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# Molecular identification and phylogenetic confirmation of *Sarcocystis* species in slaughtered camels in Al-Najif province, Iraq

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#### Abstract

**Background:** *Sarcocystis* is an intracellular parasite of particular importance as it infects many domestic animals as camels that play the role of intermediate host for the parasite.

Aim: This study aimed to identify *Sarcocystis* species in camels by molecular assay with confirmation of local isolates by phylogenetic analysis.

**Methods:** A total of 200 slaughtered camels (*Camelus dromedarius*) that were slaughtered in Al-Najaf province (Iraq) abattoirs from October (2021) to July (2022) were subjected to collect the fresh tissues from four organs (esophagus, diaphragm, skeletal muscle, and heart), to be tested later by the conventional polymerase chain reaction (PCR). Then, a total of 20 positive genomic DNA samples were sequenced, named, got specific access numbers (OP785703.1 to OP785722.1), and compared with the NCBI-GenBank isolates.

**Results:** Targeting *Cox1* gene, 80% of collected tissues were found positive by the conventional PCR assay. Phylogenetic tree analysis revealed that the local *Sarcocystis* isolates were identical to Indian *S. cameli* isolates at 99.70%–99.90%. Significantly, an increase in *Sarcocystis* infection was seen in the esophagus compared to the diaphragm, skeletal muscle, and heart; older (>4 years) than younger ( $\leq$ 4 years) camels, and in females more than males.

**Conclusion:** To the best of our knowledge, this represents the first molecular study in Iraq that identifies *Sarcocystis cameli* in camels. However, additional epidemiological and molecular studies in camel populations as well as in other domestic and wild animals appeared to be necessary.

Keywords: Sarcocystosis, S. cameli, Camelus dromedarius, PCR, Sequencing.

#### Introduction

Sarcocystis, first reported in 1843, is a genus of intracellular protozoan parasite which belongs to Sarcocystidae family, Eucoccidiorina order of Apicomplexa phylum. Worldwide, considerable production losses and health illness in animals have been recognized due to the presence of >200 species in the genus of Sarcocystis that infects a wide array of domestic and wild animals (Dubey et al., 2015a; Abbas et al., 2023). The life cycle of a parasite involves two hosts; definitive (predator) and intermediate (prey) hosts. However, Sarcocystis spp. having greater specificity to intermediate than definitive hosts (Prakas et al., 2023). In the intermediate host, the development of asexual stages of the parasite occurs usually in skeletal, cardiac, esophageal, and diaphragmatic muscles and to a lesser extent in CNS (Dubey et al., 2015a; Gajadhar et al., 2015; Prakas et al., 2021). The definitive host can be infected during ingestion of contaminated tissues with mature Sarcocystis that undergoes sexual reproduction to excrete the oocysts/

sporocysts to the environment by feces (Lindsay and Dubey, 2020; Razooqi *et al.*, 2022).

One-humped or dromedary camels (*Camelus dromedarius*) reared widely in arid regions of many world countries (Burger *et al.*, 2019; Zhu *et al.*, 2019) has been used mainly for the production of meat that is characterized by the low percentages of fats and cholesterol when comparison to meats of other animal species (Faye, 2016; Kadim *et al.*, 2018). However, the existence of parasite in camel meats might potentially downgrade their quality to be consumed by humans (Hamidinejat *et al.*, 2013; Mohammadpour *et al.*, 2020).

Although *Sarcocystis* spp. can induce significant pathology or even mortality, most diseased camels generally exhibit subclinical illness (Metwally *et al.*, 2020). Thus, the diagnosis of *Sarcocystis* spp. in camels using traditional tools represents a challenge for the researchers. Hence, molecular identification of *Sarcocystis* spp. in Iraqi camels with confirmation of local isolates by phylogeny was aimed in this study.

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# Materials and Methods

#### Study animals and sampling

Totally, 200 slaughtered camels (*C. dromedarius*) of different ages (<2 to  $\leq5$  years) and sexes (107 males and 93 females) were selected to the present study from the abattoir of Al-Najaf province (Iraq) during October (2021)–July (2022). Approximately, 100 g of fresh meat were obtained from the esophagus, diaphragm, skeletal muscle, and heart of each slaughtered camel, kept in individual plastic labeled containers, and saved frozen at -20C for later molecular examination.

# Molecular examination

#### **Conventional polymerase chain reaction (PCR)**

In accordance with the manufacturer instruction of the gSYAN DNA Extraction Kit (iNtRON Biotechnology, South Korea), the DNAs were extracted from collected samples and examined directly by the Nanodrop System (Thermo Scientific, UK) to estimate the purity and concentration of these DNAs samples (Gharban et al., 2023). According to the manufacturer instructions of GoTaqTM Green PCR MasterMix Kit (Bioneer, Korea), the Mastermix tubes were prepared at a modified final volume of 25 µl (5 µl of DNA template, 2 µl of forward primer, 2 µl of reverse primer, 4 µl of Green PCR Mastermix, and 121 of nuclease-free water) using one set of primers [(F: 5'-ATG GCG TAC AAC AAT CAT AAA GAA-3') and (R: 5-ATA TCC ATA CCG CCA TTG CCC AT-3')] targeting *Cox1* gene at a product size of 1,058 bp (Metwally et al., 2020). The PCR thermocycler was carried as follows: one cycle for initial denaturation (5 minutes/95°C); 35 cycles for denaturation (30 seconds/95°C), annealing (30 seconds/54°C) and extension (1 minute/72°C); and one cycle for final extension (5 minutes/72C). Agarose gel stained with 3 µl of Ethidium Bromide and Ladder Marker (100-2,000 bp) were analyzed by electrophoresis at 80 Volt, 80 Am for 90 minutes. PCR products of previously diagnosed and confirmed Sarcocystis spp. isolate was used as a positive control; while nuclease-free water was used as a negative control.

Finally, the DNA fragments were visualized under the UV-Transilluminator (ATTA/Korea).

#### **Phylogenetic analysis**

Sequencing of some positive DNAs was performed to confirm the species of *Sarcocystis*, and study the genetic relationship between the local and NCBI-BLAST *Sarcocystis* isolates. Targeting the *Cox1* gene, a total of 20 positive DNA samples in addition to forward primer were sent for the Macrogen Company (South Korea) to be sequenced by the AB DNA Sequencing System. First, the local isolates of *Sarcocystis* species were recorded in the National Center For Biotechnology Information (NCBI)-GenBank to obtain their access numbers; and then, the Phylogenetic tree analysis was performed by the MEGA 6.0 Software.

## Statistical analysis

GraphPad Prism (*GraphPad Software Inc., USA*) Software was served to evaluate significant variation between results of this study using the one wayanalysis of variance. Values were represented as percentages (%), and significant differences were p < 0.05 (Gharban, 2023).

### Ethical approval

The current study was licensed by the Scientific Committee of the Department of Parasitology in the College of Veterinary Medicine, University of Baghdad, Baghdad, Iraq.

#### Results

Targeting Cox1 gene, 80% (160/200) of meat samples were positive by the conventional PCR assay at a product size of 1,058 bp (Fig. 1).

The genomic DNAs of 20 positive samples were named respectively as *Sarcocystis* spp. IQN.1 to IQN.20. Comparative analysis of local isolates with the NCBI-GenBank *Sarcocystis* isolates showed that there were nucleotide alignment similarities (\*) and substitution mutations in Cox1 gene.

Comparative homology sequence identity (%) to detect genetic variations between the local *Sarcocystis* spp. IQN-isolates (total no. 20) and the NCBI-BLAST *Sarcocystis cameli* Indian (MW651858.1) isolate of *cox1* gene. This identity showed that there was 99% of similarity and 0.150%–0.50% of total genetic mutations/changes (Table 1, Fig. 2).

According to organ, significantly higher increases (p < 0.05) in the prevalence of sarcocysts were detected in the esophagus (91.43%), while the lower one (p < 0.05) was seen in the heart (62.5%). However, insignificant variation (p > 0.05) in the prevalence of *Sarcocystis* spp. between diaphragm (85.42%) and skeletal muscle (85.42%) was reported (Table 2).

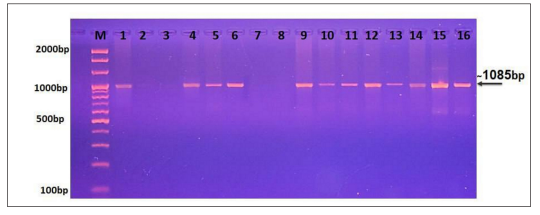
According to the age of study camels, this study has seen that the parasite is more prevalent in camels aged >4 years (96%) than those aged  $\leq$ 4 years (53.33%), (Table 3).

Regarding the sex of study camels, the infection rate of sarcocystosis was increased significantly (p < 0.05) in females (87.78%) in comparison with males (73.64%), (Table 4).

#### Discussion

*Sarcocystis* infections have been demonstrated in different animal species such as cattle, buffalo, goats, and sheep (Dakhil *et al.*, 2017; Al-Saadi *et al.*, 2020; Swar and Shnawa, 2020; Abdullah, 2021). The molecular technique is one of the most useful diagnostic methods for the identification of many pathogens, but there is low molecular data about *Sarcocystis* infection in camels. In this study, the prevalence of *Sarcocystis* spp. in slaughtered camels was 80%. This agrees with the finding reported in camels (85.5%) in Egypt (Rabie *et al.*, 2021).

In this study, the *Cox1* gene revealed a high efficacy in the detection of *Sarcocystis* spp. in tissue samples of slaughtered camels. Different studies reported that DNA barcoding system using the *Cox1* gene



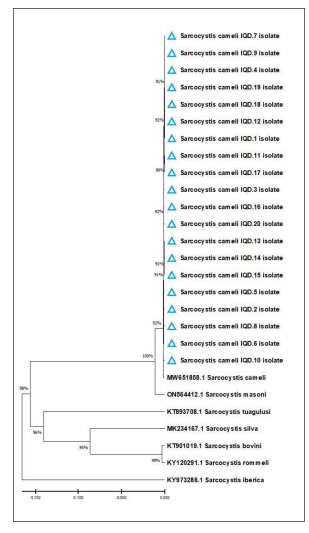
**Fig. 1.** Representative image shows the agarose-gel electrophoresis of PCR products targeting the *Cox1* gene to detect the genus of *Sarcocystis*. Lane (M): Ladder marker (100–2,000 bp); Lane (1): positive control; Lane (2): negative control; Lanes (3, 7, 8): negative samples; Lanes (4–6 and 9–16): positive samples for *Sarcocystis* spp. at 1,058.

Loca	l isolates	NCBI-GenBank isolate			
No.	Access No.	Species	Country	Access No.	Identity (%)
IQN.1	OP785703.1	S. cameli	India	MW651858.1	99.70
IQN.2	OP785704.1	S. cameli	India	MW651858.1	99.70
IQN.3	OP785705.1	S. cameli	India	MW651858.1	99.70
IQN.4	OP785706.1	S. cameli	India	MW651858.1	99.80
IQN.5	OP785707.1	S. cameli	India	MW651858.1	99.70
IQN.6	OP785708.1	S. cameli	India	MW651858.1	99.80
IQN.7	OP785709.1	S. cameli	India	MW651858.1	99.70
IQN.8	OP785710.1	S. cameli	India	MW651858.1	99.60
IQN.9	OP785711.1	S. cameli	India	MW651858.1	99.70
IQN.10	OP785712.1	S. cameli	India	MW651858.1	99.90
IQN.11	OP785713.1	S. cameli	India	MW651858.1	99.70
IQN.12	OP785714.1	S. cameli	India	MW651858.1	99.80
IQN.13	OP785715.1	S. cameli	India	MW651858.1	99.70
IQN.14	OP785716.1	S. cameli	India	MW651858.1	99.70
IQN.15	OP785717.1	S. cameli	India	MW651858.1	99.70
IQN.16	OP785718.1	S. cameli	India	MW651858.1	99.70
IQN.17	OP785719.1	S. cameli	India	MW651858.1	99.80
IQN.18	OP785720.1	S. cameli	India	MW651858.1	99.80
IQN.19	OP785721.1	S. cameli	India	MW651858.1	99.70
IQN.20	OP785722.1	S. cameli	India	MW651858.1	99.70

Table 1. NCBI-BLAST Homology sequence identity between local and GenBank isolates.

has a high efficacy in discrimination between the invertebrate and vertebrate species (Rodrigues *et al.*, 2017; Mioduchowska *et al.*, 2018; Ahmed *et al.*, 2022). Gjerde *et al.* (2015) concluded that the *Cox1* gene can work better with DNA samples derived from *Sarcocystis*, and sequences of the *Cox1* gene can

vigorously distinguish between different species of *Sarcocystis*. Rodrigues *et al.* (2017) demonstrated the suitability of *Cox1* as a gene of parasitic reorganization from those of highly related. Other studies reported that *Cox1* has a well phylogenetic character than other genes (Stöger *et al.*, 2016; Wang *et al.*, 2017). Furthermore,



**Fig. 2.** Phylogenetic tree analysis based on *Cox1* gene partial sequence in local IQN *S. cameli* isolates and NCBI-GenBank *S. cameli* isolates. Comparative identity for the genetic variations between the local isolates and NCBI-BLAST isolates showed that similarity was 98%; whereas, the total genetic mutations/changes were 0.2%–1.5%.

some authors claimed that the evolution of the *Cox1* gene enables researchers to distinguish between closely related species and investigate intraspecific variation (Changbunjong *et al.*, 2016; Viricel and Rosel, 2022). The *Cox1* has been demonstrated as the potential barcode marker for joint analysis of nuclear and mitochondrial markers of various species (Rodrigues *et al.*, 2017; Ahmed *et al.*, 2022).

Despite the discovery of five different *Sarcocystis* species in camels, multiple studies have shown that only *S. cameli* and *S. ippeni* are considered to be the real species infecting dromedary camels, namely *S. cameli*, *S. ippeni*, *S. camelicanis*, *S. camelocanis*, and *S. miescheri* (Dubey *et al.*, 2015b; Omer *et al.*, 2017; Rabie *et al.*, 2021).

# **Table 2.** Total infection rate of sarcocystosis in camels according to organ.

Orgon	Total no. –	Positives		
Organ		No.	%	
Esophagus	70	64	91.43*	
Diaphragm	48	41	85.42*	
Skeletal muscle	50	35	85.42*	
Heart	32	20	62.5	
Total	200	160	80	
Chi-square $(\chi^2)$	-	-	5.417	

\**p* < 0.043.

**Table 3.** Total infection rate of sarcocystosis in camels according to age.

	Total no	Positives		
Age (year)	Total II0. –	No.	%	
≤4	75	40	53.33	
>4	125	120	96*	
Total	200	160	80	
Chi-square $(\chi^2)$	-	-	4.425	

\*Vertical comparison between values (%) refers to significant differences at (p < 0.05).

**Table 4.** Total infection rate of sarcocystosis in camels according to sex.

Sex	Total no. –	Positives		
Sex	10tal II0. –	No.	%	
Male	110	81	73.64	
Female	90	79	87.78*	
Total	200	160	80	
Chi-square $(\chi^2)$	-	-	4.009	

\*Vertical comparison between values (%) refers to significant differences at (p < 0.05).

In this study, the phylogenetic analysis discovered that the local study isolates were identical to the global NCBI-GenBank Indian *S. cameli* isolate. This is in agreement with the results of other studies as *S. cameli* is the more prevalent species of *Sarcocystis* in camels to yet (Valinezhad *et al.*, 2008; Dubey *et al.*, 2015b; Omer *et al.*, 2017; Asopa *et al.*, 2023). For the first molecular proof of *S. cameli* in camels in Iran (Motamedi *et al.*, 2011), the bradyzoite DNA was amplified from the *18S rRNA* gene fragment using traditional PCR.

Sarcocystis species can frequently infect the muscular tissue of the heart, tongue, esophagus, and diaphragm (Wahba *et al.*, 2014; Ahmed *et al.*, 2016). Our findings revealed that *Sarcocystis* spp. has existed in all examined organs but more commonly in the

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esophagus. Some research suggested that sarcocystosis is more frequent in the cremaster muscle (Bucca et al., 2011; Sağlam and Keleş, 2016). Shekarforoush et al. (2006) demonstrated that the heart is the most infected organ; while Al-Ani and Amr (2017) noted that camel's diaphragm was the most frequently affected organ. According to Oryan et al. (2010), Sarcocystis appears to prefer the esophagus, tongue, and heart. In contrast, Gareh et al. (2020) reported that Sarcocystis can spread in various organs, mostly in the esophagus with a prevalence rate of 49%. Rabie et al. (2021) reported Sarcocystis in the esophageal, heart, and ocular muscles in addition to other organs, with a higher infection rate in the esophagus (85.5%). Asopa et al. (2023) showed that Sarcocystis spp. was prevalent in the tongue and esophagus of dromedary camels from Rajasthan's Bikaner area. This variance in the distribution of S. cameli strains may be accounted for differences in the prevalence of Sarcocystis among camel organs (Hamidinejat et al., 2013).

Another substantial risk factor for infection was age. In this study, the infection rate was considerably higher in older camels (>4 years) than in younger camels (<4 years). Other reports in Iran (Hamidinejat *et al.*, 2013), Saudi Arabia (Omer *et al.*, 2017), and Egypt (El-Bahy *et al.*, 2019) have revealed similar results. The slower development of detectable cysts may account for the low prevalence of *Sarcocystis* in young camels. Another explanation for increasing the rate of infection in older camels is that the older animals are slaughtered at a higher rate than younger ones. In addition, keeping young camels indoors may reduce their exposure to disease (Valinezhad, *et al.*, 2008, Hamidinejat *et al.*, 2013, Omer *et al.*, 2017).

Sex was discovered as a crucial factor linked to infection. Our findings were consistent with numerous studies conducted in southern Ethiopia (Woldemeskel and Gumi, 2001), Iran (Valinezhad et al., 2008), and Egypt (Rabie et al., 2021) which found that male camels were more susceptible to infection than females. Omer et al. (2017) showed that the slaughtered female camels were usually over 4 years old, whereas male camels were under 2 years old. This disparity might be explained by the fact that most males are grazed outside for hard work, while most females are kept indoors for breeding under good and hygienic management. This might make males more susceptible to infection than females (Romero et al., 2017). An insignificant difference in the frequency of sarcocystosis between male and female camels was identified by Hamidineiat et al. (2013). A lack of relationship between sex and infection rates has been shown in similar studies on camels (Woldemeskel and Gumi, 2001; Shekarforoush et al., 2006; Valinezhad et al., 2008).

# Conclusion

To the best of our knowledge, this represents the first molecular study in Iraq that identifies *Sarcocystis* 

*cameli* in camels. However, additional epidemiological and molecular studies in camel populations as well as in other domestic and wild animals appeared to be necessary.

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# **Conflict of interest**

The authors declare that there is no conflict of interest. *Author's contribution* 

OAA: Collection of tissue samples, extraction of DNAs, and preparation of Mastermix tubes; MTSA: PCR analysis, sequencing, and statistical analysis of obtained results. All authors have written the manuscript and approved the final copy of it.

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# Data availability

All data supporting the findings of this study are available within the manuscript.

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