Sarcocystis sp. parasites in the Mexican Great-tailed Grackle (Quiscalus mexicanus), Bronzed Cowbird (Molothrus aeneus), and Stripe-headed Sparrow (Aimophila ruficauda)

Abstract

The objective of this study was to describe the morphological and ultrastructural characteristics, the polymerase chain reaction and restriction fragment length polymorphism (PCR-RFLP) results, the sequences and the phylogenetic analysis of a specific fragment of internal transcribed spacer 1 (ITS-1), amplified using the 25/396 primers, of the Sarcocystis sp. parasites identified in the muscles of wild great-tailed grackles, bronzed cowbirds, and stripe-headed sparrows in Mexico. Fifteen birds with sarcocystosis in their skeletal muscles were studied: 7 great-tailed grackles (Quiscalus mexicanus), 6 bronzed cowbirds (Molothrus aeneus), and 2 stripe-headed sparrows (Aimophila ruficauda). Histopathological analysis revealed thin-walled mature parasite cysts. Ultrastructurally, the cyst wall consisted of a granular layer with villar protrusions and numerous microtubules. The bradyzoites measured 4.1 × 1.6 µm, and micronemes appeared in the anterior third of the conoid. For molecular identification, PCR-RFLP was performed using sequences of a specific fragment of internal transcribed spacer 1 (ITS-1) using the primers 25/396 and Hinf I. Hind III did not cut this fragment. The sequencing results indicated a 100% similarity among the Sarcocystis parasites from the three bird species, and a BLAST search revealed 96% sequence similarity with S. neurona. The phylogenetic analysis shows that the sequences studied are topologically distant to those sequences reported for S. neurona in the United States and in South America and are not related to any group previously reported. Although our morphological and molecular analysis data provide strong evidence that S. neurona uses these bird species as intermediate hosts, future molecular studies with additional DNA fragments, combined with biological studies, will ultimately allow us to convincingly identify these parasites. This is the first report of a Sarcocystis sp. parasite in wild birds in Mexico that may be S. neurona.

Keywords: Sarcocystis; Sarcocystis neurona; Quiscalus mexicanus; Molothrus aeneus; Aimophila ruficauda; Histopathology; Ultrastructure; PCR-RFLP, Phylogenetic analysis.
Introduction

Parasites from the genus Sarcocystis are protozoans of the phylum Apicomplexa. These parasites affect mammals, birds, reptiles, amphibians, and fish (Munday et al., 1979; Bolon et al., 1989; Hillyer et al., 1991; Dubey et al., 2001a; Dubey et al., 2003). Sarcocystis falcata is the most prevalent species in birds, which serve as intermediate hosts, with opossums (Didelphis virginiana and Didelphis albiventris) serving as the definitive hosts (Box and Duszynski, 1978; Box and Smith, 1982; Dubey et al., 2000; Dubey et al., 2001b). Until 1995, the opossum was thought to be the definitive host of only S. falcata; however, the opossum is known also to be the definitive host of S. neurona, S. speeri (Fenger et al., 1997; Dubey et al., 1998; Dubey et al., 1999) and S. lindsayi (Dubey et al., 2001c). Mansfield et al. (2008) reported S. neurona in brown-headed cowbirds. Dame et al. (1995) found a strong similarity between S. neurona and S. falcata based on an analysis of the 18S rRNA gene and, thus, suggested that these are actually the same species. Subsequent biological studies, however, confirmed that these are indeed two different species (Marsh et al., 1997a; Dubey and Lindsay, 1998). Genetic typing by various methods has established that S. neurona and S. falcata are distinct species. Polymerase chain reaction (PCR) primers targeting the SSU rRNA gene were developed that distinguished S. neurona from the Sarcocystis found in skunks, raccoons, hawks, coyotes and cats (Fenger et al., 1995). In another study, sequencing of the internal transcribed spacer 1 (ITS-1) region of the rRNA gene showed that S. falcata may be composed of a heterogeneous population and that the ITS-1 region can be used to distinguish S. neurona from S. falcata (Marsh et al., 1999). Tanhauser et al. (1999) used ITS-1 to design specific primers (25/396) and restriction enzymes (Hinf I and Hind III) to perform PCR-restriction fragment length polymorphism (RFLP) analysis to differentiate S. neurona and S. falcata. Likewise, Elsheikha et al. (2005) have used DNA markers 25/396 of ITS-1 for phylogenetic studies of S. neurona in the United States. Infections caused by S. neurona have been reported in a variety of species, such as cats (Butcher et al., 2002), raccoons (Dubey et al., 2001d), armadillos (Cheadle et al., 2001a), skunks (Cheadle et al., 2001b), seals (Miller et al., 2001), and sea otters (Dubey et al., 2003). In addition, S. neurona is an important cause of neurological problems in horses in the United States (Dubey et al., 1991; MacKay et al., 2000), although horses appear to be aberrant hosts (Dubey 2001b). Interestingly, schizonts have been observed in the brain and spinal cord and mature cysts have been observed in the skeletal muscle in a 4-year-old mare, suggesting that horses are intermediate hosts (Mullaney et al., 2005). In 2008, the presence of parasitic cysts of S. neurona in the skeletal muscles of brown-headed cowbirds (Molothrus ater) was documented, indicating that these birds may be intermediate hosts (Mansfield et al., 2008). In Mexico there are no data available on the presence of S. neurona in birds, but Yeargan et al., in 2013, found a seroprevalence of 48.5% of S. neurona in horses in northern Mexico.

The objective of this study was to describe the morphological and ultrastructural characteristics, the PCR-RFLP results, the sequences and the phylogenetic analysis of a specific fragment of internal transcribed spacer 1 (ITS-1), which was amplified using the 25/396 primers, of the Sarcocystis sp. parasites identified in the muscles of wild great-tailed grackles, bronzed cowbirds, and stripe-headed sparrows in Mexico that may be S. neurona.
Materials and methods
Fifteen wild birds (7 great-tailed grackles [Quiscalus mexicanus], 6 bronzed cowbirds [Molothrus aeneus], and 2 stripe-headed sparrows [Aimophila ruficauda]) found dead with suspected poisoning in the State of Morelos, Mexico, were submitted to the Diagnostic Laboratory and Research on Diseases of Birds of the Department of Medicine of Birds of the Faculty of Veterinary Medicine of the National Autonomous University of México (UNAM). During necropsy, parasitic structures were found in of the muscle.

Necropsy and histopathology
All birds were submitted for necropsy, and samples were taken from muscle, fixed in 10% buffered formalin, processed by routine histological techniques, embedded in paraffin, cut into 4-µm sections, and stained with hematoxylin and eosin (H&E).

Electron microscopy
Sections (3 mm²) of striated skeletal muscle with parasitic cysts were taken from great-tailed grackles, fixed in 2.5% glutaraldehyde and, subsequently, post-fixed with 1% osmium tetroxide for 2 hours. Following washing with 0.1 M cacodylate buffer solution (pH 7.2), tissues were dehydrated with increasing concentrations of acetone. Sections were then embedded in epoxy resin (Epón 812, Electron Microscopy Sciences, Industry Road Hatfield, PA) and, finally, were polymerized at 60°C for 24 hours. Afterwards, 200-µm thick semifine cuts were made using an ultramicrotome, and samples were mounted on slides and contrasted with toluidine blue (Hayat, 2000). Fine cuts were made to achieve 60-µm samples, and the samples were then mounted on copper mesh grids, contrasted with uranyl acetate and lead citrate, and observed with an electron microscope (Zeiss EM-900, Zeiss, Oberkochen, Germany) at 80 kV.

DNA extraction
To isolate the parasite DNA, muscle samples with cysts of the parasites were macerated with a pestle and suspended in 20% phosphate buffered saline (PBS). Then, 250 µl of the suspension was mixed with 250 µl of lysis buffer solution (EDTA-SDS-Tris HCl; GibcoBRL, Grand Island, NY, USA) and 25 µl of proteinase K (20 mg/ml; Fermentas Inc., Glen Burnie, MD, USA). The samples were then incubated in a 37°C water bath for 2 hours followed by DNA purification with phenol-chloroform-isooamyl alcohol. The DNA was then precipitated with ethanol and hydrated.

PCR
The following primers were used for the detection of S. falcatula or S. neurona: 25 5'-CAC ACA AAA CAC CTG AAA GTC ACG TAC TT-3' and 396 5'-CCT GCC TCA CTT CGA CAC AT-3' (Sigma-Aldrich Corp., St. Louis, MO, USA). These primers amplify a 334-bp fragment of ITS-1 of the ribosomal DNA gene. The PCR conditions were as follows: 2 µl of Taq buffer, 0.4 µl of dNTPs (0.2 mM), 1 µl of primer (0.5 mM), 1.2 µl of MgCl₂ (1.5 mM), 1 µl of Triton (0.10%), 1 µl of bovine serum albumin
(BSA) (0.015 mg/ml), 0.5 µl of Taq polymerase (2.5 U/µl; Fermentas Inc., Glen Burnie, MD, USA), 7.9 µl of diethylpyrocarbonate (DEPC) water and 5 µl (200 ng/µl) of DNA. The samples were amplified with a PCR Sprint Thermal Cycler (PCR Sprint Thermal Cycler, Thermo Fisher Scientific, Inc., Waltham, MA, USA) the following conditions: 1 cycle at 94°C for 5 minutes; 30 cycles at 94°C for 30 seconds, 62°C for 30 seconds and 72°C for 40 seconds; and 1 cycle at 72°C for 5 minutes. Then, 5 µl of each PCR reaction was electrophoresed on a 2% agarose gel, which was then stained with ethidium bromide and observed under ultraviolet light.

**Sequencing**

The approximately 334-bp PCR fragment that was observed on the agarose gel was cut out and purified with a QIAquick® Gel Extraction Kit (QIAGEN, Hilden, Germany) following the manufacturer’s instructions. The purified fragment was visualized in a 1% agarose gel that was stained with ethidium bromide, along with the molecular weight marker GeneRuler (Fermentas Inc., Glen Burnie, MD, USA). PCR sequencing was performed with a BigDye® Terminator version 3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA, USA) following the manufacturer’s instructions. The sequencing reactions were purified in CENTRI-SEP™ Spin Columns (CS-901, Princeton Separations, Adelphia, NJ, USA) following the manufacturer’s instructions. Subsequently, the sequencing reactions were read with a 3130 Genetic Analyzer (Applied Biosystems Foster City, CA, USA). Sequencing of double-stranded DNA for the marker 25/396 was carried out to confirm the nucleotide sequence, and the primers 25/396 were used to obtain sense and antisense sequences.

Raw sequences (sense and antisense) were edited using MEGA version 4 software (Tamura et al., 2007). The consensus sequence yielded a file corresponding to each of the samples, and an alignment among the three sequences - the stripe-headed Sparrow (G1), the bronzed cowbird (T1), and the great-tailed grackle (ZS) - was carried out.

**Phylogenetic analysis**

Phylogenetic analysis was based on sequences of the 25/396 DNA marker obtained from G1, T1 and Z5 in addition to the sequences obtained from GenBank. Sequences were aligned, and a phylogenetic tree was constructed with the neighbor-joining (NJ) method using the Kimura 2-parameter (K-2P) model. Bootstrap support for the results of the NJ analysis was based on 1,000 replicate datasets generated from the original multiple sequence alignment.

**RFLP**

The approximately 334-bp PCR fragment that was observed on the agarose gel was digested with two restriction enzymes, Hinf I (Life Technologies, Carlsbad, CA, USA) and Hind III (Roche Applied Science, Indianapolis, IN, USA). The following RFLP conditions were used: 10 µl of PCR product, 3 µl of buffer, 1 µl of enzyme (Hinf I or Hind III), and 16 µl of DEPC water. The samples were incubated for 12 hours in a water bath at 37°C. Then, 15 µl of the reaction was electrophoresed on a 3% agarose gel that was stained with ethidium bromide and observed under ultraviolet light.
Results

The bodies of all birds were in good condition, and intramuscular parasitic cysts were found in the striated muscle of the breast, wings, and legs. These cysts were white in color, elongated, and undulating, and their dimensions ranged from 0.3 to 0.5 cm in length (Fig. 1). No pathological changes were evident in other organs.

Microscopic examination of the muscle tissue sections of the 15 birds showed multiple round (20 to 200 µm in diameter), elongated (100 to 4000 µm in length) cyst structures, each with a continuous thin wall (less than 2 µm) eccentrically compressing the myofibrils. Inside, the Sarcocystis parasites contained abundant bradyzoites that in some areas were separated by septa. In the central part of the larger cysts, an acellular, amorphous pale eosinophilic material that corresponded to degenerated protozoa was observed (Fig. 2). In 3 great-tailed grackles, discrete, multifocal, inflammatory cell aggregates composed of lymphocytes and plasma cells were observed around the cysts.

Electron microscopy

Striated skeletal muscle sections exhibited sarcoplasmic, thin-walled parasitic cysts consisting of a granular layer with villar protrusions containing numerous electron-dense microtubules that extended from the point to the base and beyond the underlying granular layer. The parasitic cysts showed some electrolucid merozoites beneath their walls, whereas toward the central portion, numerous mature bradyzoites were grouped together and separated by prolongations of the granular layer. Bradyzoites measured 4.1 µm in length and 1.6 µm in width and exhibited numerous micronemes in the anterior third of the conoid. The majority of nuclei are round with abundant electron-dense heterochromatin, which adhered to the internal nuclear sheath, and occasionally, a prominent nucleolus was noted (Fig. 3 y 4).

PCR and sequencing

Following amplification with the 25/396 primer pair, 11 of the 15 skeletal muscle samples were positive (73%): 5 Mexican great-tailed grackles, 4 bronzed cowbirds, and 2 stripe-headed sparrows.
Raw sequences (sense and antisense) of one specimen from each species were included in the study. The sequences were edited, and the fragment lengths were 338 bp for G1, 338 bp for T1, and 323 bp for Z5. Alignment among the 3 sequences was carried out, and 100% similarity was observed (fig. 5). In fact, all 3 sequences were identical. As the sequencing results contained only 323 bp for the Gt5 sequence, the final 15 nucleotides from the other 2 samples were not aligned with this sequence.

Using the Basic Local Alignment Search Tool (BLAST) from GenBank, a comparison of the sequences obtained in this study and the sequences contained in the database was performed. A 96% similarity was found between the sequences here and the sequences corresponding to the following GenBank accession numbers: AY627839, AY627841, AY627842, AY627845, AY627848, AY627850, AY627851, AY627859, AO93159 and AY627852. All sequences corresponded to the fragment of the genome of Sarcocystis neurona with the exception of AY627852, which corresponded to a fragment of the genome of Toxoplasma gondii and AF093159, which corresponded to a fragment of the genome of S. falcata.

**Phylogenetic and restriction fragment length polymorphism (RFLP) analysis**

The sequences obtained in this study show a 100% homology between them. You may also notice that they are topologically more distant among the sequences.
Figure 5. Sequences of a specimen from each species included in the study: stripe-headed sparrow (G1), bronzed cowbird (T1), and great-tailed grackle (Z5). Sequences were edited, and the lengths of the sequences were 338 bp for G1, 338 bp for T1, and 323 bp for Z5. Alignment between the three sequences was carried out, and 100% similarity was observed.

Sarcocystis were easily visualized in striated muscle that exhibited thin-walled parasitic cysts that measured between 0.3 and 0.5 cm in length. These observations differ from those reported by Mansfield et al. (2008). In that study, the investigators analyzed 381 brown-headed cowbirds (Molothrus ater) from the United States and noted macroscopically visible Sarcocystis only in the legs. In addition, the histopathological examination revealed 2 types of cysts: thin-walled and thick-walled. By electron microscopy, the thick-walled cysts were identified as S. falcatula, the thin-walled cysts were identified as S. neurona, and these findings were confirmed by PCR-RFLP. In the present study, macroscopic detection of Sarcocystis sp. parasites was common, possibly because the majority of the birds presented with chronic parasitic infection with mature Sarcocystis in the muscle fibers and degenerate bradyzoites in the interior, similar to the pattern in birds infected with
Sarcocystis in wild birds of Mexico

S. falcatula. Presentation can be acute or chronic, depending on the affected bird. The acute presentation is generally observed in Old World psittacines and in pigeons and causes high mortality associated with pneumonia and encephalitis without the development of parasitic cysts. The chronic presentation occurs in American Passeriformes, which are intermediate hosts. No mortality is observed, and the disease is characterized by the formation of parasitic cysts in skeletal muscle, without a consequent inflammatory reaction (Villar et al., 2008). Histopathologic identification is subjective because S. neurona and S. falcatula have similar morphological characteristics, and literature data reports are inconclusive regarding the wall thickness of the parasitic cyst. Previously, however, Dubey et al. (2001e) reported the presence of a mature parasitic cyst with a 1 to 1.5-µm wall in the cerebellum of an ibis (Carphibis spinicollis), corresponding to Sarcocystis neurona-like parasites and

Figure 6. Neighbor-joining phylogenetic tree constructed from a matrix generated by the Kimura 2-parameter method using 1000 bootstrap replicates. The obtained sequences are identical and are phylogenetically distant from any previously reported sequence.
similar to the protozoa described in this study. In this work, electron microscopy was only used to visualize Sarcocystis in great-tailed grackles, and the ultrastructural characteristics of the wall and bradyzoites coincide with the descriptions of *S. neurona* in bronzed cowbirds (Mansfield et al., 2008), horses (Mullaney et al., 2005), mice, and cell culture (Speer and Dubey, 2001). In these reports, the main characteristic for the ultrastructural identification of *S. neurona* is the accumulation of micronemes in the anterior third of the conoid end and a granular wall with numerous microtubules in villar protrusions. These studies suggest that the Sarcocystis parasites observed in pectoral muscles, legs, and wings of the three species of birds (wild great-tailed grackles, bronzed cowbirds, and stripe-headed sparrows) are similar to *S. neurona*.

Regarding the molecular identification of Sarcocystis in great-tailed grackles, bronzed cowbirds, and stripe-headed sparrows using PCR-RFLP (Tanhauser et al., 1999) with the 25/396 primers and *Hinf* I and *Hind* III restriction enzymes, a positive amplification product of 334 bp was observed. These products contained two *Hinf* I cutting sites in the same positions as the sequences of *S. neurona*. However, no recognition sites for *Hind* III were present in the 334-bp band. This finding is consistent with the banding pattern of *S. neurona* that was previously reported (Tanhauser et al., 1999; Mansfield et al., 2008; Mullaney et al., 2005; and Elsheikha et al., 2005).

Alignment among the 3 sequences was carried out, and 100% similarity was observed. A 96% similarity was found between the sequences studied and *S. neurona*. An additional cutting site that was not detected using PCR-RFLP was detected by sequencing. The generated fragments were approximately 140, 108, 62, and 24 bp. The additional cut by the *Hinf* I enzyme may be due to a mutation or insertion of bases, due to independent genetic evolution during geographic isolation, as shown in the phylogenetic tree where the analyzed sequences are topologically separated between the US and South America. A similar result was observed by Elsheikha et al. (2005). These investigators examined the sequences of 25/396 fragments from 10 *Sarcocystis neurona* samples isolated and found mutations and base insertions, providing evidence for the presence of closely related genetic variants of *S. neurona* existing within the US and South America. Monteiro et al. (2013) suggest that it is possible that genetic groups of *S. neurona* and *S. falcatula* may exchange highly divergent alleles in sexual recombination.

However, because 25/396 marker sequence similarity is highly conserved among apicomplexan protozoans (*Sarcocystis neurona*, *S. falcatula*, *Toxoplasma gondii*, *Neospora caninum*), errors can occur in the interpretation of phylogenetic relationships. This finding emphasizes the importance of using more than just genetic or DNA markers for a robust phylogenetic analysis. In addition, these DNA studies should be supplemented with biological studies because, for example, mice

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**Figure 7.** RFLP analysis of a *Sarcocystis* sp. parasite. A 334-bp fragment was amplified with the 25/396 primers. Lanes 1 and 4: PBR322/DNA/BsuRI (HaeIII) molecular weight marker; lanes 2 and 5: bronzed cowbird and great-tailed grackle samples treated with HindIII; lanes 3 and 6: bronzed cowbird and great-tailed grackle samples digested with *Hinf* I.
are more susceptible to *S. neurona* but are resistant to *S. falcata* (Dubey and Lindsay, 1998), while Australian parakeets are susceptible to *S. falcata* and are not affected by *S. neurona* (Marsh et al., 1997b).

**Conclusions**
The morphological, ultrastructural and PCR-RFLP characteristics suggest that the parasitic cysts observed in the studied birds are *S. neurona*. However, in the phylogenetic tree, sequences are topologically distant from the published sequences of *S. neurona* from the United States and South America, which suggests that this may be a new subspecies of *S. neurona*.

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**Conflicts of interest**
Gary García Espinosa is head of the Departamento de Medicina y Zootecnia de Aves, Facultad de Medicina Veterinaria y Zootecnia. The other authors declare that they have no conflicts of interest.

**Author contributions**
Félix Domingo Sánchez Godoy: Designed the research and wrote the article.
Fernando Chávez Maya and Adriana Méndez Bernal: Contributed new reagents and analytical techniques.
Gary García Espinosa and Elizabeth Morales Salinas: Analyzed the data and wrote the article.
Cristina Guerrero Molina and Néstor Ledesma Martínez: Analyzed the data.

**References**
2) Box ED, Duszynski DW. 1978. Experimental transmission of *Sarcocystis* from icterid birds to sparrows and canaries by sporocysts from the opossum. *Journal of Parasitology* 64:682-688.
4) Butcher M, Lakritz J, Halaney A, Branson K, Gupta GD, Kreeger J, Marsh AE. 2002. Experimental inoculation of domestic cats (Felis domesticus) with Sarco-

5) Cheadle MA, Tanhauser SM, Dame JB, Sellon DC, Hines M, Ginn PE, MacKay
RJ, Greiner EC. 2001a. The nine-banded armadillo (Dasypus novemcinctus) is an intermediate host for Sarco-

6) Cheadle MA, Yowell CA, Sellon DC, Hines M, Ginn PE, Marsh AE, MacKay RJ,
Dame JB, Greiner EC. 2001b. The striped skunk (Mephitis mephitis) is an intermediate host for Sarco-

7) Dame JB, MacKay RJ, Yowell CA, Cutler TJ, Marsh A, Greiner EC. 1995. Sarco-
cystis falcata from passerine and psittacine birds: Synonymy with Sarco-

8) Dubey JP, Lindsay DS. 1998. Isolation in immunodeficient mice of Sarco-
cystis neurona from opossum (Didelphis virginiana) faeces, and its differentiation from Sarco-

9) Dubey JP, Davis SW, Speer CA, Bowman DD, De Lahunta A, Granstrom DE,
Topper MJ, Hamir AN, Cummings JF, Suter MM. 1991. Sarco-

10) Dubey JP, Johnson GC, Bermudez A, Suedmeyer KW, Fritz DL. 2001e. Neural Sarcocystosis in a Straw-necked Ibis (Carphibis spinicollis) associated with a Sarco-
cystis neurona-like organism and description of muscular Sarco-
cystis of an unidentified Sarco-

11) Dubey JP, Lindsay DS. 1999. Sarco-

12) Dubey JP, Lindsay DS, Rosenthal BM, Kerber CE, Kasai N, Pena HF, Kwok OC,
Shen SK, Gennari SM. 2001b. Isolates of Sarco-
cystis falcata-like organisms from South American opossums Didelphis marsupialis and Didelphis albiven-
tris from Sao Paulo, Brazil. Journal of Parasitology 87:1449-1453.

13) Dubey JP, Lindsay DS, Saville WJ, Reed SM, Granstrom DE, Speer CA. 2001a. A review of Sarco-

14) Dubey JP, Lindsay DS, Venturini L, Venturini C. 2000. Characterization of Sar-

15) Dubey JP, Rosenthal BM, Speer CA. 2001c. Sarco-

16) Dubey JP, Saville WJ, Stanek JF, Lindsay DS, Rosenthal BM, Oglesbee MJ, Rosyp-
al AC, Njoku CI, Stich RW, Kwok OC, Shen SK, Hamir AN, Reed SM. 2001d. Sarco-
cystis neurona infections in raccoons (Procyon lotor): evidence for natural infection with sarcocysts, transmission of infection to opossums (Didelphis virginiana), and experimental induction of neurologic disease in raccoons. Veterinary Parasitology 100:117-129.


30) Monteiro RM, Keid LB, Richtzenhain LJ, Valadas SY, Muller G, Soares RM. 2013. Extensively variable surface antigens of *Sarcocystis* spp. infecting Brazilian marsupi-
als in the genus *Didelphis* occur in myriad allelic combinations, suggesting sexual recombination has aided their diversification. *Veterinary Parasitology* 196:64-70.


