An Unidentified Filarial Species and its Impact on Fitness in Wild Populations of the Black-Footed Ferret (*Mustela nigripes*)

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AN UNIDENTIFIED FILARIAL SPECIES AND ITS IMPACT ON FITNESS IN WILD POPULATIONS OF THE BLACK-FOOTED FERRET (MUSTELA NIGRIPES)

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ABSTRACT: Disease can threaten the restoration of endangered species directly by substantially decreasing host survival or indirectly via incremental decreases in survival and reproduction. During a biomedical survey of reintroduced populations of the highly endangered black-footed ferret from 2002 to 2005, microfilariae discovered in the blood were putatively identified as Dirofilaria immitis, and widespread screening was initiated using a commercially available antigen-based ELISA test. A subset of animals (n = 16) was screened for D. immitis using a highly sensitive PCR-based assay. Microfilariae were also molecularly and morphologically characterized. Of 198 animals at six reintroduction sites, 12% had positive results using the ELISA test. No antigen-positive animals which were screened via PCR (n = 11) had positive PCR results, and all antigen-positive animals (n = 24) were asymptomatic. No significant differences were found in body mass of antigen-positive (male: 1223 ± 82 g [mean ± SD], female: 726 ± 75 g) vs. antigen-negative (male: 1,198 ± 119 g, female: 710 ± 53 g) individuals (P = 0.4). Antigen prevalence was lower in juveniles (3%) than adults (12%; P = 0.03), and higher in in situ, captive-reared individuals (33%) than wild-born individuals (10%; P = 0.005). Morphologic analysis of microfilariae revealed they were neither D. immitis nor any other previously characterized North American species. PCR amplification of the 5S spacer region of rDNA revealed that the filarial sequence shared only 76% identity with D. immitis. This previously unidentified filarial sequence was present in all antigen positive animals (11 of 11 tested). It appears that black-footed ferrets were infected with a previously undescribed species of filaria whose antigen cross-reacted with the ELISA assay, although further analysis is needed to make a conclusive statement. Nonetheless, this previously undescribed filaria does not appear to threaten recovery for this highly endangered mammal.

Key words: Black-footed ferret, cross-reactivity, Dirofilaria immitis, heartworm antigen, microfilaria, molecular characterization, Mustela nigripes, seroprevalence.

INTRODUCTION

Conservation and wildlife biologists recognize disease as a potential threat to the restoration of endangered species (Deem et al., 2001). Infectious diseases and the resulting pathology can have a direct negative effect on host survival. Epizootics have been implicated in the decline or extirpation of such species as the lion (Panthera leo) from Serengeti National Park (Roelke-Parker et al., 1996), albatross (Diomedea chlororhynchos, D. amsterdamensis, and Phoebetria fusca) from the Southern Ocean (Weimerskirch, 2004), and Hawaiian hon-
extirpated by epizootics of the canine distemper virus and sylvatic plague (Yersinia pestis; Thorne and Williams, 1988). Twenty-one years of captive breeding (from 1985 to 2006) and 15 yr of reintroduction efforts (from 1991 to 2006) have resulted in a population increase from a nadir of 18 captive individuals to more than 500 individuals in the wild and 240 individuals in the captive breeding program (Conservation Breeding Specialist Group, 2003). Although the black-footed ferret is no longer in immediate risk of extinction, disease remains a persistent threat to the success of reintroduction and species recovery efforts. Besides the omnipresent threats of canine distemper virus and sylvatic plague, other diseases have the potential to threaten or impede the recovery of the black-footed ferret.

To further our understanding of the health and disease threats to the reintroduced populations, a biomedical survey of wild black-footed ferrets was conducted to assess health, disease, immunology, reproduction, and genetics (Howard et al., 2006). From 2002 through 2006, a total of 253 black-footed ferrets were captured at reintroduction sites in South Dakota, Montana, Wyoming, Arizona, Utah, Colorado, and Mexico. Each animal was trapped and anesthetized for a physical examination and blood collection, then returned to the capture site. Blood smears prepared for hematology revealed the presence of microfilariae in two black-footed ferrets from northeast Utah. Superficial similarity of the microfilariae to those of Dirofilaria immitis and positive results from a commercially available, enzyme-linked immuno-sorbent antigen assay (ELISA) led us to initially identify the species as Dirofilaria immitis. Dirofilaria immitis is a ubiquitous filarial nematode with an expanding geographic range throughout the Americas (Sacks and Caswell-Chen, 2003; Labarthe and Guerrero, 2005). This filaria infects multiple companion species including dogs, cats, and domestic ferrets and has been implicated as an infective agent of many free-ranging wildlife worldwide (e.g., Weissman, 1992; Marks and Bloomfield, 1998; Nakagaki et al., 2000; Sacks and Blejwas, 2000; Pence et al., 2003; Bakker et al., 2006). Domestic ferrets have been shown to be a highly suitable and susceptible host of Dirofilaria immitis (McCall, 1998). Symptoms, based on clinical observations of naturally infected domestic ferrets, include coughing, severe dyspnea, moderate to severe exercise intolerance, moderate cyanosis, severe pleural effusion, heart murmur, decreased appetite, and weight loss consistent with adult worms residing in the heart and associated vessels (Antinoff, 2001). Given the substantial pathology of infected domestic ferrets, infection in the endangered black-footed ferret could have a significant impact on the recovery of this species.

Prior to the initiation of this biomedical survey, Dirofilaria immitis had not been observed in the black-footed ferret and was not considered a threat to the wild population. To assess the prevalence of the filarial infection, widespread testing at six reintroduction sites using a commercially available antigen-based ELISA test for Dirofilaria immitis was initiated. Rates of infectivity among juveniles vs. adults and captive-reared vs. wild-born individuals were compared. As a crude assessment of fitness, body mass among antigen positive and antigen negative animals were compared. Results of the ELISA test were validated with molecular tests and morphologic characterization of the filariae in black-footed ferrets.

**MATERIALS AND METHODS**

**Animals, anesthesia, and blood analysis**

Black-footed ferrets were evaluated during the fall or spring monitoring surveys at six reintroduction sites in Buffalo Gap National Grasslands and Badlands National Park, South Dakota, USA (43°15’N, 109°06’W); Janos, Chihuahua State, Mexico, (31°04’N, 107°51’W); Aubrey Valley, Arizona, USA (35°32’N, 113°10’W), Coyote Basin, Utah and Wolf Creek, Colorado, USA...
(40°15′N, 109°06′W), Shirley Basin, Wyoming, USA (42°07′N, 106°03′W), and Charles M. Russell National Wildlife Refuge, Montana, USA (47°37′N, 107°46′W). All animals were captured in the wild but were born either in the wild or in captivity and then reintroduced. Trapping and anesthesia protocols for the black-footed ferret are well established (Scheets, 1972; Kreeger et al., 1998). Animals were cage-trapped at night and returned to the same trap location following examination and recovery from anesthesia. Additionally, 21 ex situ black-footed ferrets from the U.S. Fish and Wildlife Service–National Black-Footed Ferret Conservation Center in Wheatland, Wyoming (41°46′N, 105°19′W) were sampled. All animals received detailed physical examinations by veterinarians. All trapping and handling was authorized, coordinated, and conducted by the U.S. Fish and Wildlife Service’s Black-Footed Ferret Recovery Implementation Team as part of routine population monitoring for reintroduction sites.

Each black-footed ferret was transferred to an anesthesia induction chamber and anesthetized via inhalation using isoflurane. A physical examination was conducted and weight obtained. Whole blood was collected from the jugular vein and aliquoted into EDTA tubes (for hematology) and serum separator clot tubes (for serum chemistry). Following clotting, serum tubes were centrifuged, and serum was recovered and frozen on site. The blood sample in EDTA and the blood smears prepared from each EDTA sample were shipped within 24 hr of collection to the Wyoming State Veterinary Laboratory (Laramie, Wyoming) for assessment of a complete blood count (CBC) with differential evaluation of white blood cells. Blood smears also were evaluated for microscopic detection of circulating microfilariae. This test is less sensitive than other approaches (e.g., modified Knott test) for evaluating prevalence of microfilariae, and likely underestimated the number of individuals with circulating microfilariae. Serum samples were maintained frozen and later shipped to Wyoming for diagnostic testing. Both plasma and serum samples were evaluated for heartworm using a commercially available ELISA kit designed to detect adult *D. immitis* antigen (DIROCHEK®; Synbiotics, Inc., San Diego, California). Plasma was used initially for the ELISA test; tests were reevaluated using serum.

To determine if antigen-positive animals had less body mass than antigen-negative animals, a univariate analysis of variance (ANOVA) on body with sex as an additional treatment was conducted. For this analysis, only wild-born adult animals were used; the analysis was conducted with and without location as a treatment. To test if adults and juveniles had significantly different frequencies of antigen-positive ELISA tests, a Yates corrected chi-square ($\chi^2$) test of independence was used for in situ animals. The same statistical test was used to determine whether the prevalence of antigen-positive animals was different for in situ animals that were captive-reared or wild-born.

**Molecular and morphologic identification of filarial DNA**

An aliquot of whole blood (250 µl) from ferrets was stored in 1 ml standard lysis buffer and stored at 4° C until DNA was extracted with a Qiagen Tissue Kit (Qiagen Inc., Hilden, Germany) following the manufacturer’s protocol. We conducted two polymerase chain reaction (PCR)–based screens on a subset of the DNA samples extracted from ferret blood; nine individuals in which microfilariae were seen and ELISA tests were positive, two individuals that had positive ELISA tests but no microfilariae were seen on blood smears, and five individuals that were negative for both ELISA tests and microfilariae. We used universal primers to amplify the coding region of the 5S rDNA of all filarial parasites (Xie et al., 1994). In addition, we used a highly sensitive, nested PCR protocol with *D. immitis* specific primers to more accurately diagnose infection in these animals (US Patent No. 6268153). PCR was conducted with positive and negative controls and products were run on a 2.5% agarose gel to visualize the presence of amplified DNA. All PCR reagents except primers were obtained from Applied Biosystems (Foster City, California). The PCR reaction contents included: 5 µl 10× reaction buffer, 8 µl 1.25 mM dNTP mixture, 20 pmoles of each primer, and 1 unit of *TagGold* DNA polymerase. The PCR cycling conditions were 95 C 12 min, followed by 30 cycles of 95 C 1 min, 55 C 1 min, 72 C 1 min, and a 10 min final extension at 72 C.

The amplified 5S spacer DNA was sequenced using the same primers as used for PCR. Sequencing reactions used the Big Dye v. 3.1 Cycle Sequencing Mix (Applied Biosystems). The 5S spacer sequences were aligned using Megalign software (Lasergene, DNASTAR, Inc., Madison, Wisconsin). Phylogenetic trees using the unknown ferret filarial 5S sequence and all of the filarial 5S ribosomal spacer sequences previously published (Xie et al., 1994) were calculated using the ClustalW “slow/accurate” method (Thompson et al., 1994) in Lasergene.
The morphology of microfilariae collected from two black-footed ferrets at the Utah/Colorado location was examined by one co-author (OB) using light microscopy and accessioned into the Muséum National d’Histoire Naturelle (Accession No. 156 JW). Specimens were keyed to family based on microfilaria morphology. Genus and species cannot be identified solely by microfilariae; however, certain taxa could be eliminated on the basis of their morphologic characteristics. Particular attention was paid to species known to occur in North America.

RESULTS

Antigen prevalence and physical examination

Two hundred twenty plasma or serum samples collected from 198 free-ranging individuals at the six reintroduction sites from 2002 to 2005 were tested. Twenty-four individuals (12%\textsuperscript{a}) distributed among five of the six ex situ reintroduction sites had positive ELISAs; no positive animals were found at the breeding center. Nine individuals, all from the Utah/Colorado reintroduction site, had microfilariae visible in the blood; this site also had the highest frequency (78\%) of positive antigen tests (Table 1). Twenty-two individuals were recaptured; two individuals seroconverted between captures, and three antigen-positive individuals were recaptured on multiple occasions. Two of these individuals dispersed 17 km between captures (Table 2).

All animals appeared free of symptoms associated with *D. immitis* infection. No evidence of dyspnea, cyanosis, or unusual heart rhythms was detected. Animals with chronic infection, as detailed by our recaptures, appeared healthy at each capture (up to 1 yr between captures). No substantial weight fluctuation in these animals was found (Table 2). For the comparison of body mass between antigen-positive and antigen-negative animals, results did not differ between models that included or excluded location as a factor; therefore, only results of the ANOVA model conducted without location effect were reported. No significant difference in body mass of wild-born adults that were antigen-positive vs. negative was found; all of the variance in the model was explained by the difference in body mass between sexes as expected in this sexually dimorphic species (mean\(\pm\)S.D. for antigen-positive females: 726\(\pm\)75 g; antigen-negative females: 710\(\pm\)53 g; antigen-positive males: 1,223\(\pm\)82 g; antigen-negative males: 1,197\(\pm\)119 g; overall model: \(F_{3,103}=237, P<0.001\), antigen: \(F_{1,103}=0.7, P=0.4\), sex: \(F_{1,103}=381, P<0.001\)).

The prevalence of antigen-positive juveniles (2 of 77) was significantly less than that of adults (15 of 122; \(\chi^2=4.5, df=1, P=0.03\)); however, in situ, the prevalence of antigen-positive captive-reared individuals (7 of 21) was significantly higher than wild-born individuals (17 of 178; \(\chi^2=7.9, df=1, P=0.005\)). No antigen-positive animals were found in the ex situ captive breeding population.

<table>
<thead>
<tr>
<th>Location</th>
<th>Sample size</th>
<th>No. antigen-positive</th>
<th>No. microfilaria positive</th>
<th>No. sequence positive\textsuperscript{a}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arizona</td>
<td>45</td>
<td>6</td>
<td>0</td>
<td>4 (5)</td>
</tr>
<tr>
<td>Utah/Colorado</td>
<td>9</td>
<td>7</td>
<td>9</td>
<td>8 (9)</td>
</tr>
<tr>
<td>Mexico</td>
<td>9</td>
<td>3</td>
<td>0</td>
<td>0 (0)</td>
</tr>
<tr>
<td>South Dakota</td>
<td>101</td>
<td>5</td>
<td>0</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Montana</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0 (2)</td>
</tr>
<tr>
<td>Wyoming</td>
<td>32</td>
<td>3</td>
<td>0</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Captive facility</td>
<td>21</td>
<td>0</td>
<td>0</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

\textsuperscript{a} These 16 samples were also PCR-assayed for *Dirofilaria immitis*. No positive samples were found.
Molecular identification of a filarial species

All 5S spacer sequences obtained from samples of 16 ferrets were identical and amplified in 12 of 16 samples (Table 1). The sequence was 360 base pairs long (GenBank Accession No. EU055542) and did not match published *D. immitis* 5S spacer sequences. In fact, the ferret parasite sequence and *D. immitis* sequence were very dissimilar. When compared with all published filarial 5S spacer sequences, the unknown ferret parasite sequence was most similar to *Acanthocheilonema viteae* (97% identity, Fig. 1). All of the other filarial species (*Litomosoides sigmodontis*, *Loa loa*, *Mansonella perstans*, *Brugia pahangi*, *Brugia buckleyi*, *Wuchereria bancrofti*, *Onchocerca volvulus*, *Onchocerca gutturosa*, and *D. immitis*) were much more divergent than *A. viteae*. In this data set, the *D. immitis* 5S spacer sequence was the least similar to the unknown ferret parasite sequence (76% identity). The ClustalW alignment and the resulting phylogenetic tree confirmed the distant relationship between *D. immitis* and the filaria found in black-footed ferrets (Fig. 1). None of the 16 animals, whether antigen-positive or negative, were positive for the PCR-based *D. immitis* test. All positive controls amplified at the expected length of 155 base pairs, which indicated that negative results from ferret samples were due to a lack of *D. immitis* DNA in the blood, not because of PCR failure.

**Morphology**

Microfilariae from both black-footed ferrets had similar morphologic characters, thus they were considered to be the same species. Microfilariae were observed in these individuals at each capture.

**Table 2. Chronology of infection, sex, and body mass for five black-footed ferrets that were recaptured during the biomedical survey.**

<table>
<thead>
<tr>
<th>Individual</th>
<th>Date of initial capture</th>
<th>Body mass (g)</th>
<th>Serology (+/−)</th>
<th>Date of second capture</th>
<th>Body mass (g)</th>
<th>Serology (+/−)</th>
<th>Date of third capture</th>
<th>Body mass (g)</th>
<th>Serology (+/−)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arizona 8, male</td>
<td>October 2004</td>
<td>1,212</td>
<td>(−)</td>
<td>March 2005</td>
<td>1,196</td>
<td>(+)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arizona 3, female</td>
<td>October 2004</td>
<td>688a</td>
<td>(+)</td>
<td>March 2005</td>
<td>614</td>
<td>(−)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>South Dakota 109, male</td>
<td>September 2002</td>
<td>678a</td>
<td>(−)</td>
<td>March 2003</td>
<td>1,124</td>
<td>(+)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Utah 3, maleb</td>
<td>August 2004</td>
<td>1,024</td>
<td>(+)</td>
<td>March 2005</td>
<td>1,080</td>
<td>(+)</td>
<td>August 2005</td>
<td>1,188</td>
<td>(+)</td>
</tr>
<tr>
<td>Utah 4, femaleb</td>
<td>August 2004</td>
<td>732</td>
<td>(+)</td>
<td>April 2005</td>
<td>850</td>
<td>(+)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Utah 5, maleb</td>
<td>March 2005</td>
<td>1,228</td>
<td>(+)</td>
<td>August 2005</td>
<td>1,180</td>
<td>(+)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

These individuals were juveniles (less than 1 yr old) at date of first capture.

Microfilariae were observed in these individuals at each capture.

*a* These individuals were juveniles (less than 1 yr old) at date of first capture.
Figure 1. Unrooted phylogenetic tree of the 5S spacer sequence of nine known filarial species and the unknown filarial sequence isolated from the blood of a black-footed ferret. The tree was constructed using the slow/accurate algorithm of computer program, ClustalW. Branch tips are labeled with the filarial species, the Genbank accession number of the sequence used to build the tree, and percent similarity (base pair congruence) between the ferret filarial 5S spacer sequence and the 5S spacer sequence from other filarial parasites.

Figure 2. A. *Dirofilaria immitis* from a blood smear of a domestic dog. B. Unidentified microfilariae from a blood smear of a black-footed ferret. Bar in lower right corner of each photograph represents 50 μm.
A comparison of morphologic features among filarial species reported in North America (Table 3), allowed exclusion of these species. The first 15 species listed in Table 3 have longer microfilariae than the unidentified filariae. Furthermore, in the subfamily Dirofilariinae, *D. immitis* and *D. tenuis* were conclusively excluded because they lack an internal body. *Mansonella* (*M.*) *llewellyni*, *M. (M.)* *interstitium* were excluded because they have no internal body. The two *Molinema* species had a longer caudal filament. *Acanthocheilonema reconditum*, as well as *A. mephitis* and *A. procyonis* were excluded because they lack an internal body. Microfilariae of *Monanema* are not found in blood but in skin (in marmots) and are sheathed.

**DISCUSSION**

The identity of the infective agent of antigen-positive animals remains elusive, but we hypothesize that the antigen of an unidentified filaria, whose microfilariae were observed in some animals, cross-reacted with the *D. immitis* ELISA test, and thus the widespread positive results were attributable to a wide-ranging unidentified filariae. We base our hypothesis on four lines of evidence: 1) PCR-based *D. immitis* tests were negative for all antigen-positive animals that we tested (n = 11) suggesting that the ELISA test produced false positives; 2) unidentified filariae sequence was found in 10 of 11 antigen-positive animals, including eight of the nine animals in which we visually identified circulating microfilariae, which suggests that the unidentified filariae cross-reacted with the ELISA test; 3) the ferret population with the highest prevalence of circulating microfilariae (Utah/Colorado) also had the highest prevalence of antigen-positive animals and the highest prevalence of unidentified filariae based on molecular sequence data; 4) all antigen-positive animals were asymptomatic, including six animals which were captured 6 mo to 1 yr later. Given the highly debilitating nature of *D. immitis* in domestic ferrets, symptoms of dyspnea and cyanosis during the stress of capture and anesthesia were expected to occur.

![](https://i.imgur.com/3.png)

<table>
<thead>
<tr>
<th>Filarial species</th>
<th>Length</th>
<th>Width</th>
<th>Sheath</th>
<th>I. b.</th>
<th>C. f.</th>
<th>References</th>
<th>Host</th>
</tr>
</thead>
<tbody>
<tr>
<td>?</td>
<td>165–185</td>
<td>5</td>
<td>—</td>
<td>9–13</td>
<td>9–13</td>
<td>Present study</td>
<td>Mustela nigripes</td>
</tr>
<tr>
<td><em>Dirofilaria immitis</em> (Leidy 1856)</td>
<td>233–270</td>
<td>5.4–6.5</td>
<td>—</td>
<td>—</td>
<td>25</td>
<td>Fuellebom, 1912</td>
<td>Canis familiaris</td>
</tr>
<tr>
<td></td>
<td>307–322</td>
<td>6.7–7.1</td>
<td>—</td>
<td>—</td>
<td></td>
<td>Newton and Wright, 1956</td>
<td>Canis familiaris</td>
</tr>
<tr>
<td><em>Dirofilaria striata</em> (Molin 1858)</td>
<td>230–240</td>
<td>2–3</td>
<td>—</td>
<td>ND</td>
<td>++</td>
<td>Orihel and Ash, 1964</td>
<td>Lynx rufus</td>
</tr>
<tr>
<td></td>
<td>327–371</td>
<td>4–5</td>
<td>—</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Dirofilaria tenuis</em> (Chandler 1942)</td>
<td>220–250</td>
<td>2–3</td>
<td>—</td>
<td>—</td>
<td>22–25</td>
<td>Orihel and Beaver, 1965</td>
<td>Procyon lotor</td>
</tr>
<tr>
<td><em>Dirofilaria subdermata</em> (Mönnig 1924)</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>No reference</td>
<td>Erethizon dorsatum</td>
</tr>
<tr>
<td><em>Loa loa</em> (Price 1957)</td>
<td>285±</td>
<td>6</td>
<td>+</td>
<td>35</td>
<td>0</td>
<td>Eberhard and Orihel, 1984</td>
<td>Sylvilagus floridanus</td>
</tr>
<tr>
<td><em>Pelecitus scapiceps</em> (Leidy 1886)</td>
<td>220–250</td>
<td>5</td>
<td>+</td>
<td>35</td>
<td>0</td>
<td>Bartlett, 1983</td>
<td>Lepus americanus</td>
</tr>
<tr>
<td><em>Dirofilaria formia pulmoni</em> (Davidson 1975)</td>
<td>213–288</td>
<td>3–4</td>
<td>—</td>
<td>—</td>
<td>15</td>
<td>Davidson, 1975</td>
<td>Sciurus carolinensis</td>
</tr>
<tr>
<td><em>Brugia beaveri</em> (Ash and Little 1964)</td>
<td>300±</td>
<td>6</td>
<td>+</td>
<td>45</td>
<td>0</td>
<td>Ash and Little, 1964</td>
<td>Procyon lotor</td>
</tr>
<tr>
<td><em>Brugia lepori</em> (Eberhard 1984)</td>
<td>210–266</td>
<td>4–6</td>
<td>+</td>
<td>35</td>
<td>0</td>
<td>Eberhard, 1984</td>
<td>Sylvilagus aquaticus</td>
</tr>
<tr>
<td></td>
<td>275–330</td>
<td>5–7</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Mansonella (Mansonella) llewellyni</em> (Price 1962)</td>
<td>290±5±</td>
<td>2.5</td>
<td>—</td>
<td>0</td>
<td>15</td>
<td>Price, 1962</td>
<td>Procyon lotor</td>
</tr>
<tr>
<td><em>Molinema arbata</em> (Highby 1943)</td>
<td>280–297</td>
<td>6</td>
<td>—</td>
<td>—</td>
<td>40</td>
<td>Highby, 1943</td>
<td>Erethizon dorsatum</td>
</tr>
<tr>
<td><em>Molinema spreti</em> (Anderson 1953)</td>
<td>400–450</td>
<td>8</td>
<td>—</td>
<td>80</td>
<td>35</td>
<td>Anderson, 1953</td>
<td>Castor canadensis</td>
</tr>
<tr>
<td><em>Dasypodifilaria averyi</em> (Eberhard 1984)</td>
<td>254–394</td>
<td>7–9</td>
<td>+</td>
<td>ND</td>
<td>5</td>
<td>Eberhard, 1984</td>
<td>Dasypus novemcinctus</td>
</tr>
<tr>
<td><em>Acanthocheilonema reconditum</em> (Grassi 1890)</td>
<td>269–283±</td>
<td>4.3–4.8</td>
<td>—</td>
<td></td>
<td>17</td>
<td>Newton and Wright, 1956</td>
<td>Canis familiaris</td>
</tr>
<tr>
<td><em>Acanthocheilonema mephitis</em> (Webster and Beauregarde 1964)</td>
<td>186–218</td>
<td>3–3.8</td>
<td>—</td>
<td></td>
<td>6.6–9.5</td>
<td>Webster and Beauregarde, 1964</td>
<td>Mephitis mephitis</td>
</tr>
<tr>
<td><em>Acanthocheilonema procyonis</em> (Price 1955)</td>
<td>129–149</td>
<td>3.6–4.9</td>
<td>—</td>
<td>—</td>
<td></td>
<td>Smith, 1980</td>
<td>Procyon lotor</td>
</tr>
</tbody>
</table>

*Measurements from fixed blood or female uteri, otherwise in blood smears, all in μm.*
but were not observed. Recaptured, antigen-positive animals exhibited no weight loss, and in two cases, animals had dispersed substantial distances (~17 km) between captures. Additionally, no difference in body mass among antigen-positive and antigen-negative adults was observed. These observations suggest that the antigen-producing filariae did not produce obviously debilitating symptoms, and is likely not a major threat to black-footed ferret recovery. Although our results are highly suggestive, further testing with larger sample sizes is warranted.

**Filaria identification**

Both molecular and morphologic analysis indicated that the microfilariae seen in blood smears were not *D. immitis*. The molecular sequence of this species has not been previously described. The filaria shares 97% sequence similarity with *Acanthocheilonema viteae*, but this level of divergence indicates that the filaria is a separate species. Furthermore, the morphology was not consistent with *A. viteae*. Morphologic identification of the adult filariae remains undetermined, because adult worms have yet to be located. Clearly, infected individuals of an endangered species cannot be sacrificed to locate and collect adult worms. Further complicating matters, black-footed ferrets are an elusive, fossorial animal; dead individuals are rarely found above ground. Monitoring of captive individuals will continue, and necropsies will include a search for adult worms.

Several suppositions can be proposed about the life history of the antigen-producing filariae, whatever its identity may be, and the effect it has on its host. All juvenile black-footed ferrets were captured between the ages of 2 and 4 mo of age, based on an estimated June 1st birth date. The low antigen prevalence of the filariae in 2–4-mo-old juveniles suggests that either the life cycle of the filariae had not been completed or that juveniles were not infected at that age. A low prevalence of *D. immitis* infection in host juveniles was found in the coyote (*Canis latrans*, Sacks and Blejwas, 2000), which was due to the 6–7 mo prepatent period of *D. immitis*. The life cycle of the filariae found in our study remains unknown, yet the high prevalence of microfilariae in Utah/Colorado animals (100%) suggests that the black-footed ferret is a suitable host for the completion of the filariae’s life cycle.

Free ranging individuals that were captive-reared had a significantly higher prevalence of infection (33%) than wild-born individuals (10%). Two explanations are possible for this increased prevalence. The captive environment could have increased an animal’s risk of exposure to the vector. A sample of captive individuals revealed no antigen-positive animals; however, our sample size (*n* = 21) was small and confined to one of the six captive breeding facilities. Alternatively, captive-reared individuals that were reintroduced to the wild could have been more susceptible to infection in the wild. Recently released animals likely spend more time above ground in an attempt to find resources and establish territories, which could increase their risk of exposure to vectors of filariae. In addition, the stress of the release could suppress the immune system making these animals more susceptible to disease and infections (Wingfield and Romero, 2001).

**Management implications**

It appears that the ubiquitous prevalence of this filarial species across the range of the black-footed ferret does not pose an immediate threat to the recovery of this highly endangered species. The debilitating effects of heartworm infection on the survival of animals were not observed; however, the study was not designed to look at the other important contributor to fitness, reproduction. Parasitic infections can cause subtle, but important decreases in reproduction, which in turn can increase the risk of extinction (Anderson and May, 1978). For
the black-footed ferret, decreases in any vital rate that limits the growth rate could have devastating consequences to species recovery. The increased prevalence of infection in captive-reared free-ranging animals merits further investigation.

Antigen-based ELISA kits may not be a reliable test for diagnosing heartworm in free-ranging animals. Indeed, it was suspected that these kits also provided false-positives when used in the Channel Island fox (Roemer et al., 2000). Future assessments of prevalence of *D. immitis* in black-footed ferrets should utilize the PCR-based assay.

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