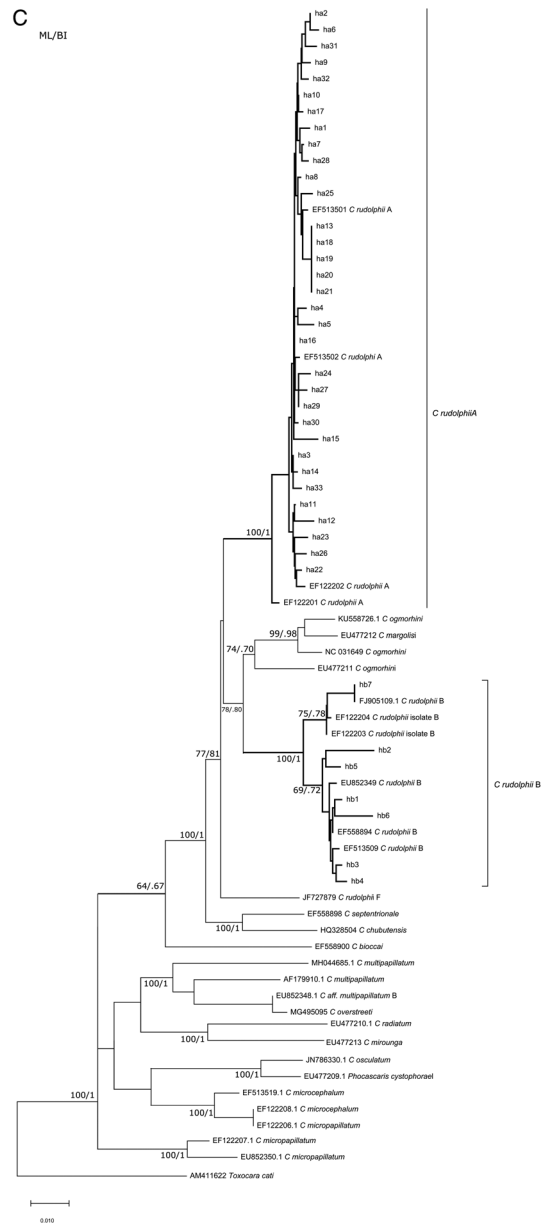


Fig. 5. Continued.

the clade containing [*C. rudolphii* complex (A, B, C, D, E and F) + *C. ogmorhini*] and appeared as the most basal species within this clade. These findings agree with results based on protein electrophoretic data (Orrechia *et al.*, 1994), 28S (Nadler *et al.*, 2000) and mitochondrial markers (Lin *et al.*, 2012). Within the obtained topologies, strong BI inferences and bootstrap values supported the position of *C. ogmorhini* (parasite of an otiariids) as sister group of the clades of avian parasites. Moreover, the *C. osculatum* complex, *P. cystophorae*, *C. radiatum* and *C. miroungae* (all from phocid hosts) appeared sharing a common ancestor with avian species namely *C. microcephalum*, *C. micropapillatum* and *C. multipapillatum*.

More sequences were available for ITS and *cox2* trees construction, and the obtained topology showed a major clade including several sister groups: (*C. rudolphii* complex + *C. variegatum* + *C. ogmorhini* + *C. eudyptulae* + *C. septentrionale*) and (*C. osculatum* complex + *P. cystophorae*, *C. radiatum* + *C. miroungae* + *C. microcephalum* + *C. micropapillatum* + *C. multipapillatum*). This topology suggests the paraphyly of the *C. rudolphii* complex and the avian *Contracecum* species, which is consistent with the

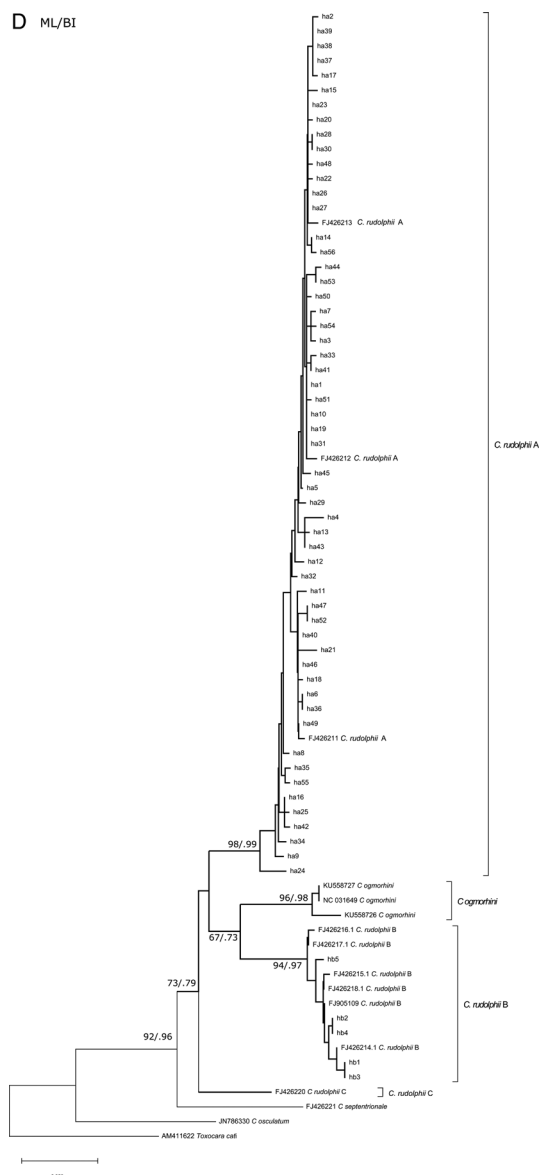


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findings inferred from the nuclear large subunit ribosomal DNA analysis (Nadler *et al.*, 2000).

In conclusion, the present results support the paraphyly of the avian *Contracaecum* species, with in addition *C. ogmorhini*. This is consistent with previous molecular studies (Nadler *et al.*, 2000; Szostakowska and Fagerholm, 2007, 2012; Farjallah *et al.*, 2008a; Dzido *et al.*, 2012; Lin *et al.*, 2012; Delgado and Garcia, 2015; Szudarek *et al.*, 2017) and allozyme data (Mattiucci *et al.*, 2008). These results were previously proposed by Fagerholm and Gibson (1987) on the basis of morphological characters, especially the similarities of caudal papillae patterns, which suggested *C. ogmorhini* as closely related to the avian *Contracaecum*, in contrast to Berland (1964), which stated avian *Contracaecum* as a monophyletic clade.

As previously suggested by Nadler (2000), these phylogenetic results, based on nuclear and mitochondrial markers, support the urgent need for a nomenclatural revision of these ascaridoid nematodes. This task requires sampling covering the maximum of species, from various hosts and localities, and the analysis of

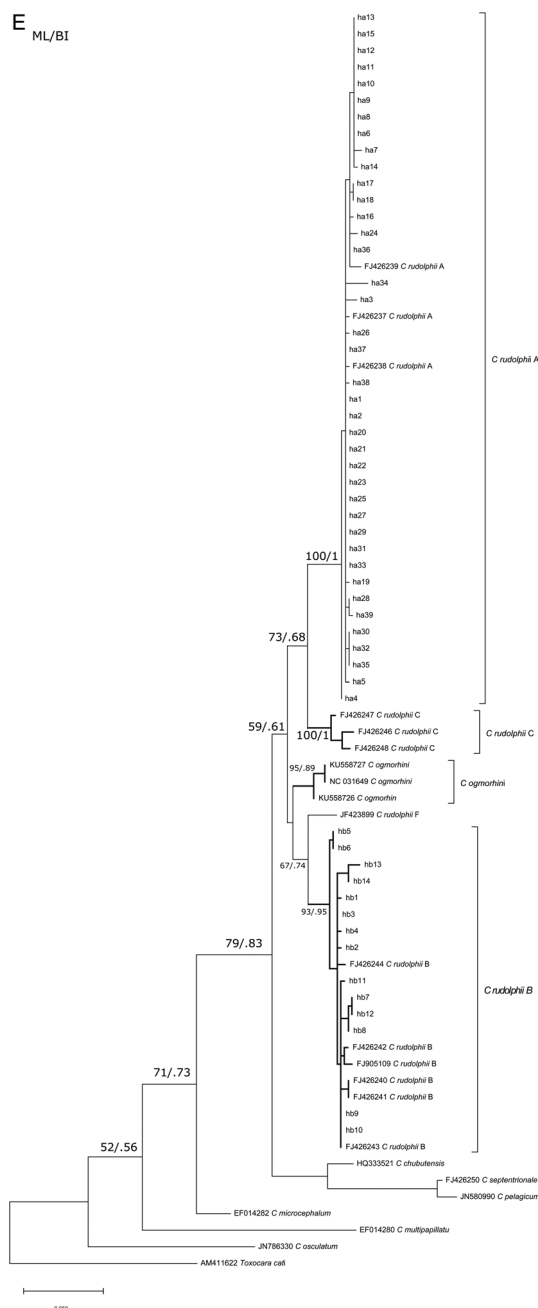


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several nuclear and mitochondrial markers that can provide a significant number of characters necessary to clarify the taxonomic status of each group and the phylogenetic relationships between them.

Supplementary material. The supplementary material for this article can be found at <https://doi.org/10.1017/S0031182020001341>

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Conflict of interest. None.

Ethical standards. None.

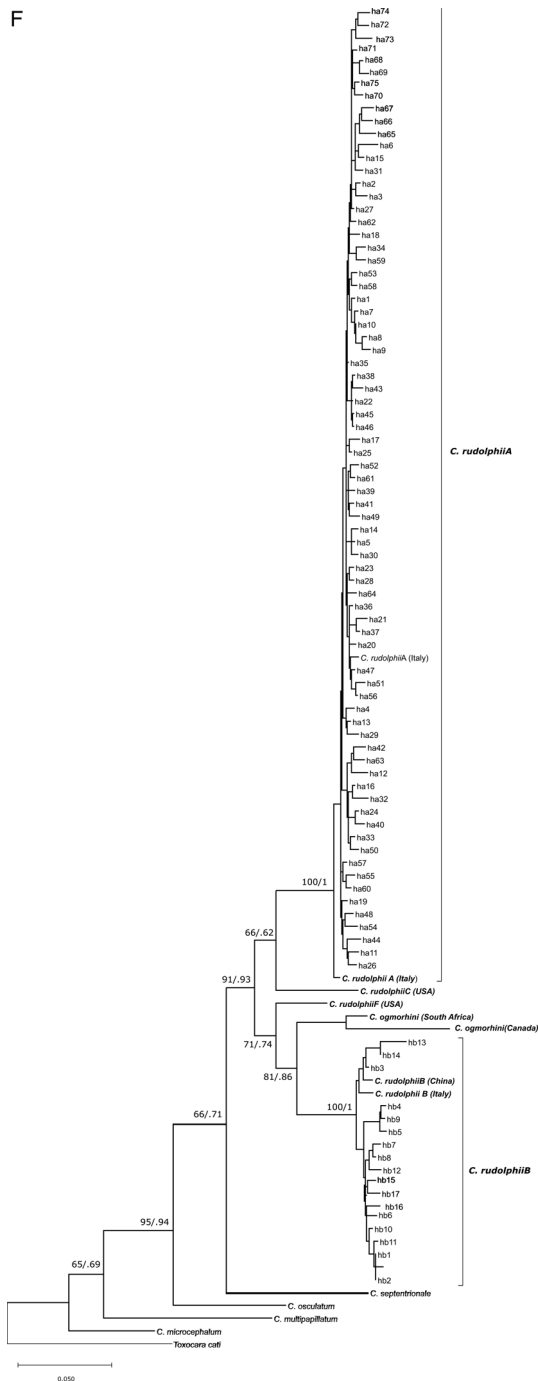


Fig. 5. Continued.

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