**Sarcocystis capracanis** infection in Philippine Domestic Goats (*Capra hircus*): Ultrastructural Studies

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Published reports of *Sarcocystis* species in the goat *Capra hircus* are relatively few (Foreyt 1989; Singh et al. 1990; Saito et al. 1996; Woldemeskel & Gebreab 1996). Three species have been reported in domestic goats worldwide, namely: *Sarcocystis capracanis* Fischer, 1979, *Sarcocystis moulei* Neveu-Lemaire, 1912 (Syn: *Sarcocystis caprafelis* El-Refaii, Abdel-Baki and Sevin, 1980), and *Sarcocystis hircicanis* Heydorn and Unterholzner, 1983 (Dubey 1980; Dubey et al. 1984; Dubey et al. 1989). In the Philippines only the bubaline *Sarcocystis fusiformis* (Railliet, 1897) Bernard and Bauche, 1912 (Tongson & Molina, 1979; Parairo et al. 1988; Claveria & Cruz 1999), the redescribed *Sarcocystis levinei* (Dissanaike and Kan, 1978) Huang, Dubey and Ugglia, 1997 (Huong et al. 1997; Claveria & Cruz, 2000), the cattle *Sarcocystis cruzi* (Hasselman, 1926) Wenyon, 1926 (Claveria et al. 2001a) and the swine *Sarcocystis miescheriana* (Kuhn 1865) Labbe, 1899 (syn: *Sarcocystis suicanis* Erber, 1977) (Claveria et al. 2001b) have been identified to date. This paper presents the first documented case of *Sarcocystis capracanis* infection in Philippine domestic goats.

**Methods**

Muscle tissue samples were obtained from twenty six 3-to5-yr-old domestic goats grown in the provinces of Batangas, south of Manila and Iloilo, Visayas and slaughtered in Metro Manila. The examination for sarcocysts was limited to muscles of the esophagus, neck, diaphragm and fore- and hind limbs.
Tissue samples in labeled-plastic bags inside a cooler were brought to the Zoology and Parasitology Research Laboratory at the Science and Technology Research Center, De La Salle University-Manila, and were examined on the day of collection. Tissue samples were initially examined for the presence of macrosarcocysts, and thin muscle slices were then teased lightly and examined for microsarcocysts. Those positive for sarcocysts were fixed in 10% formalin and processed following the standard histologic and H & E staining procedure, and some were fixed in 2.5% cold glutaraldehyde for transmission electron microscopy (TEM). Semithin sections stained with toluidine blue were prepared to aid in the localization of sarcocysts in tissue samples prior to the preparation of ultrathin tissue sections for TEM (Joel-1010, Japan) at the Biomedical Research Center, St. Luke’s Medical Center, Quezon City, Manila. The sarcocyst ultrastructure particularly the cyst wall was characterized using the cyst wall sarcocysts types established by Dubey et al. (1989).

Results and Discussion

Eleven goats had microsarcocysts. Muscle tissue samples had few to several fusiform and/or rounded sarcocysts, with the predominance of the fusiform form in the limb muscle. Fusiform sarcocysts measured 100-200µm long and 7-31µm across, and exhibited compartmentalized zoites (Fig. 1), while rounded

Figures. 1-4. Sarcocystis capracanis in limb muscle tissue. 1. Fusiform-shaped sarcocyst showing zoite compartmentalization. 2. Rounded non-septate sarcocyst. 3. Semithin tissue section showing mature sarcocyst (S) and intravascular schizonts (Box). 4. Histologic tissue section with intravascular schizonts showing prominent zoites (arrows). Muscle fibers (M).
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Figures 5-9. *Sarcocystis capracanis*. 5. Cyst wall with palisade-like villar protrusions and compartmentalized arrangement of zoites (Z) and metrocytes (Me). 6. Portion of the cyst wall showing spacing of villar protrusions (VP) and serrated VP wall. Note marked damage to surrounding host tissues (HT). Sections of zoites (Z) and metrocyte (Me). 8. Cyst wall with VP and numerous lightly-stained amylopectin granules in zoites (Box). 9. Cyst wall showing VP and primary cyst wall with undulations/tubular extensions (arrows) and granular layer (GL).

Sarcocysts about 15-30µm across exhibited no apparent zoite compartments (Fig. 2). In both histologic and semithin tissue sections intravascular schizonts and parasites enroute to parasitization of muscle fibers were evident (Figs. 3 & 4).

The sarcocysts possess distinct striated cyst wall (Fig. 5), 2.2-3.17µm thick with tightly packed palisade-like villar protrusions (VP) (Figs. 6 & 8). Inside the sarcocysts are groups of densely-stained zoites and lightly-stained metrocytes, showing distinct septal partitions (Figs. 5 & 7). Most VP are spaced 1.1-1.4µm apart and possess microfilaments and electron-dense wavy wall (Figs. 6 & 8). The basal and distal segments of the VP measured 0.41-0.85µm and 0.15-0.29µm, respectively. The 0.15-0.18µm thick primary cyst wall (PCW) bears prominent and evenly distributed undulations or tubular extensions interspersed between
the VP (Fig. 9). The granular layer (=ground substance) 0.66-0.77\(\mu\)m thick is confluent with the villar core and septae enclosing the zoites (Figs. 8 & 9). The sarcocyst of \textit{S. capracanis} typifies the Type 14 cyst wall (Dubey et al. 1989).

The presence of sarcocysts in esophageal and diaphragm muscle corroborates earlier reports in goats (Singh et al. 1990; Woldenmeskel & Gebreab 1996). Immature and fully developed \textit{S. capracanis} has been demonstrated in the spinal cord and liver of goats (Collins et al. 1980; Dubey et al. 1984). Considering the limitation in the tissue samples examined, we do not preclude the possibility of infection of other internal organs as earlier reported. The fusiform sarcocysts isolated in the present study were smaller (100-200\(\mu\)m by 7-13\(\mu\)m) compared to those sarcocysts (35-1000\(\mu\)m by 20-100\(\mu\)m) reported by Dubey et al. (1984). Moreover, the rounded sarcocysts we noted bearing unclear septal partitions may represent another etiologic species of goat sarcocystosis or possibly a developmental form of \textit{S. capracanis}.

The striated palisade shape cyst wall of \textit{S. capracanis} we observed bears similarities with earlier studies (Fischer, 1979; Aryeetey et al. 1980; Dubey et al. 1984; Saito et al. 1996). The VP length and width (1.1-3\(\mu\)m by 0.15-0.85\(\mu\)m) approximate those of the sarcocysts (1.8-2.8 \(\mu\)m by 0.41-0.61\(\mu\)m) isolated at 64 days post-inoculation (DPI) by Dubey et al. (1984). Thickness of the granular layer of sarcocysts isolated approximates that of the sarcocysts obtained at 64 DPI (Dubey et al. 1984), though thicker compared to the sarcocysts isolated from goats at 56, 92 and 118 DPI (Aryeetey et al. 1980). The variations may have been influenced by differences in the source of sarcocysts, i.e from naturally-infected goats in the present study as against experimentally-exposed goats in earlier reports. In the presence of morphological variations in the wall of \textit{Sarcocystis} belonging to the same species or different strains, Dubey et al. (1989) attribute this to possible influence of age and developmental stage of the parasite, host cell parasitized and difference in muscle tissue processing employed.

Ultrastructurally, \textit{S. capracanis} exhibits resemblance with other species identified as having Types 14, 15, 16 or 17 sarcocyst wall, such as: \textit{Sarcocystis gracilis} Ratz, 1908 of the roe deer (\textit{Capreolus capreolus}); \textit{Sarcocystis odoi} Dubey and Lozier, 1983, and \textit{Sarcocystis odocoileocanis} Crum. Fayer and Prestwood, 1981 of the white-tailed deer (\textit{Odocoileus virginianus}); \textit{Sarcocystis tarandivulpes} Gjerde, 1984, \textit{Sarcocystis rangiferi} Gjerde, 1984, and \textit{Sarcocystis tarandi Gjerde}, 1984 of the reindeer (\textit{Rangifer tarandus tarandus}) (Dubey et al. 1989). While our study represents the first documented case of sarcocystosis capracanis in the country, it would be valuable to continue to identify other probable etiologic species infecting goats, and in the near future to carry out experimental exposure studies to establish the parasite’s definitive host(s).

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