Molecular evidence for *Mycobacterium bovis* infection in wild Chilean hawk (*Accipiter chilensis*)

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**ABSTRACT.** The present study reports a case of tuberculosis in a juvenile Chilean hawk (*Accipiter chilensis*). Granulomas were identified in the liver, intestine and mesentery. Several of these granulomas contained acid-fast bacilli suggesting a mycobacterial infection through ingestion. Molecular diagnosis detected *Mycobacterium bovis* as etiological agent. Whether *M. bovis* infection in this Chilean hawk represents a risk for other wildlife, livestock or humans remains to be determined.

*Key words:* Chilean hawk, infection, *Mycobacterium bovis*, spillover.

*Mycobacterium bovis* (*M. bovis*), a slow-growing and an obligate intracellular bacterium, produces a chronic infection principally of the respiratory system (Michel et al 2010) affecting most domestic mammals, wild species and humans (Neill et al 2005), and is transmitted via air but also through the digestive tract (Kaneene and Pfeiffer 2006). An important aspect is that *M. bovis* is considered a major zoonotic agent, associated with human tuberculosis (Neill et al 2005).

Transmission of *M. bovis* from domestic animals to wildlife (spillover) and vice versa (spillback) has been facilitated by anthropogenic factors, which have been considered in *M. bovis* infection eradication programs worldwide (Nugent et al 2012).

Avian mycobacterial infection is an ubiquitous disease in domestic, captive, and wild birds caused by *M. avium* complex and *M. genavense*. Diseases of birds of prey kept in captivity as well as free-living raptors are well documented (Millán et al 2010, Vidal et al 2017). Serological studies show that free-living birds are exposed to pathogens through contact with poultry waste and run-off from farms, or by ingestion of contaminated carcasses (Höfle et al 2002, Cunha et al 2017), first being susceptible hosts, thereafter potential sources of the infection. The aim of the present study was to provide evidence of *M. bovis* infection in a wild Chilean hawk specimen found dead in southern Chile, where the major part of the Chilean livestock production system is situated.

According to phenotypic features (Pavez 2004), a juvenile *Accipiter chilensis* (order Accipitriformes), commonly known as Chilean hawk, was found dead in a rural area near the city of Valdivia, Chile (39° 48’ S, 73° 15’ W). The animal was submitted to Universidad Austral de Chile for further analysis. A complete necropsy was performed. The bird showed little subcutaneous and intracoelomic fat. Multiple nodules of approximately 2-5 mm diameter, whitish in color and of firm consistency, were observed on the surface and inside the liver parenchyma, as well as in the mesentery and wall of the small intestine (figure 1A). Multiple sections of small intestine and liver were fixed in 10% neutral buffered formalin and embedded in paraffin, sectioned at 5 µm, and stained with hematoxylin and eosin (HE) and Ziehl-Neelsen (ZN) stain. The liver and the intestinal submucosa showed multiple granulomas with central necrosis, surrounded by histiocytic infiltrate with presence of multiple giant cells and few heterophils (figure 1B). The ZN stain showed large numbers of red acid-fast bacilli (AFB) on the inside and in the periphery of the granulomas (figure 1C). Two paraffin blocks containing hepatic tissue and tissue from the small intestine wall were selected for molecular confirmation. To prevent cross contamination of the samples for PCR, the microtome and each paraffin block were disinfected with a 10% chlorine solution and thereafter with 100% ethanol. Ten to twenty sections measuring 5 µm were cut and transferred into a sterile 1.5 mL tube for paraffin removal (Miller et al 1997). After the dewaxing process of the embedded specimen, the samples were subjected to DNA extraction, PCR confirmation and molecular typing. The High Pure PCR Template Preparation kit (Roche) was used as DNA extraction-purification protocol, following the manufacturer instructions. A real-time PCR system was used to confirm suspected infectious agents. Primarily, the insertion element IS901 sequence was used as target for *Mycobacterium avium* subsp. *avium* (Maa) (Pradenas et al 2014). Secondly, a real-time TaqMan PCR was used to target a specific for the 65-kD heat shock protein gene from *M. genavense* and MAC organisms (Tell et al 2001). Thirdly, a real-time PCR to identify *M. bovis* species was performed using a published protocol (Sweney et al 2007), specific for the RD4 genomic region.
repeats (TRs) from within the genomic DNA of *M. bovis*. The target loci included two exact tandem repeats (ETR A and C), six mycobacterial interspersed repetitive-units (MIRU 2, 10, 20, 23, 26, 39) and one Queen’s University of Belfast (Qub23) (Supply 2006). PCR amplicons were analysed by fragment analyser capillary electrophoresis system (Labgene Scientific SA). The electrophoretogram showed a VNTR code 7-5-2-5-2-4-7-1-6, in agreement with *M. bovis* ETR-A, ETR-C, MIRU 2, 10, 20, 23, 26, 39 and QUB23 loci, respectively (table 1). The presence of MIRU 10 with two repetitions instead of five and of MIRU 26 with seven repetitions instead of two to six identified a new *M. bovis* VNTR pattern. The electrophoretogram picks for the two mentioned VNTRs were clearly defined.

The Chilean hawk is a raptor living exclusively in the southern temperate forests of Chile and Argentina (33°-56° S) (Figueroa et al 2004). The bird is well adapted to this environment, feeding mainly on birds, a few rodents and insects (Trejo et al 2006). This hawk species is included in Appendix II of the Convention on International Trade in Endangered Species of Wild Fauna and Flora, CITES. The present study reports lesions consistent with tuberculosis and *M. bovis* detection and confirmation in a dead Chilean hawk. The latter represents an uncommon finding, since avian mycobacterial infections are mainly associated with *M. avium*, although a few other mycobacterial species have been identified (Steele 1995). The PCR negative results regarding no *Maa* and *M. genavense* detection, provide more supportive evidence that *Mycobacterium bovis* was the bacteria causing the lesions, and assure that there was no cross reaction with avian mycobacterial pathogens that can cause similar granulomatous lesions.

Mycobacterial infections in wild birds are difficult to detect and usually have a low prevalence, causing clinical disease only in a few cases (Tell et al 2001, Sánchez et al 2016). Moreover, clinical manifestations in wildlife are rarely found, and are usually identified only if a necropsy is performed (Hoop et al 1996), as was the case in the present study. Typical tuberculosis granulomas were identified from the liver, intestines and mesentery of the hawk and several of these granulomas contained acid-fast bacilli, suggesting an infection transmission by ingestion. The latter becomes an acceptable hypothesis, as Chilean hawk has experienced a decrease in the amount of prey species dwelling within the temperate native forest, which has been fragmented due to farming activities (Jaksic and Jiménez 1986). First, the hunting and consumption of infected rodents or birds and maybe scavenging behaviour of infected ruminants in farming grassland areas could therefore explain the present finding. In the area where the hawk was found dead, a bovine tuberculosis eradication program is currently underway and under control. However, high rates of infection in some areas and in some medium sized to large herds still remain (Max et al 2011).
Table 1. Number of copies of *M. bovis* VNTR for the nine loci analysed.

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<tr>
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<th>ETR</th>
<th>MIRU</th>
<th>QUB</th>
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<tr>
<td>Amplicons base pair</td>
<td>719</td>
<td>504</td>
<td>161</td>
</tr>
<tr>
<td>Nº repetitions</td>
<td>7</td>
<td>2</td>
<td>23</td>
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ETR: exact tandem repeats.
MIRU: Mycobacterial Interspersed Repetitive Units.
QUB: Queen’s University of Belfast.

The real-time PCR systems to confirm suspected infectious agents used specific probes and primers. Besides, the protocol considered a negative and positive controls in order to surveillance the proficiency of the reaction as well as DNA extraction negative and positive controls. Therefore, the odds of false-positive results were not expected. The typing information correlates with a *M. bovis* VNTR genotype with a SB0140 spoligotype, the most commonly found genotype in cattle in southern Chile (Zumárraga et al. 2013). This report expands the diversity of wildlife species that can become actively infected by *M. bovis*. Whether *M. bovis* infection in this Chilean hawk represents a risk of infection for livestock, humans or just a dead end host, remains to be determined.

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REFERENCES


