Unique case of disseminated toxoplasmosis and concurrent hepatic capillariasis in a ring-tailed lemur: first case description

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Received: 30 January 2015 – Revised: 2 April 2015 – Accepted: 8 April 2015 – Published: 24 April 2015

Abstract. A unique co-infection with Toxoplasma gondii and Capillaria hepatica is reported in a semi-free-living ring-tailed lemur (Lemur catta). In this case acute toxoplasmosis, characterised by severe necrotising inflammation in different organs, was the leading cause of death, whilst accompanying chronic capillariasis was regarded as a predisposing factor. A concise description of both disease manifestations is given.

1 Introduction

Toxoplasma (T.) gondii, a coccidian parasite with worldwide distribution, can infect nearly all warm-blooded vertebrate species (Tenter et al., 2000). Although infection may be common in many mammalian species, including several nonhuman primate (NHP) species, clinical disease is rare. Interestingly, some species are highly susceptible to infection and often develop fatal toxoplasmosis. These include New World monkeys, prosimians, mountain hares, and Australian marsupials (Dubey et al., 1985; Gustafsson et al., 1997; Juan-Salles et al., 1997). Clinical signs of disease depend on the infected species and the affected organ systems. Highly susceptible species often acquire overwhelming infection and die peracutely without premonitory signs (Dietz et al., 1997). Among primates, lethal toxoplasmosis has been reported mostly in New World monkey species and lemurs; thus both of them seem to be highly susceptible to the disease (Epiphanio et al., 2003; Yabsley et al., 2007).

Hepatic capillariasis is caused by the parasitic metazoan nematode Capillaria (C.) hepatica that infects a broad range of primate species (Stidworthy et al., 2009). Despite the fact that deaths due to heavy parasitic burden have been reported, hepatic capillariasis is a mainly silent condition, presenting no actual symptoms, mostly diagnosed post mortem (Pizzi et al., 2008). In this case report, an interesting combination of disseminated toxoplasmosis with C. hepatica co-infection in a ring-tailed lemur is presented.

2 Case history

An adult male ring-tailed lemur (Lemur catta) from a zoo-logical garden (with access to both indoor and outdoor enclosures) was found dead without any overt signs of disease prior to death. The animal was subsequently submitted to the German Primate Center for necropsy and further investigation. Prominent gross pathologic lesions were found in the small intestine and the mesenteric lymph nodes. Within the jejunum, multifocal ulcerations with hyperaemic margins were present, and the mesenteric lymph nodes were enlarged.

3 Histology

The main histologic findings focused on the intestine, the liver, and the lymphatic system. The small intestine presented segmental intestinal necrosis and multifocal, transmural, ulcerative jejunitis. Ulcers were observed only in the je-
Figure 1. Transmural, ulcerative enteritis, jejunum, ring-tailed lemur (Lemur catta). Transmural ulcer characterised by amorphous, eosinophilic material comprising of cellular debris originating from mucosa, submucosa, and muscularis areas together with inflammatory cell relics. Composition of several low-power (magnification: ×5) photomicrographs. Haematoxylin and eosin (H&E).

Figure 2. Parasitic, necrotising hepatitis, liver, ring-tailed lemur (Lemur catta). Ovoid structures (sectioned at different planes) represent C. hepatica unembryonated bipolar eggs accompanied by multiple foci of hepatocellular necrosis and mild inflammatory cell infiltrate. Magnification: ×10. H&E.

Figure 3. Parasitic, necrotising hepatitis, liver, ring-tailed lemur (Lemur catta). Necrotic focus circumscribed by inflammatory cell infiltrate. A single toxoplasmic tissue cyst is present on the boundaries between healthy and necrotised tissue (arrow). Magnification: ×20. H&E.

Figure 4. Parasitic, necrotising hepatitis, liver, ring-tailed lemur (Lemur catta). Extracellular T. gondii tissue cyst (arrows) inducing inflammatory response and hepatocellular necrosis. Magnification: ×40. H&E.

The spleen and mesenteric lymph nodes were highly lymphocyte-depleted and diffusely necrotic. Splenic white pulp showed neither follicles nor perivascular lymphatic sheaths; cortical follicular architecture in the lymph nodes was also effaced. Furthermore, marked necrosis extended into the splenic red pulp and the medulla of the lymph nodes, while T. gondii cysts were frequently observed within the lymphoid-depleted areas. Finally, the central nervous system featured no signs of meningoencephalitis.
4 Immunohistochemistry

In order to prove the presence of the protozoan, associate it with the lesions, and to study its potential distribution among other organs, a *T. gondii*-specific immunohistochemistry (IHC) was performed (Figs. 5–6). In more detail, a commercial mouse monoclonal primary antibody against *T. gondii* P30 membrane protein was used (Novocastra, NCL-TG) in a 1:20 dilution. A biotinylated secondary antibody, streptavidin, and the colour indicator (alkaline phosphatase red detection kit; Ventana, Cat. 760-031) were applied according to the supplier’s instructions. Two negative controls were performed. The first included replacement of the primary antibody with sheep serum; the second utilised tissue sections from a healthy animal. Infected tissue served as a positive control. For counterstaining Mayer’s haematoxylin was used.

Liver, spleen, lymph node, intestine, pancreas, and lung specimens presented positive signals revealing both the presence of protozoan cysts and/or tachyzoites. While no lesions were noticed in routine histological evaluation, adrenal gland, myocardium, and prostate gland presented minor but nevertheless typical IHC staining consistent with tachyzoites.

5 Discussion

The present case confirms the high susceptibility of lemurs to *T. gondii* infection. There have been sporadic reports of deaths in captive ring-tailed lemurs induced by toxoplasmosis (Spencer et al., 2004). In lemurs, and New World monkeys toxoplasmosis leads to acute disease, illness, and death of both host and parasite (synnecrosis), in contrast to Old World monkeys and great apes, which are only minimally affected (parasitism) (Innes, 1997). Attempting to offer an explanation for this difference, some authors state evolutionary reasons. Evolutionary differences in habit and habitat between the definitive hosts and New World monkeys are mentioned as the main reasons that precluded the latter from developing protective immune responses (Carme et al., 2009; Dietz et al., 1997; Epiphanio et al., 2003; Innes, 1997; Yabbsley et al., 2007). According to this hypothesis, the extreme susceptibility may be attributed to the arboreal habits of some New World monkey species and prosimians. Arboreal monkeys are not exposed to oocysts shed with felid faeces to the same extent terrestrial species are. Since parasitism is an evolutionary phenomenon (Epiphanio et al., 2003), arboreal species may have thus failed to develop resistance to toxoplasmosis, in contrast to their non-arboreal counterparts (Carme et al., 2009; Innes, 1997).

Most of the pathomorphologic findings of the present case regarding toxoplasmosis confirmed those already reported for New World monkeys (Brack et al., 1995; Epiphanio et al., 2003). Often there are no gross lesions at necropsy. Pulmonary oedema, congestion, and consolidation may be the only significant necropsy findings. Microscopically, necrotising lesions within liver, lung and heart, accompanied by nonsuppurative meningoencephalitis, are the predominant histologic findings. The distribution of *T. gondii* organisms in the tissue is variable. Bradyzoites, tachyzoites, and cysts are often widely disseminated, as shown in the present case by immunohistochemistry. Diagnosis is based on demonstration of these organisms in tissue samples by light and electron microscopy. *T. gondii* tachyzoites are 2–6 µm long, crescent shaped, while cysts are 5–100 µm round thin-walled structures containing few to several hundred bradyzoites (Dubey et al, 1998).

In the present case, diagnosis was confirmed by specific immunohistochemical staining. Immunohistochemical labelling for *T. gondii* antigen was positive in samples of small intestine, liver, spleen, and mesenteric lymph nodes, suggesting the gut as the route of entry for this infection. Furthermore, several other tissues were found to be immunohistochemical positive, which is indicative of systemic infection, despite the variations in the intensity of labelling.
Acute phase toxoplasmosis features random, multifocal, necrotising hepatitis, directly related to the rapid proliferation of tachyzoites. Chronic capillaritis, which was also diagnosed in this case, provokes hepatocellular necrosis and mild granulomatous response, while granulomatous hepatitis has been recorded only in more severe cases (Pizzi et al., 2009). In this case however, acute disseminated toxoplasmosis is considered to be the actual cause of death. Histological and immunohistochemical analyses emphasise the role of T. gondii in the pathogenesis of the aforementioned parasitic hepatitis. Even though the current C. hepatica burden is considered too low to be lethal, its contribution requires further investigation. The liver alterations induced by C. hepatica may have induced a higher susceptibility for the manifestation of T. gondii infection within the organ, leading to severe liver pathology and subsequent spread of the agent to other organs. Based on our own observations in similar cases, concomitant liver hemosiderosis may also act – through the iron overload – as a major predisposing factor for disease development in susceptible species. As suggested by our findings, the liver plays a key role in the pathogenesis, clinical presentation, and pathology of systemic toxoplasmosis. Therefore, there are concerns regarding the substantial influence of chronic subclinical liver alterations in the outcome of the infection.

While the source of the T. gondii agents in this case remains undetermined, the most likely scenario is an oral route of infection. Several alterations within the small intestine induced by T. gondii support this assumption. Felids are the only known definitive hosts for T. gondii, and ingested oocysts shed by cats are the source of infection for several mammalian species acting as intermediate hosts. The role of nondomestic felids in the epidemiology is unknown. Free-roaming house cats defecating near or within the enclosures could serve as a potential source of infection, as they might contaminate stored wood chips intended for bedding materials, or fruits and vegetables prepared for feeding. If possible, it is advisable to keep an appropriate distance between the housings of felids and susceptible species. A strict hygiene practice and a feeding programme without raw meat can further reduce the risk of infection.

Finally, this is the first report of T. gondii and C. hepatica co-infection in a ring-tailed lemur to our knowledge, underscoring this species’ vulnerability to similar infections, along with the need of strict surveillance to avoid similar devastating conditions.

Acknowledgements. We are grateful to W. Henkel, E. Lischka, S. Wienstroth, N. Schminke, and L. Hummel for excellent technical assistance.

The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

The authors received no financial support for the research, authorship, and/or publication of this article.

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