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Source: Journal of Wildlife Diseases, 34(2) : 317-324

Published By: Wildlife Disease Association

URL: https://doi.org/10.7589/0090-3558-34.2.317
GASTRIC SPIRAL BACTERIA AND INTRAMUSCULAR SARCOCYSTS
IN AFRICAN LIONS FROM NAMIBIA

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ABSTRACT: Six juvenile male, one adult male, and three adult female African lions (Panthera leo) from Etosha National Park, Republic of Namibia were presented for necropsy. Two of four adults and one of six juveniles had moderate numbers of gastric spiral bacteria. Additionally, four of four adults had sarcocysts. All juveniles had enteric Sarcocystis sp. oocysts, but no sarcocysts. The gastric spiral bacteria were located extracellularly in fundic and pyloric glands, and also apparently intracellularly within parietal cells in the fundic region. The organisms were 4 to 8 μm long, 0.63 μm wide, with a periodicity of 0.60 μm. The bacteria had blunt ends with multiple flagella. No periplasmic fibrils were observed. The histologic and ultrastructural characteristics of the bacteria were considered most consistent with species in the genus Helicobacter or incompletely identified Helicobacter-like organisms. Gastric inflammation did not differ significantly between infected and uninfected individuals. The bacteria may be commensals, or an opportunistic pathogen. The sarcocysts were observed in hindlimb skeletal muscle of four individuals, with one individual also containing a single sarcocyst within glossal musculature. All observed cysts were mature, and were contained within individual myocytes. The cyst wall consisted of a 44 to 66 nm, granular, electron dense parasitophorous membrane with subjacent, 0.8 to 1.3 μm thick, granular and fibrillar ground substance which also extended into the cyst interior as thin septa. The membrane was folded and lined irregularly spaced, 0.8 to 1.3 μm tall villi centrally containing ground substance. The membrane was continuous in the villar projections, but divided into discrete aggregations of the electron dense material between the villi. Bradyzoites within the interior of the cyst were 3 by 12 μm. The sarcocysts were determined to be Sarcocystis felis based on the characteristic ultrastructural appearance of the cyst wall.

Key words: African lion, gastric spiral bacteria, Helicobacter sp./Helicobacter-like organism, Panthera leo, Sarcocystis felis, sarcocysts.

INTRODUCTION

Gastric spiral bacteria of the genus Helicobacter and incompletely identified Helicobacter-like organisms (HLO) are Gram negative, tightly coiled, 4 to 10 μm long organisms with bipolar flagella. They have been described in many host species, including man, non-human primates, ferrets (Mustela putorius), pigs (Suis scrofa), domestic cats (Felis domesticus), cheetahs (Acinonyx jubatus), a cougar (Felis concolor) (Lee et al., 1988; Lee and O’Rourke, 1993; Eaton et al., 1993a; Hill et al., 1997), and various species of small felids (M. J. Kinsel, unpublished data). Most HLO are assumed to be closely related to the type species from man alternately as Gastrospirillum hominis or Helicobacter heilmanii (Solnick et al., 1993). Phylogenetic characterization of two human derived organisms indicated they were closely related to Helicobacter felis (Solnick et al., 1993), and presumably HLO observed in animals also are related to Helicobacter sp. Definitive identification of HLO is hampered by their general failure to grow in vitro, although recent data suggest that in vitro culture is possible if optimal conditions are met (Hanninen et al., 1996). Other procedures which may prove useful for identifying Helicobacter spp. or HLO in biopsy or necropsy samples include rapid urease tests, and various molecular biology techniques including in situ hybridization, and the polymerase chain reaction (PCR) (Van den Berg et al., 1989; Valentine et al., 1991; Megraud, 1996).

Both Helicobacter spp. and HLO have been observed in domestic and captive felids with gastric inflammation (Eaton et al., 1993a; Geyer et al., 1993; Lee and
O’Rourke 1993; Hermanns et al., 1995; Hill et al., 1997), and positive correlations between gastric spiral bacterial infection and gastritis have been reported (Eaton et al., 1993a, b; Hermanns et al., 1995; Happonen et al., 1996). Additionally, the capacity of related organisms, particularly Helicobacter pylori in humans, to incite clinically significant gastric pathology ranging from gastritis to gastric neoplasia is well documented (Dubois, 1995). However, definitive association of gastritis in felids with HLO remains tenuous as controlled pathogenesis studies are lacking, and HLO can be present without gastritis.

To date, study of gastric spiral bacteria has been restricted to domestic, laboratory, and captive species. The incidence and health implications of gastric spiral bacterial infection in free-ranging animal populations remains unknown. However, a widespread distribution similar to that observed in domestic and captive species would seem likely.

Conversely, Sarcocystis spp. are a group of Apicomplexan parasites whose worldwide distribution and ability to induce clinically significant disease in domestic, captive, and wild animals are well documented (Fowler, 1986; Dubey et al., 1989; Hullard, 1993). These organisms have a complex life cycle involving sexual reproduction in the intestinal tract of a definitive host, and asexual reproduction culminating in sarcocysts in intermediate hosts. Usually the intermediate host is a prey species, typically a bird or herbivore, while definitive hosts are generally carnivores. However, carnivores can occasionally act as intermediate hosts in the sarcocystis life cycle (Dubey et al., 1989). Presumably the definitive host in this scenario would be a species that at least intermittently engages in scavenging behavior, although no determinations have been made (Greiner et al., 1989). Sarcocysts consistent with, or identified as, Sarcocystis felis have been observed in domestic cats (Felis rufus) (Anderson et al., 1992; Dubey et al., 1992), cougars and Florida panthers (Felis concolor coryi) (Greiner et al., 1989), and one African lion (Panthera leo) (Dubey and Bwangamoi, 1994). The definitive host for this species has not been determined.

The present study used descriptive methods to document the presence of gastric spiral bacteria and sarcocysts in free-ranging lions from Etosha National Park, Republic of Namibia. Ancillary procedures, such as bacterial culture, rapid urease tests, fecal floatations, and molecular biology methods were not performed due to constraints of field conditions.

MATERIALS AND METHODS

Six 1-17-old male, one 4-17-old male, one 7-17-old female, and two 14-17-old female African lions from Etosha National Park, Republic of Namibia (19°25’S, 15°45’E) were presented for postmortem examination. Field necropsy procedures were performed 12 to 18 hr postmortem.

Routine, representative samples of all organ systems were collected for histopathology. Tissues were fixed in 10% neutral buffered formalin, processed routinely, embedded in paraffin, sectioned at 5 μm, and stained with hematoxylin and eosin (H&E). Sections of stomach were also prepared with a modified Steiner silver impregnation (Chrunikian and Garvey, 1990).

Samples for transmission electron microscopy were identified from H&E slides and removed from paraffin blocks. Tissues were deparaffinized, post-fixed in osmium tetroxide, and embedded in epoxy (Lx112, Ladd Research Industries, Inc., Burlington, Vermont, USA). Thick sections for light microscopy were cut at 0.5 μm and stained with toluidine blue and basic fuchsin (Hoffmann et al., 1983). Thin sections were cut at 90 nm, stained with lead citrate and uranyl acetate, and examined with a Hitachi H-600 transmission electron microscope (Hitachi, Ltd., Tokyo, Japan) operated at 75 KV.

In the following discussion measured values of organisms and structures are followed by the mean and standard deviation in parentheses.

RESULTS

At necropsy, all carcasses were moderately autolyzed. Gross lesions were limited to gunshot trauma and macroscopic evi-
sisted of mild to occasionally moderate, multifocal accumulations of lymphocytes and plasma cells in the deep lamina propria. The female also had minimal, diffuse, eosinophil infiltrate in the deep lamina propria, and the male had few, moderate to markedly reactive lymphoid follicles in the pyloric region. Similar inflammation was observed in uninfected adults.

In the juvenile lion, bacteria were most prominent in the pyloric region, and were observed extracellularly in the crypts and mid-zonal aspects of the gastric glands. In the fundus bacteria were rarely observed extracellularly in the apical aspects of the gastric glands and also apparently intracellularly within parietal cells. Gastric inflammation in this individual was limited to occasional, mild to moderate, mid-zonal lymphoplasmacytic aggregations in the pylorus, and few, mildly hyperplastic lymphoid follicles in the fundic region. Uninfected juveniles had similar gastric inflammation.

Ultrastructural evaluation of affected pyloric stomach revealed numerous extracellular spiral bacteria. The bacteria were 0.53 to 0.73 μm (0.63, 0.05) wide, with a periodicity of 0.50 to 0.67 μm (0.59, 0.06), with blunted ends, and multiple polar flagella. Periplasmic fibrils were not observed (Fig. 2).

Hematoxylin and eosin sections of hind-limb skeletal muscle in four of four adult lions contained rare to moderate numbers of cysts consistent with Sarcocystis sp. One female had a single cyst in glossal musculature. The sarcocysts were 40 by 70 μm to 50 by 260 μm, round to ellipsoid, and were located within individual myofibers (Fig. 3). There was no inflammatory reaction associated with these sarcocysts, although one individual did have rare necrotic myocytes with minimal to mild mononuclear inflammation distant to observed sarcocysts. Interestingly, none of the juvenile lions had sarcocysts, but all had large numbers of Sarcocystis sp. oocysts within small intestinal lamina propria macrophages. Identification of the oocysts to species was not possible.
Ultrastructurally, the sarcocysts were mature with a well developed cyst wall (Fig. 4). The cyst wall consisted of a parasitophorous vacuolar membrane with subjacent ground substance. The membrane was comprised of electron dense, granular material 44 to 66 nm (56, 11) thick folded into 0.8 to 1.3 μm (1.1, 0.2) tall, irregularly spaced villar projections. The membrane was continuous in the villar projections, but was divided into discrete aggregations of the granular material between the villi. Underlying the membrane was a 0.9 to 2.0 μm (1.3, 0.4) thick layer of ground substance comprised of finely granular and fibrillar material which also occupied the central aspect of the villous projections and occasionally extended deep into the interior of the cyst as thin septa. The cyst wall occasionally had small invaginations at the origins of the septa. The cyst contained numerous, approximately 3 by 12 μm bradyzoites but no metrocytes. The ultrastructural appearance of the sarcocysts was consistent with that of *S. felis* (Dubey et al., 1992).

**DISCUSSION**

Gastric spiral bacteria have been observed in many species including domestic cats and captive felids (Lee et al., 1988; Eaton et al., 1993a; Lee and O’Rourke, 1993; Hill et al., 1997), but the presence or incidence of this class of organism in wild animal populations has not been re-
reported. This is the first report of gastric spiral bacteria in a free-ranging species.

Gastric spiral bacteria have been associated with gastritis in felids, with both *Helicobacter* spp. and HLO implicated (Eaton et al., 1993a; Hermanns et al., 1995; Happonen et al., 1996). Mixed *Helicobacter* sp. and HLO infections also have been observed in felids with gastritis (Eaton et al., 1993a; Happonen et al., 1996). However, positive correlations between gastritis and HLO or *Helicobacter* sp. infection are observed only variably in felids (Eaton, et al., 1993a, b; Geyer et al., 1993; Hermanns et al., 1995), and confusion about what level of gastric inflammation constitutes gastritis in felids also complicates interpretation (Happonen et al., 1996). Further association between HLO obtained from felids and their ability to induce gastritis is limited to passage of the organisms into laboratory animals, with resultant gastritis (Eaton et al., 1995). Thus the role of HLO in the development of gastritis in felids remains unclear.

In the lions in this report, gastric spiral bacteria were observed in 3 of 10 individuals, one juvenile, and two adults. All lions examined had similar gastric inflammation regardless of the presence or absence of spiral bacteria. The spiral bacteria may have a commensal relationship with the lion, or may be an opportunistic pathogen. Commensal or opportunistic pathogenic associations between HLO and host species has been suggested previously (Fox and Lee, 1989). Alternatively, the spiral organisms may cause gastritis in only a limited number of infected individuals, which has been observed in other species (Doolay, 1993; Lee and O’Rourke, 1993; Dubois, 1995).

The gastric spiral bacteria in this report are considered morphologically compatible with HLO. Organism length, tight coiling, polar flagellar tufts, and lack of periplasmic fibrils are consistent with previous reports of HLO morphology. Additionally, an affinity for parietal cells has also been reported for HLO (Dubois et al., 1991; Lee and O’Rourke, 1993), and spiral bacteria were occasionally observed apparently within parietal cells in the affected lions.

The possibility of a greater prevalence of infection in this group of lions does exist, as other testing procedures often used for the detection and identification of gastric *Helicobacter* sp. or HLO in support of histologic findings were not utilized in this study. Rapid urease tests, bacterial culture, and PCR can be used to elucidate on the presence of gastric spiral bacteria even in the face of negative histology due to patchy distribution or low numbers of bacteria. Urease tests and bacterial culture were impractical given field conditions, and PCR is not widely available (Jenkins and Bassett, 1997). Additionally, histology remains a highly sensitive test with adequate sample size, and is the only method by which gastritis can be diagnosed (Jenkins and Bassett, 1997).
Sarcocysts have been previously described in a variety of felids including one African lion (Dubey and Bwangamoi, 1994). Immunosuppression with abnormal or aberrant infection was initially suggested as a cause for sarcocysts in felids (Kirkpatrick et al., 1986; Edwards et al., 1988) but it is now thought that some Sarcocystis sp. use carnivores as intermediate hosts in the course of their normal life cycle. The life cycle and transmission of most of these Sarcocystis sp. have yet to be determined; however, scavenger species are assumed to be the definitive hosts (Greiner et al., 1989). Sarcocystis felis is thought to be one of the species utilizing carnivore intermediate hosts.

Some Sarcocystis spp. can be identified with relative certainty based on ultrastructural characteristics of the cyst wall, including S. felis (Dubey et al., 1992). Previous reports where sarcocysts in felids have been described ultrastructurally are all consistent with S. felis, which suggests a worldwide distribution. The sarcocysts observed in the lions in this report also were consistent with S. felis as determined by ultrastructural architecture of the cysts. Sarcocysts from a springbok (Antidorcas marsupialis), a common prey species of the lions in Etosha, were ultrastructurally distinct from the lion sarcocysts (M. J. Kinzel, unpublished data). All lions had markedly reactive lymphoid tissue, suggesting normal immune function, and aberrant migration of a Sarcocystis sp. secondary to

Figure 4. Ultrastructural architecture of Sarcocystis felis cyst wall from an African lion. Parasitophorous membrane (p) is folded into irregularly spaced villi and consists of discrete aggregates of the electron dense material between villi (arrowhead). Ground substance (g) extends into cyst interior as septa (s). Bradyzoites (b) are present in the cyst interior. Uranyl acetate and lead citrate. Bar = 1 μm.
immunosuppression was considered unlikely. The *S. felis* sarcocysts were observed in all adult lions, but not in juveniles, implying infection is acquired in maturity, and is widespread in the adult population. Additionally, sarcocysts distinct from *S. felis* have been observed in domestic dogs (*Canis familiaris*), African wild dogs (*Lycaon pictus*), and black-backed jackals (*Canis mesomelas*) in areas of Africa where they have been found also in lions (Bwangaomi et al., 1993; Wesemeier et al., 1995). This may imply that utilization of carnivores as intermediate hosts by *Sarcocystis* spp. is more prevalent than previously suspected.

In summary, gastric spiral bacteria consistent with HLO were observed in 30% of lions examined. To the authors’ knowledge this is the first report of gastric spiral bacteria in a free-ranging wild species. No correlation between HLO infection and gastric inflammation could be made. Additionally, sarcocysts consistent with *S. felis* were observed in 40% of the lions examined and 100% of the adults. These findings suggest infection is strongly associated with maturity and common in the adult population. This is the second report of *S. felis* in African lions.

**ACKNOWLEDGMENTS**

The authors acknowledge the excellent technical assistance of the histopathology and electron microscopy sections of the University of Illinois Laboratories of Veterinary Diagnostic Medicine, and the Ministry of Environment and Tourism, Republic of Namibia. Partial funding for this project was obtained from the Chicago Zoological Society and the University of Illinois College of Veterinary Medicine.

**LITERATURE CITED**


Received for publication 2 May 1997