Sarcocystis cruzi (Hasselmann, 1923) Wenyon, 1926: Redescription, molecular characterization, and deposition of life cycle stages specimens in the Smithsonian Museum*

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Running Head: Redescription of Sarcocystis cruzi

*Dedicated to Dr. Ronald Fayer, ex-Director of Animal Parasitology Institute (now Animal Parasitic Diseases Laboratory), Beltsville Agricultural Research Center, Beltsville, MD 20705-2350, USA for his seminal contributions to the biology of Sarcocystis infections of livestock.

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Abstract

Currently, seven named *Sarcocystis* species infect cattle: *Sarcocystis hirsuta*, *S. cruzi*, *S. hominis*, *S. bovifelis*, *S. heydorni*, *S. bovini*, and *S. rommeli*; other, unnamed species also infect cattle. Of these parasites of cattle, a complete life cycle description is known only for *S. cruzi*, the most pathogenic species in cattle. The life cycle of *S. cruzi* was completed experimentally in 1982, before related parasite species were structurally characterized, and before the advent of molecular diagnostics; to our knowledge, no archived frozen tissues from the cattle employed in the original descriptions remain for DNA characterization. Here, we isolated DNA from a paraffin-embedded kidney of a calf experimentally infected with *S. cruzi* in 1980; we then sequenced portions of 18S rRNA, 28S rRNA, COX1 and Acetyl CoA genes and verified that each shares 99% to 100% similarity to other available isolates attributed to *S. cruzi* from naturally infected cattle. We also reevaluated histological sections of tissues of calves experimentally infected with *S. cruzi* in the original description, exploiting improvements in photographic technology to render clearer morphological detail. Finally, we reviewed all available studies of the life cycle of *S. cruzi*, noting that *S. cruzi* was transmitted between bison (*Bison bison*) and cattle (*Bos taurus*) and that the strain of parasite derived from bison appeared more pathogenic than the cattle strain. Based on these newfound molecular, morphological, and physiological data we thereby redescribed *S. cruzi* and deposited reference material in the Smithsonian Museum for posterity.

Keywords: *Sarcocystis cruzi*, life cycle, history, cattle, molecular, bison
Introduction

Cattle are routinely infected with parasites in the genus *Sarcocystis*. The taxonomy of *Sarcocystis* species in cattle remains debated; at least seven named species occur: *S. cruzi*, *S. heydorni*, *S. bovini*, *S. hirsuta*, *S. rommeli*, *S. hominis*, and *S. bovifelis*. Molecular evidence suggests the occurrence of additional species (reviewed in Dubey and Rosenthal, 2023). Among these species of bovine *Sarcocystis*, a complete description of the life cycle is known only for *S. cruzi*. The life cycle of *S. cruzi* was completed in 1982 (Dubey, 1982a) experimentally in cattle before the advent of molecular diagnostics (Dubey et al., 1989); to our knowledge, no archived frozen tissues from the cattle employed in the original descriptions remain for DNA characterization. Current DNA characterization is based on naturally infected cattle (Rosenthal et al., 2008; Gjerde, 2016; reviewed in Dubey and Rosenthal, 2023). For genotyping, several genes have been employed to characterize *S. cruzi* from naturally infected cattle including small subunit ribosomal RNA (18S rRNA) (Holmdahl et al., 1993; Rosenthal et al., 2008; Jehle et al., 2009; Gjerde, 2016; Gjerde et al., 2016), large subunit ribosomal RNA (28S rRNA) (Gjerde, 2016); internal transcribed spacer of rDNA (ITS1 and ITS2) (Rosenthal et al., 2008; Gjerde, 2016), and mitochondrial genes, cytochrome c oxidase subunit (COX1) (Gjerde, 2013; Gjerde, 2016; Gjerde et al., 2016). Surveys based on these were summarized previously (Dubey et al., 2016; Dubey and Rosenthal, 2023). Additionally, Doi et al. (2023) reported Acetyl-CoA synthetase gene marker in *S. cruzi* from the cardiac muscles of cattle from Saitama Prefecture.

Here, we reevaluated studies on the life cycle of *S. cruzi*, identified additional morphological details of the developmental stages, redescribe the parasite, provide the first molecular characterization from experimentally infected cattle, and deposit life cycle stages in the Smithsonian Museum for future reference.
History and background

Before the discovery of the obligatory two-host life cycle of *Sarcocystis* species in 1972, only one species of *Sarcocystis* (*Sarcocystis fusiformis*) was recognized in cattle (Heydorn and Rommel, 1972; Rommel and Heydorn, 1972; Rommel et al., 1972). Then, researchers believed *S. fusiformis* parasitized both cattle (*Bos taurus*) and water buffalo (*Bubalus bubalis*). Currently, the name *S. fusiformis* is restricted to the macroscopic sarcocyst-forming species in water buffalo (*Bubalus bubalis*); no species of *Sarcocystis* sharing its morphology has yet been identified in cattle (Dubey et al., 2014; Dubey et al., 2015b).

Subsequent investigations in Germany, identified three species of *Sarcocystis* infecting cattle: one transmitted exclusively by canids, one exclusively by cats, and the third exclusively by humans. Based on these findings, Heydorn et al. (1975b) proposed new names for species of *Sarcocystis* in cattle; *S. bovicanis* for the species transmitted via dogs, *S. bovifelis* for the species transmitted by cats, and *S. bovihominis* for the species transmitted by humans. An intense debate followed the proposal to rename these species. Scientists from Germany and their collaborators (reviewed in Dubey, 2022) sought a rational system, combining information from the intermediate and definitive host (e.g., *S. bovicanis*). They suggested new names for *Sarcocystis* species of livestock (reviewed in Dubey, 2022). However, Levine (1977) upheld the principle of nomenclatural priority. This principle seeks stability in scientific communication by retaining valid names for described species. Levine (1977) concluded that the previously used names for *Sarcocystis* species must be retained, even though the descriptions of species were incomplete; he assigned *S. cruzi* for *S. bovicanis*, *S. hirsuta* for *S. bovifelis*, and *S. hominis* for *S. bovihominis*.

History and biology of *S. cruzi*

Levine (1977) credited Hasselmann (1926) with naming *S. cruzi*. Levine was unaware of an
earlier paper by Hasselmann who named it as *Miescheria cruzi* in a brief note in 1923 (reviewed in Dubey and Rosenthal, 2023). Neither of these papers by Hasselmann (1923, 1926) contains a morphological description of the parasite. Hasselmann found the parasite in all 55 hearts from Salvador (Bahia) and São Paulo, Brazil (Hasselmann, 1923). He called the parasite *cruzi* because it resembles *Trypanosoma cruzi* in histological sections. Hasselmann was unaware of *Sarcocystis* species in cattle. Wenyon (1926) transferred the genus *Miescheria* to the genus *Sarcocystis*, resulting in *Sarcocystis cruzi* (Hasselmann, 1923) Wenyon, 1926. Hasselmann, a pathologist, focused on host reactions rather than parasite morphology. In his 1926 paper, Hasselmann (1926) called the parasite *Miescheria crusi* but the name *cruzi* has precedence over *crusi* (Dubey and Rosenthal, 2023). *Sarcocystis cruzi* sarcocysts are thin walled (<1.0 μm) and commonly parasitize the myocardium. Until 2014, all sarcocysts with thin walls were considered *S. cruzi*. In 2015, a new thin-walled species, *S. heydorni*, was recognized; importantly, its definitive host is human, not canid (Dubey et al., 2015a).

Most experiments involving the life cycle of *S. cruzi* were performed in Germany and USA; here we review these studies.

**Experimental transmission of bovine Sarcocystis to dogs in Germany**

In the pioneering study by scientists in Germany, muscles from cattle esophagi naturally infected with *Sarcocystis* were homogenized and fed to 12 dogs (Heydorn and Rommel, 1972). Dogs excreted sporocysts after a prepatent period of 9 or 10 days and patent periods of 57-71 days; sporocysts were 13.9-11.7 x 6.2-10.8 μm. The sporocysts were in the lamina propria of small intestine; the same dogs were refed infected beef when the sporocysts were no longer detected in feces; dogs excreted sporocysts 10-12 days later after ingesting infected beef for the second time. In these studies, the parasite was called *Isospora bigemina* (Heydorn and Rommel, 1972).
Experimental infection of cattle with *Sarcocystis* was performed by Gestrich *et al.* (1975). The proposal to name this species *S. bovicanis* by (Heydorn *et al.*, 1975b) was primarily based on this experiment. This paper does not have an English summary; therefore, we provide details here.

Eight, 7 to 8-weeks-old calves were orally inoculated with 50,000 (1 calf), 100,000 (2 calves), or 2,000,000 (5 calves) sporocysts; the calves were necropsied days 28 (2 calves), 29 (2 calves), 30 (1 calf), 48 (1 calf), 62 (1 calf) post inoculation (p.i.), and muscle biopsy were taken from 1 calf, days 160 and 194 p.i. (Gestrich *et al.*, 1975). All five calves inoculated with 2,000,000 sporocysts died of acute sarcocystosis; schizonts were found in blood vessels. Thin walled (<0.5 µm) sarcocysts were detected in muscles of examined days 48, 62, and 98 p.i.; no mention was made concerning days 160 and 194 p.i. (Gestrich *et al.*, 1975).

Subsequently, development of *S. cruzi* sarcocyst was studied ultrastructurally by Heydorn *et al.* (1975a) and Mehlhorn *et al.* (1975) using four calves inoculated with 50,000 to 80,000,000 sporocysts. Details of inoculation, dose and necropsy are provided here for the benefit of readers who may not have access to these papers in German. Calf 1 was inoculated with 80,000,000 sporocysts and necropsied day 27 p.i. Calf 2 was inoculated with 50,000 sporocysts, and reinoculated with 100,000 on 14 days later; the same calf received 250,000 sporocysts weekly for 7 weeks, and 2,000,000 on day 60 and then necropsied day 150 from the initial inoculation of sporocysts (Mehlhorn *et al.*, 1975). The third calf was inoculated with 100,000 sporocysts and with 2,000,000 sporocysts on day 42; the calf was necropsied on day 76 p.i. (day 34 after the second dose). The fourth calf received 100,000 sporocysts and was necropsied day 62 p.i. Sarcocyst formation was reported beginning day 34 p.i. but the duration of infection remains unclear because the calf was inoculated several times. At day 62 p.i., sarcocysts contained mainly metrocytes and a few bradyzoites. Sarcocysts at day 76 p.i. contained bradyzoites. Details of sarcocyst development were given by Mehlhorn *et al.* (1975).
and Heydorn et al. (1975a) described structures of bradyzoites (they called them merozoites); both studies are based on the same calves.

**Summary of research on S. cruzi infections in cattle at USDA**

Pioneering research on bovine sarcocystosis was performed at the Animal Parasitology Institute (now Animal Parasitic Diseases Laboratory, APDL), United States Department of Agriculture (USDA) Beltsville, Maryland under the direction of Dr. Ronald Fayer (now retired); these studies are summarized here (Table S1). These studies established for the first time that *S. cruzi* is pathogenic for cattle. Calves orally inoculated with 250,000 to 1,000,000 sporocysts developed acute systemic disease and intravascular schizonts were found in almost all organs of calves (Fayer and Johnson, 1973). Cows inoculated with sporocysts from dogs aborted and developed acute sarcocystosis (Fayer et al., 1976a). Results of this experiment also linked the mysterious Dalmeny disease reported by Corner et al. (1963) to acute *S. cruzi* infection. Other studies on the biology of experimental *S. cruzi* infection are also summarized in Table S1.

Fayer and associates also described gametogony and sporogony of *S. cruzi* in intestines of dogs fed beef from experimentally or naturally infected calves and its transmission to different hosts (Fayer, 1974; Fayer, 1977a; Fayer et al., 1976b; Sheffield and Fayer, 1980); these studies are discussed later.

In the 1980’s, one of us (JPD), studied the full life cycle of a *S. cruzi* transmissible from coyotes to cattle in Bozeman, Montana, USA. For the life cycle studies with the Bozeman isolates of *S. cruzi*, newborn calves and coyotes were raised in captivity to strictly monitor their diets (Dubey, 1982a) (Table S2).

While reevaluating histologic slides from the 1982 study (Dubey, 1982a), a paraffin block of a kidney of a calf 383 fed 150,000,000 *S. cruzi* sporocysts from coyote no. 27
(Table S2) and euthanized day 24 p.i. was located, and DNA was extracted. Here, we report the first genetic characterization of *S. cruzi* from this experimentally infected kidney.

**Materials and methods**

**Specimens studied**

In view of the recent development and recognition of several *Sarcocystis* species in cattle, we reevaluated histologic sections and obtained new morphological and genetic data from parasites used by Dubey (1982a). Life cycle stages were photographed using a digital DP73 camera (Olympus Optical Ltd.) fitted on an Olympus AX 70 microscope (Olympus Optical Ltd., Tokyo, Japan), and deposited in the Smithsonian Museum.

**Inoculation of calves at Bozeman**

Materials and methods were the same as described in detail previously (Dubey, 1982a). Briefly, 38 Holstein-Friesian newborn calves were inoculated with 275,000 to 500,000,000 sporocysts of seven isolates (BI-B7) of *S. cruzi* obtained from laboratory-raised coyotes fed naturally infected hearts of cattle raised in Montana (Tables S2, S3). Calves were necropsied between days 4 and 153 p.i. (Table S2). Parasitemia was determined as described elsewhere (Dubey, 1982c).

A total of 19 coyotes were used in this study and they were obtained from dens within a week of birth and were raised on canned milk and canned or dry dog food (Dubey, 1982c). Four coyotes were necropsied for histological studies. For the study of early stages of gametogenesis, one coyote was killed at 6 h and another at 12 h after ingesting a single meal of infected meat. A third coyote was fed two meals, 6 h apart, of infected meat and killed 18 h after ingesting the last meal. The infected meat consisted of tongue, heart, esophagus, and other skeletal muscles of experimentally inoculated calf 422 (Table S3). A fourth coyote was
euthanized 16 days after feeding experimentally infected beef (Table S3).

To assess the potential of the coyote as definitive host of *S. cruzi* in the wild, 15 coyotes fed infected beef (Table S3) were euthanized, the small intestine mucosa and the submucosa was scraped from the muscular layers, homogenized in a blender, and sporocysts were collected as described before (Dubey, 1980). Additionally, sporocysts were collected from sugar flotation of the entire daily feces of each coyote (Table S3).

*Transmission of dog transmitted *S. cruzi* infected with Beltsville strain to coyote and cattle*

To study if the Beltsville strain of *S. cruzi* derived from a cattle-dog cycle was also transmissible to coyotes, beef from a calf inoculated with the dog-derived *S. cruzi* sporocysts was sent cold by air by Dr. Fayer from Beltsville, Maryland to Bozeman, Montana where it was fed to a laboratory-raised coyote; the coyote started excreting sporocysts day 9 p.i. The coyote was euthanized on day 12 p.i. and 50,000,000 sporocysts from the intestinal digest were fed to a newborn calf and the calf was euthanized on day 15 p.i.

At necropsy, virtually all internal organs of cattle were studied histologically as described (Dubey, 1982a). Portions of tissues were fixed in 10% (v/v) neutral buffered formalin (NBF). Similar procedures were followed for tissues of coyotes. Selected tissues were also fixed in Bouin's fluid (BF) or Helly's fixative (HF). Paraffin-embedded sections were cut at 5 µm. Selected tissues were embedded in glycol methacrylate (called plastic here) and sectioned at 3 µm. Sections were stained with hematoxylin and eosin (HE), Heidenhain's iron hematoxylin (IH) or periodic acid-Schiff's-hematoxylin (PASH). A similar process was used for tissues of coyotes. The histological slides evaluated were those from the 1980 study.

*Molecular characterization*

Recently, we performed molecular characterization of parasites that had been embedded, four
decades ago, in a paraffin block containing kidney tissue of experimentally infected calf no. 383 (Table S2). Genomic DNA extraction was performed using the Qiagen DNeasy® Blood and Tissue Kit and Qiagen QIAmp® DNA FFPE Tissue Kit according to the manufacturer’s instructions and stored at -20°C for further use. The quality and quantity of the DNAs were calculated using a Qubit 4 Fluorometer (Invitrogen, ThermoFisher Scientific, USA).

For genotyping, *S. cruzi* specific primers (18S rRNA, 28S rRNA, COX1 and ACS genes) were designed using Primer3 version 4.1.0 (Untergasser *et al.*, 2012) and NCBI primer blast using *S. cruzi* as reference sequence from NCBI (Table S4). The 25 µL of PCR mix consisted of a 2 µL DNA template (120 ng/µL), 12.5 µL of Platinum Hot Start PCR Master mix (Invitrogen, USA), 1 µL of 10 pmol/µL of each primer (IDT, USA) (Table S4) and 8.5 µL of molecular grade water. Templates were denatured at 94°C for 3 min, followed by 35 cycles of denaturation at 94°C for 30 s, annealing at 60°C for 30 s, and elongation at 68°C for 20 s min; these cycles were followed by a terminal elongation at 68°C for 5 min. The PCR products were analyzed on a 2 % agarose gel running at 100 V for 30 mins and the size of the amplicons was estimated by comparison with the 100 bp DNA Ladder (Promega, USA).

The PCR products were purified using the ExoSAP method (Bell, 2018) and sequenced using the Big Dye Terminator v3.1 cycle sequencing kit (Applied Biosystems™, USA), following the manufacturer’s instructions, in an ABI 3100 genetic analyzer (Applied Biosystems™, USA). The sequences were deposited in the Genbank database and accession numbers were obtained (Table S4).

**Phylogenetic analysis**

Phylogenetic analyses were performed independently on nucleotide sequences of 18 rRNA, 28S rRNA, COX1 and acetyl-CoA synthetase genes to evaluate the evolutionary relationship between *S. cruzi* and other closely related *Sarcocystis* spp. The segments obtained were
trimmed aligned by MAFT alignment in Geneious software followed by ClustalW multiple alignment in MEGA7 (Kumar et al., 2016). Any ambiguous bases were clarified using the respective chromatograms. Subsequently, the consensus sequence of the SSU rDNA was analyzed using a standard online Basic Local Alignment Search Tool (BLAST) (https://blast.ncbi.nlm.nih.gov/Blast.cgi) against the genetic dataset of various S. cruzi species available at NCBI. For generating consensus trees, we first blasted the edited sequences against the NCBI database and downloaded the most closely related Sarcocystis species. This allowed us to characterize a well-supported clade of S. cruzi to which our sequences belong. The sequences were subjected to phylogenetic analysis using MEGA 7 software (Kumar et al., 2016). A codon-based ClustalW multiple alignment of all sequences was performed in MEGA X. Neighbor-Joining trees were obtained. To assess the robustness of the phylogenetic trees, 1000 bootstrap replications were employed in all analyses. All codon positions were used.

The molecular analysis was performed against the existing partial sequences previously obtained from different isolates of S. cruzi from cattle, and eight haplotypes from Sarcocystis species identified in Bovidae and Cervidae. All sequences were truncated slightly at both ends to preserve the homologous nucleotide positions for further analysis. Neospora caninum and Toxoplasma gondii were used as out-groups. For Acetyl CoA synthetase gene, we downloaded the sole available haplotype (acc. no. LC729541) and found it identical to our isolate of S. cruzi.

**Results**

The following description of the life cycle stages is from the coyote-cattle cycle of S. cruzi as reported previously (Dubey, 1980; Dubey, 1982a; Dubey, 1982c; Dubey et al., 1982), and in the present study.
Intravascular stages

Terminology used

Different terms have been used to describe *S. cruzi* asexual stages. Historically, coccidian (*Eimeria* spp.) asexual stages are called schizonts (Dubey, 2020). In schizogony, the sporozoite nucleus divides into four or more nuclei (> 1000 in *Eimeria bovis*) before merozoites are formed. However, in some other coccidians (e.g., *Toxoplasma gondii*) some stages divide into two daughter organisms by endodyogeny. To encompass all coccidians, Levine (1973), an expert on coccidian parasites, called asexual stages as meronts instead of schizonts. Dubey (1982a) followed Levine and called *S. cruzi* intravascular stages as meronts. However, other authors (Fayer and Johnson, 1973; Gestrich *et al*., 1975) called them schizonts. For uniformity, we have now used the term schizonts for *S. cruzi*. We are aware of the differences in asexual cycle in *Eimeria* versus *Sarcocystis*. Unlike *Eimeria*, *S. cruzi* schizonts divide by endopolygeny, where the parasite nucleus becomes multilobed but remains interconnected (Dubey *et al*., 2016).

In *S. cruzi*, at least three generations of intravascular stages were identified. To trace the development of earliest stages, two calves each fed 500,000,000 sporocysts were euthanized on day 4 and 7. No development was found earlier than day 11 p.i. At day 7 p.i., sporozoite-like zoites were found within intravascular leukocytes and in endothelial cells of arteries within mesenteric lymph nodes. After an intensive search, a total of 36 zoites were found in numerous histological sections of the mesenteric arteries; 31 of these were in endothelial cells, 4 were in macrophage-like leukocytes in vascular lumen and 1 was apparently extracellular (Dubey, 1983). Immature schizonts were found in mesenteric lymph node arteries day 11 p.i. (Figure 1A) and mature first generation schizonts were found days 15-26 p.i. (Figure 1B). The development of schizonts was asynchronous. Schizonts averaged 41.0 x 17.5 µm and contained up to 350 merozoites.
Second generation schizonts were found in capillary endothelial cells of many organs but were most numerous in kidneys days 19-46 p.i. (Figure 2). The shape and size of schizonts varied, depending on the tissue parasitized; renal schizonts (19.6 x 11.0 µm) were shorter than intramuscular schizonts (25.0 x 11.0 µm).

*Parasitemia and multiplication in blood.*

Two waves of parasitemia were recognized. Few merozoites were found in blood day 17 p.i., coincident to maturation of first generation schizonts. The second wave of parasitemia occurred days 24-46 p.i. Most (97.2%) merozoites were within lymphocytes or monocytes (Figure 3A). The number of merozoites ranged from 0.14 to 9.8 per ml of peripheral blood (Dubey, 1982c). A few merozoites divided in monocytes, apparently by endodyogeny (division of into 2 nuclei). All dividing merozoites were intracellular these merozoites were 6.8 x 3.2 µm in Giemsa-stained smears of blood.

Sarcocysts were found within myocytes, most commonly in the heart and tongue. The earliest sarcocysts were found day 42 p.i. and contained 1 or 2 metrocytes. At day 46 p.i., sarcocysts were mostly in the myocardium and contained up to 8 metrocytes. At day 55 p.i., sarcocysts were < 20 µm and contained only metrocytes (Figure 3C, D). Metrocytes were round to elongate, and usually stained faintly with H and E stain. At day 67 p.i., sarcocysts were up to 360 µm long and 30 µm wide; they had only metrocytes. At day 86 p.i., sarcocysts contained both metrocytes and bradyzoites and were infectious for coyotes. The sarcocyst wall was thin (< 1 µm) and smooth. Sarcocysts remained microscopic in all calves, including the last day of observation, day 153 p.i. (Figure 3 E, F). Bradyzoites were 11 x 3.0 µm in smears and 8.2 x 3.0 µm in sections (Figure 4A). Degenerating sarcocysts were not found.

Only a few sarcocysts were found in brain and spinal cord tissues. Immature sarcocysts containing metrocytes were found in smooth muscles of the small intestine, urinary bladder,
rumen, and abomasum (Dubey, 1982a).

Only one morphologic type of sarcocysts, corresponding to *S. cruzi*, was found in calves inoculated with sporocysts from coyotes and the development correlated with day p.i. The identity of *S. cruzi* sarcocysts was confirmed ultrastructurally (Dubey, 1982a).

**Gametogony**

Gamonts were seen in parasitophorous vacuoles of goblet cells in the epithelium of the small intestine at 6 h p.i. (Figure 4 B, C). The host cell nucleus was indented, misoriented, but not hypertrophied (Figure 4 B, D, E, G). Numerous free bradyzoites were seen in contents of stomach, small intestine, and colon (Figure 4A). At 6 h, gamonts were found throughout the tips of the villi of the small intestine of the coyote. Most of the round to elongate gamonts were found in goblet cells; up to three gamonts were seen in a single goblet cell (Figure 4B). A few gamonts were observed in the lamina propria just below the basement membrane (Figure 4H). Fewer than 5% of gamonts were microgamonts. All stages of microgametogony, from the single nucleated stage to fully developed gametes were seen (Figure 4F-I). Mature microgamonts were 7.1 x 4.8 μm and contained 3-11 slender gametes, which were 3.5-4.0 μm long and less than 0.5 μm wide (Figure 4I). Macrogamonts were 8.0 x 7.3 μm and contained PAS-positive granules.

At 12 h, stages resembled those at 6 h. Additionally, two oocyst-like structures were seen in Giemsa-stained smears of the intestine. By 24 h, most gamonts had become oocysts. In the coyote killed day 16 p.i., only oocysts were found; they were in the lamina propria (Figure 5 A-D). Unsporulated oocysts contained a central or a band nucleus (Figure 5B). Oocysts grew to about 20 x 19 μm before sporulation began. All stages of sporulation, from binucleated sporocysts to fully sporulated sporocysts were seen day 16 p.i. (Figure 5D). Living sporozoites were around 10 x 3 μm whereas those in sections were around 9 x 2 μm.
The prepatent period was 9 days in all 16 coyotes (9 coyotes fed naturally infected beef, 7 coyotes fed experimentally infected beef, including 1 coyote fed the beef infected with the Beltsville strain) (Table S3).

Transmission of dog transmitted S. cruzi infected with Beltsville strain to coyote and cattle in Bozeman

Coyote No. 30 fed beef infected with the Beltsville strain of S. cruzi started excreting sporocysts on day 9 p.i. and 300,000,000 sporocysts were recovered in intestinal digest when the coyote was euthanized (Table S3). The newborn calf orally inoculated with 50,000,000 sporocysts remained asymptomatic. Many first generation schizonts were found in histological sections of arteries in kidneys and mesenteric lymph nodes when the calf was euthanized day 15 p.i. These schizonts were structurally similar to those described by Fayer (1977b).

Molecular characterization

The sequencing of S. cruzi from the kidney of experimentally infected calf resulted in sequences of 1391 bp for 18S rRNA gene; 583 bp for 28S rRNA; 960 bp for COX1 and 1743 bp for ACS gene with a guanine-cytosine-GC content of 41.6%, 43.2%, 51.9% and 54.8%, respectively. In the phylogenetic analyses conducted using the Neighbor-Joining (NJ) method based on partial 18S rRNA gene sequences (Figure 6), the placement of the two major clades comprising Sarcocystis spp. of cattle, showed some variation depending on the sequence variability and alignment settings, particularly the gap opening and gap extension penalties.

BLAST analysis showed 99% to 100% similarity of the present isolate to other available isolates attributed to S. cruzi from naturally infected cattle. Phylogenetic reconstruction of variation in 18S rRNA sequences, using the Neighbor Joining method, confirmed its uniquely close relationship to other S. cruzi isolates, to the exclusion of other congeneres (Figure 6). The
newly sequenced isolate belongs to a group comprised only of other isolates of *S. cruzi*, with strong bootstrap support, and distinct from sequences from other parasites of cattle including *S. rangi, S. arietianis, S. hircianis, S. tenella, S. alces, S. suihominis, S. bovifelis* and *S. truncata* (Figure 6). Relationships among species generally accord with prior findings (Gjerde, 2013; Gjerde, 2016).

*Bison (Bison bison) as an intermediate host of S. cruzi*

Bison can act as intermediate host of *S. cruzi*. A laboratory-raised coyote (No.13) excreted sporocysts 11 days after feeding bison meat in Montana (Dubey, 1980). Three cattle calves inoculated with 100,000 (1 calf), 700,000 (2 calves) sporocysts from coyote 13 developed patent *Sarcocystis* infections (Fayer *et al.*, 1982). Two calves had diarrhea days 19-25 p.i. One of the two calves fed 700,000 sporocysts became moribund day 28 p.i. and was necropsied. Hemorrhages were found in many organs, particularly on the cecum; there is no record of histological examination of tissues in the paper and these specimens are no longer available for retrospective evaluation. Sarcocysts characteristic of *S. cruzi* were found in histological sections of the two calves necropsied days 63 and 89 p.i. (Fayer *et al.*, 1982).

For experimental transmission of the cattle strain of *S. cruzi*, two newborn laboratory-raised bison were inoculated orally with sporocysts of B1 isolate of *S. cruzi* originally isolated from a laboratory-raised coyote (coyote 25, see Table S3) fed naturally infected beef in Montana. A second passage of this isolate was used to inoculate each bison. For this, sporocysts from coyote 25 were fed to a newborn cattle calf and muscles from the calf were fed to laboratory-raised coyotes (Nos. 38 and 43, see Table S3). Sporocysts from coyotes 38 and 43 were used as inocula for bison (Dubey, 1982b). One six-day old bison was fed 10,000,000 sporocysts. This bison developed transient diarrhea and fever day 14 p.i. then remained asymptomatic until day 25 p.i. The bison was euthanized day 28 p.i. because it had severe
diarrhea and was unable to swallow. Second generation schizonts were found in histological sections of the adrenal cortex and in the lamina propria of small and large intestines. Merozoites were detected in peripheral blood days 25 to 28 p.i.

The second bison was inoculated with 100,000 sporocysts of the B1 isolate when 7-days old; this bison had fever on days 16, and 18 and then again on days 27-30 p.i. (Dubey, 1982b). Sarcocysts were found in muscles when the bison was necropsied day 73 p.i. Merozoites, schizonts and sarcocysts found in the bison tissues were like those of *S. cruzi* in cattle tissues. The unusual findings regarding bison infections were: diarrhea, and the site of second generation schizonts; in cattle, the kidney is the preferential site for second generation schizonts whereas in the bison intestines and adrenal were the parasitized sites. Diarrhea is not a typical clinical feature of acute sarcocystosis in cattle.

In summary, *S. cruzi* was transmissible from bison to cattle and cattle to bison via coyotes. The bison isolate of *S. cruzi* was apparently more pathogenic than the cattle strains and its schizonts were in the intestines than in the kidney most often parasitized by cattle strains. Nothing is known of the genetic features that define strain differences in *S. cruzi*.

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*Taxonomic summary of Sarcocystis cruzi*

*Diagnosis:*

*Intermediate host: Cattle (Bos spp.), Bison (Bison spp.)*

Asexual development in bovine tissues. Two morphologically distinct generations of schizonts directly in endothelial cells of blood vessels, particularly in kidneys 11-46 days p.i. followed by parasitemia and development of merozoites in peripheral blood monocytes. Sarcocysts in parasitophorous vacuoles of myocytes, particularly in the myocardium. Sarcocysts microscopic, < 1 mm long, with thin (< 0.5 um thick) sarcocyst wall.
Definitive hosts: Domestic dog (*Canis domesticus*), coyote (*Canis latrans*), red fox (*Vulpes vulpes*) confirmed.

Gamonts in epithelium, primarily in goblet cells of small intestine. Sporogony in the lamina propria of small intestine. Prepatent period 9 days or more. Sporulated sporocysts ~16 x 10 μm.

Specimens deposited

The specimens were deposited in United States National Parasite Collection in the Division of Invertebrate Zoology and National Museum of Natural History, Smithsonian Institution, Museum Support Center, MRC 534, 4210 Silver Hill Road, Suitland, Maryland 20746, USA, under numbers USNM-(Table 2).

DNA sequences were deposited in Genbank (Table S4).

Discussion

Intravascular stages

In general, the intravascular stages of *S. cruzi* found in dogs fed the Beltsville strain resemble those reported to have developed in coyotes fed infected beef of the Bozeman strains. The minor differences observed likely relate to dose, histological methods, time spent searching for early stages, and age of calves. For example, only a few first generation schizonts were found in calves infected at Beltsville and these were immature (Fayer, 1977b). Using the Beltsville strain cycled through a coyote, numerous first generation schizonts were detected in arteries in kidneys and mesenteric lymph nodes and schizonts contained fully developed merozoites; these observations are probably related to the age of calves and dose; the calves were 3 to 4.5 months old and inoculated with 200,000 sporocysts at Beltsville whereas, a newborn calf was inoculated with 50,000,000 sporocysts at Bozeman.
Here, we summarized development of the ox-coyote cycle of *S. cruzi*. After the sporocysts excyst in cattle gut, the sporozoites reach arteries in the mesenteric lymph nodes, probably via portal circulation. The zoite found in blood day 4 p.i. was probably a sporozoite, as were those found day 7 p.i. The first generation meronts have a predilection for arteries, whereas second generation meronts occur predominantly in capillaries. Both first and second generation meronts develop in cells between the tunica intima and the endothelium (Dubey *et al.*, 1980).

Second generation schizonts were first detected day 19 p.i. and some had matured by day 24 p.i. This indicates a generation time of five or more days. Most of the second generation schizonts were found between day 24 and 29 p.i. Whether more than one multiplication cycle of second generation schizonts occurs was not determined.

Intramuscular schizonts were longer than renal meronts. Whether this phenomenon was related to the stage of development or to the type of host cell parasitized was not determined. Such elongated meronts were not reported by Fayer and Johnson (1973). In addition, individual merozoites were found in muscles days 33-45 p.i. in the present study; these were not reported to occur in the calves examined by Fayer and Johnson (1973), who found individual merozoites in the kidneys.

*Sarcocysts*

Sarcocysts are formed around the sixth week after inoculation of sporocysts. Dubey (1982a) did not find sarcocysts until day 45 p.i. After reevaluation of histological sections 35-44 days p.i., a few sarcocysts were detected days 42 and 44 p.i (present study). Fayer and Johnson (1973) found sarcocysts containing up to 50 zoites in a calf killed day 33 p.i.; it is likely that the sarcocyst observed were from a spontaneous infection acquired from eating contaminated hay. Distinction of sarcocysts from intramuscular schizonts is sometimes difficult unless
sections are stained optimally. However, a parasitophorous vacuole (PV) around the zoites can distinguish them; sarcocysts are located in a PV whereas schizonts are located directly in the host cell cytoplasm without a PV. Structurally, second generation schizonts could be differentiated from early sarcocysts. The PAS-positivity is also a distinguishing feature. Schizonts are PAS-negative but zoites in sarcocysts are PAS-positive. However, PAS positivity of metrocytes in early sarcocysts varies, depending on fixation; Bouin's fluid was better than formalin for preserving PAS-positive polysaccharide granules.

Mehlhorn et al. (1975) also studied the development of sarcocysts. Beginning day 34 p.i., only sarcocysts were found and the sarcocyst wall remained thinner than 1 µm even at day 150 p.i. Sarcocysts at day 76 p.i. were structurally mature but the infectivity to calves was not tested (Mehlhorn et al., 1975). Fayer and Johnson (1974) stated that the sarcocysts on day 54 p.i. contained only metrocytes; no mention was made when bradyzoites developed or when sarcocysts were infectious for dogs. Sarcocysts became infectious between days 67 and 86 p.i. in calves fed sporocysts from coyotes (Dubey, 1982a).

**Gametogony and sporogony**

Gametogony was completed in the coyote intestine between 6 and 12 h p.i. It is likely that gamonts are formed earlier. It is noteworthy that bradyzoites convert to gamonts in goblet cells while still in the surface epithelium. Subsequent development occurs in unidentified cells in the lamina propria. How *S. cruzi* gamonts reach the lamina propria is unknown. Initial gamonts are intracellular in PV, mostly in goblet cells. One hypothesis is that the infected host cells drop down to the lamina propria. Fayer (1974) and Sheffield and Fayer (1980) first described the gametogony and sporogony of *S. cruzi* in the small intestine of dogs fed infected beef. By using transmission electron microscopy, Sheffield and Fayer (1980) documented fertilization of gametes and the formation of the oocyst wall beginning at 12 h. They also mentioned that
gametes were extracellular within the lamina propria but provided no quantitative data. Female gametes contain polysaccharide granules and if the fertilized gametes are indeed extracellular then these likely provide energy for sporulation and survival in the environment. There are no ultrastructural studies on the sporulation of Sarcocystis oocysts. However, Fayer (1974) described sporulation of S. cruzi in intestines of dogs by light microscopy. Macrogamonts were detected day 2 through 13 p.i. He found small PAS positive granules in macrogamonts on day 5 and these grew in size and number until day 13. Sporulating oocysts were seen day 7-13 p.i.. On day 8 p.i., a nuclear band was seen across the center of some oocysts and binucleate sporocysts were identified day 8 p.i. The presence of this nuclear band is confirmed here (Figure 5B). Sporulated oocysts/sporocysts were first detected on day 9 p.i. (Fayer, 1974).

**Prepatent and patent periods**

As stated earlier, Heydorn and Rommel (1972) reported that dogs fed naturally infected beef excreted S. cruzi sporocysts 9 or 10 days later. Fayer (1974, 1977a) evaluated this topic in detail. He reported prepatent periods of 9-33 days, irrespective whether the beef was from experimental or naturally exposed infections. Most dogs excreted sporocysts between day 11 and 16 p.i. The dogs excreted sporocysts intermittently up to 60 days p.i. (Fayer, 1977a). After feeding 500 g of naturally infected beef, 11 dogs excreted a total of between 861,000 and 3,764,000 sporocysts. More sporocysts were excreted by dogs after ingesting experimentally infected beef than naturally exposed beef (Fayer, 1977a). In one dog as many as 20,077,000 sporocysts were excreted (Fayer, 1977a).

*Sarcocystis* infections in beef are common (Dubey et al., 2016). One of the explanations is that dogs are common on cattle farms. We are not aware of any surveys for the presence of S. cruzi sporocysts in feces of farm dogs. In a study of dogs from a Humane Center in Ohio, USA *Sarcocystis* spp. sporocysts were found in 1.8% of 500 dogs; most of these dogs were
probably domestic pets (Streitel and Dubey, 1976).

In contrast to these findings in dogs, all 16 coyotes (9 fed naturally infected beef and 7 fed experimentally infected beef) started excreting sporocysts on day 9 p.i. (Table S3). Thus, the minimum prepatent period is 9 days (not 8 days mistakenly reported in Dubey [1982a]). In the coyote-cattle study at Bozeman, patent period was not determined because coyotes were euthanized days 12-27 p.i. and sporocysts were collected from the digest of entire small intestines. As many as 265,000,000 sporocysts were recovered from coyotes fed naturally infected beef and 300,000,000 sporocysts were present in a coyote fed experimentally infected beef (Table S3). Differences in these studies may be related to the dose, method of sporocyst detection, and duration of observation period. For the study in coyotes, the entire daily feces were sugar-float and tested microscopically. Additionally at necropsy, the entire small intestine was homogenized and digested to recover sporocysts. Sarcocystis sporocysts are trapped in the lamina propria and are released in the intestinal tract intermittently. No inflammatory response to sporocysts in the lamina propria was observed.

The 9-day prepatent period coincides with the completion of sporulation on day 9 (Fayer, 1977a). However, sporogony is asynchronous because at day 16 p.i., unsporulated oocysts were observed in the lamina propria of infected coyotes (Figure 5). Only sporulated oocysts, and free sporocysts are found in feces of dogs or coyotes. Thus, transport of oocysts from the lamina propria to the intestinal lumen is likely regulated by the parasite because no sporocysts were detected in feces until day 9 p.i.

In cattle and coyotes infected at Bozeman, only thin-walled S. cruzi sarcocysts were found; no other coccidian oocysts were detected in the feces of these coyotes (Dubey, 1982a). The unsporulated oocysts seen by Fayer (1974) and Heydorn and Rommel (1972) in feces of dogs fed Sarcocystis infected beef belong to another coccidian, Hammondia heydorni; it cycles through ruminants and canids (Dubey et al., 2002).
**Definitive hosts of S. cruzi**

The domestic dog, coyote, and red fox are proven definitive hosts of *S. cruzi* because sporocysts from these hosts were infectious to cattle, and infected beef proved infectious to laboratory raised dog, coyote, and fox (Table 1). There is one report of a crab-eating fox from Brazil as a definitive host for *S. cruzi*, but it needs confirmation. Wolves and raccoon dogs fed naturally infected beef also excreted *S. cruzi*-like sporocysts, but infectivity of wolf or raccoon-derived sporocysts has not been tested (Table 1). Adult raccoons excreted *S. cruzi*-like sporocysts after ingesting *S. cruzi* infected beef; however, the raccoons were adults when trapped (Fayer et al., 1976b). Subsequently, two studies could not confirm these findings (Table 1). Therefore, we do not consider raccoons as an established definitive host of *S. cruzi*. Although jackal (*Canis aureus*) was mentioned as a definitive host for *S. cruzi* (Dubey and Rosenthal, 2023) and sporocysts have been found in feces of naturally infected jackals (Kirkova et al., 2011; Tulov, 2013; Gherman and Mihalca, 2017), there is no evidence that the species was *S. cruzi*.

Recently, mustelids in Lithuania were reported as possible definitive hosts for four *Sarcocystis* species infecting cattle (*S. cruzi, S. bovifelis, S. hominis, S. hirsuta*). DNA was obtained from sporocysts separated from intestinal scrapings of mustelids and characterized by nested PCR of partial COX1 gene (Prakas et al., 2021). High detection rates of two species, *S. bovifelis* (89.3%) and *S. cruzi* (73.8%) were observed, while *S. hirsuta* (3.6%) and *S. hominis* (1.2%) have been found in only a few animals. The DNA of *S. cruzi* was identified in 77.5% of 40 American mink (*Neovison vison*), 75% of four Beech marten (*Martes foina*), 75% of 20 European pine marten (*Martes martes*), 20% of five European badger (*Meles meles*), and 80% of 15 European polecat (*Mustela putorius*). However, none of these carnivores were previously identified as definitive hosts for *Sarcocystis* species of cattle in other countries (Dubey et al., 2016).
There are only a few reports of the prevalence of *Sarcocystis* sporocysts in wild canids. *Sarcocystis* sporocysts were detected in feces of 21 of 150 (8.0%) coyote feces from Utah and Idaho (Fayer and Johnson, 1975) and in feces of 89 of 169 (52.7%) coyotes from Montana (Dubey *et al*., 1978). In both studies, sporocysts were infectious to cattle.

**Molecular characterization**

Based on our molecular results, we found 100% identity in all the four genes from experimentally infected calf No. 383 with other *S. cruzi* spp., confirming that the life stages described belong to *S. cruzi*.

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**Competing interests.**

The author declares there are no conflicts of interests.
Ethical approval.
Not applicable

Author’s contributions.
J. P. Dubey conceived and wrote the initial draft. Aditya Gupta, Larissa S. de Araujo, and Asis Khan performed molecular investigations, Oliver Kwok helped with bibliography, and B. M. Rosenthal edited and provided encouragement to complete this study. All authors approved the final manuscript.
References


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**Gjerde B** (2016). Molecular characterisation of *Sarcocystis bovifelis*, *Sarcocystis bovini* n. sp.,
Sarcocystis hirsuta and Sarcocystis cruzi from cattle (Bos taurus) and Sarcocystis sinensis from water buffaloes (Bubalus bubalis). Parasitology Research. 115, 1473-1492. doi: 10.1007/s00436-015-4881-5. PMID: 26677095.


Streitel RH and Dubey JP (1976). Prevalence of *Sarcocystis* infection and other intestinal


Figure 1. Development of first generation schizonts of *Sarcocystis cruzi* in arteries of calves. (A) Longitudinal section of an artery in mesenteric lymph node of a calf 36, day 11 p.i. Note an early schizont (arrow). The infected endothelial cell is protruding in the lumen. Note the central nucleus (arrowhead) of the schizont. HE. (B) Cross section of a renal artery showing three schizonts (a, b, c) in sequential stages of development. The schizonts have occluded the
lumen of the artery. Plastic-embedded 3 µm section. HE. (a) Early immature schizont with lobulation of nucleus with nucleoli (arrowheads). (b) Maturing elongated schizont with merozoite formation, each nuclear lobe is incorporated in a merozoite. (c) An almost mature schizont with radially arranged merozoites.
Figure 2. Development of second-generation *Sarcocystis cruzi* schizonts in kidney of calf 383, day 24 p.i. Plastic-embedded 3 µm section. HE. (A) Early immature narrow multilobed
schizont (arrow) in interstitium; note prominent nucleoli (arrowhead). (B) Immature schizont (arrow) in glomerulus with nuclear lobes separated by clefts. (C) Schizont (arrow) with randomly arranged nuclear lobes. (D) Two schizonts in glomerulus. (a) Immature schizont (arrow) in glomerulus with nucleoli (arrowhead) in nuclear lobes. (b) A mature schizont with radially arranged merozoites (arrow).
**Figure 3.** Asexual stages of *Sarcocystis cruzi.* (A) Merozoite in buffy coat smear of peripheral blood of calf 412, day 28 p.i. For size comparison, erythrocytes and thrombocytes are presented. Giemsa stain. (B) Early sarcocyst (arrow) in parasitophorous vacuole (PV) of myocardium of calf 601, day 42 p.i. HE. (C) An immature sarcocyst (arrow) with six metrocytes (arrowheads) in PV of diaphragm of calf 460, day 55 p.i. HE. (D) Cross section of an immature sarcocyst in myocardium of calf 460, day 55 p.i. Plastic-embedded 3 µm section. HE. Note sarcocyst wall (arrows). The metrocytes (arrowheads) are globular. (E, F) Sarcocysts in tongue of calf 488, day 153 p.i. HE. Sarcocysts have thin walls (arrow). The eosinophilic area surrounding the sarcocysts is degenerated host tissue giving false impression that the cysts are thick walled. (E) Cross section of two sarcocysts (a, b), apparently within one myocyte. The sarcocyst on the left is immature and contained metrocytes are separated by septa (arrowheads). (F) Longitudinal section of a sarcocyst at its one end. Note intensely stained bradyzoites (br) and lightly stained metrocytes (me), and thin septa (arrowheads).
**Figure 4.** Development of sexual stages of *Sarcocystis cruzi* in sections of small intestine of coyotes. The epithelial brush border is oriented up towards the intestinal lumen in B-G. (A) Bradyzoite (arrow) in intestinal lumen. The apical end is stained intensely (arrowhead) and the nucleus (arrow) is central. Plastic-embedded 3 µm section. Coyote 34, 6 h p.i. HE. (B) Three organisms (a, b, c) in a goblet cell. The zoite in (a) is probably a bradyzoite and (b, c) are
macrogamonts. Arrowhead points to host cell nucleus. Coyote 34, 6 h. HE. (C). Two organisms in a goblet cell. The organism marked by arrow is probably a bradyzoite and the one marked by arrowhead is a macrogamont with dividing nucleus. Coyote 44, 12 h p.i. (D) Two macrogamonts within a goblet cell (gc) intensely stained red with PASH. Coyote 34, 6 h p.i. (E) A macrogamont with central nucleus. Arrowhead points to host cell nucleus. Coyote 34, 6 h p.i. (F) A binucleate microgamont (arrow) in a goblet cell (gc). Arrowhead points to host cell nucleus. Coyote 34, 6 h p.i. (G) A microgamont with gametes at the periphery. Arrowhead points to host cell nucleus. Coyote 34, 6 h p.i. (H) Mature microgamont (arrow) in the lamina propria. Coyote 44, 12 h p.i. (I) Two microgametes (opposing arrowheads) apparently free in a cell. Also note 3 macrogamonts (white arrows). Coyote 44, 12 h p.i.
Figure 5. Sporogony of *Sarcocystis cruzi* in small intestine of a coyote 43, day 16 after ingesting infected beef. Plastic-embedded 3 μm section. HE. (A) Unsporulated oocyst with a central nucleus (nu). The outer layer of oocyst wall is partly formed (opposing arrowhead). The inner layer is thin (arrow). (B) Unsporulated oocyst. The outer layer of oocyst wall is partly formed (arrowhead). The inner layer is thin (opposing arrows). Note a central band of basophilic nucleus (arrow). (C) Sporocysts with polar nuclei (arrowheads). (D) Sporulated oocysts/sporocysts (arrowheads) in the lamina propria, beneath the epithelium (double
arrowheads). Note sporozoites (arrow).

Figure 6. Phylogenetic relationships of *Sarcocystis cruzi* with selected members of the Sarcocystidae inferred from various genetic markers (A = 18S rRNA, B = 28S rRNA, C = COX1, D = Acetyl CoA genes ACS) under Neighbor-Joining criterion (1000 bootstrap values; p-distance model; pairwise deletion). Branch supports are indicated near the corresponding nodes. Asterias is indicated against the species under study and the highlighted part shows the *S. cruzi* cluster with highest nodal support.
Table 1. Definitive hosts of *Sarcocystis cruzi* and development of gametogony

<table>
<thead>
<tr>
<th>Definitive host</th>
<th>Country</th>
<th>Inoculum</th>
<th>Prepatent Period (days)</th>
<th>Patent period</th>
<th>Gametogony</th>
<th>Infectivity from sporocysts to calves</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dog (<em>Canis familiaris</em>)</td>
<td>Germany</td>
<td>Naturally infected beef</td>
<td>9 or 10</td>
<td>70 days</td>
<td>Not studied, oocysts in lamina propria</td>
<td>Yes</td>
<td>Heydorn and Rommel (1972)</td>
</tr>
<tr>
<td>Dog (<em>Canis familiaris</em>)</td>
<td>USA</td>
<td>Naturally infected beef</td>
<td>9 - 22</td>
<td>3 - 16 days</td>
<td>Sporogony in lamina propria</td>
<td>Yes</td>
<td>Fayer (1974)</td>
</tr>
<tr>
<td>Dog (<em>Canis familiaris</em>)</td>
<td>USA</td>
<td>Experimentally infected beef</td>
<td>9 days</td>
<td>Not studied</td>
<td>Gametogony and sporogony in lamina propria</td>
<td>Yes</td>
<td>Sheffield and Fayer (1980)</td>
</tr>
<tr>
<td>Coyote (<em>Canis latrans</em>)</td>
<td>USA</td>
<td>Feces of naturally infected coyotes fed to calves</td>
<td>9 days</td>
<td>Not studied</td>
<td>Gametogony and sporogony in lamina propria</td>
<td>Yes</td>
<td>Fayer and Johnson (1975)</td>
</tr>
<tr>
<td>Wolf (<em>Canis lupus</em>)</td>
<td>Germany</td>
<td>Naturally infected beef</td>
<td>9 or 10</td>
<td>&gt; 6 weeks</td>
<td>Not studied</td>
<td>Not tested</td>
<td>Rommel <em>et al.</em> (1974)</td>
</tr>
<tr>
<td>Species</td>
<td>Location</td>
<td>Source of Infection</td>
<td>Number</td>
<td>Duration</td>
<td>Study Information</td>
<td>Cyst Results</td>
<td>Reference(s)</td>
</tr>
<tr>
<td>-------------------------------</td>
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<td>--------</td>
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<td>-------------------</td>
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<td>----------------------------------------</td>
</tr>
<tr>
<td>Red fox (Vulpes vulpes)</td>
<td>Germany</td>
<td>Naturally infected beef</td>
<td>9 or 10</td>
<td>&gt; 6 weeks</td>
<td>Not studied</td>
<td>Not tested</td>
<td>Rommel et al. (1974)</td>
</tr>
<tr>
<td>Red fox (Vulpes vulpes)-subspecies V. fulva</td>
<td>USA</td>
<td>Naturally infected beef</td>
<td>11</td>
<td>Not studied</td>
<td>Not tested</td>
<td></td>
<td>Fayer et al. (1976b)</td>
</tr>
<tr>
<td>Red fox (Vulpes vulpes)</td>
<td>USA</td>
<td>Experimentally infected beef</td>
<td>9</td>
<td>Not studied</td>
<td>Yes</td>
<td></td>
<td>Dubey (1983)</td>
</tr>
<tr>
<td>Raccoon (Procyon lotor)</td>
<td>USA</td>
<td>Naturally infected beef</td>
<td>8</td>
<td>Not studied</td>
<td>Not studied</td>
<td>Not tested</td>
<td>Fayer et al. (1976b)</td>
</tr>
<tr>
<td>Raccoon (Procyon lotor)</td>
<td>USA</td>
<td>Experimentally infected beef</td>
<td>No sporocysts</td>
<td></td>
<td>Not tested</td>
<td></td>
<td></td>
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<tr>
<td>Crab-eating fox (Cerdocyon thous)</td>
<td>Brazil</td>
<td>Naturally infected feces infective to a calf, muscles from calf infective to a dog</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Raccoon dog (Nyctereutes)</td>
<td>Japan</td>
<td>Naturally infected beef</td>
<td>9</td>
<td>&gt; 6 weeks</td>
<td>Not studied</td>
<td></td>
<td>Saito and Itagaki (1994)</td>
</tr>
</tbody>
</table>
Two calves fed sporocysts (135,000 or 720,000) from feces of naturally infected coyotes from Utah developed acute sarcocystosis.

Raccoons were adults when trapped.

Table 2. Details of *Sarcocystis cruzi* (Bozeman isolates) specimens deposited in the Smithsonian Museum.

<table>
<thead>
<tr>
<th>Specimen No.</th>
<th>Host</th>
<th>Histology Case No.</th>
<th>Days (d) or hours (h) p.i.</th>
<th>Tissue</th>
<th>Figure No.</th>
<th>Stain</th>
<th>Museum No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Calf 36</td>
<td>19252</td>
<td>11 d</td>
<td>Mesenteric lymph node</td>
<td>1A</td>
<td>HE</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Calf 38</td>
<td>19253</td>
<td>15 d</td>
<td>Kidney</td>
<td>1B</td>
<td>HE</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Calf 383</td>
<td>451-80</td>
<td>24 d</td>
<td>Kidney</td>
<td>2 A, B, C</td>
<td>IH</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Calf 412</td>
<td>None</td>
<td>28 d</td>
<td>Blood</td>
<td>3A</td>
<td>Giemsa</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Calf 601</td>
<td>469-81</td>
<td>42 d</td>
<td>Heart</td>
<td>3B</td>
<td>IH</td>
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<td>6</td>
<td>Calf 460</td>
<td>171-81</td>
<td>55 d</td>
<td>Diaphragm</td>
<td>3C</td>
<td>HE</td>
<td></td>
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<tr>
<td>7</td>
<td>Calf 460</td>
<td>171-81</td>
<td>55 d</td>
<td>Heart</td>
<td>3D</td>
<td>HE</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>Calf 488</td>
<td>414-81</td>
<td>153 d</td>
<td>Tongue</td>
<td>3E, F</td>
<td>HE</td>
<td></td>
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<tr>
<td>9A</td>
<td>Coyote 34</td>
<td>C-177-81</td>
<td>6 h</td>
<td>Small intestine</td>
<td>4A, B, E, F, G</td>
<td>HE</td>
<td></td>
</tr>
<tr>
<td>9B</td>
<td>Coyote 34</td>
<td>C-177-81</td>
<td>6 h</td>
<td>Small intestine</td>
<td>4D</td>
<td>PASH</td>
<td></td>
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<tr>
<td>10</td>
<td>Coyote 44</td>
<td>C-178-81</td>
<td>12 h</td>
<td>Small intestine</td>
<td>4 C, H, I</td>
<td>HE</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>Coyote 43</td>
<td>C-190-81</td>
<td>16 d</td>
<td>Small intestine</td>
<td>5 A-D</td>
<td>HE</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>Calf 383</td>
<td>451-80</td>
<td>24 d</td>
<td>Kidney</td>
<td>Paraffin block</td>
<td>None</td>
<td></td>
</tr>
</tbody>
</table>