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**LIFE CYCLE OF CALYPTOSPORA FUNDULI (APICOMPLEXA: CALYPTOSPORIDAE)**

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**ABSTRACT:** The taxonomic status of the extraintestinal piscine coccidium *Calyptospora funduli* is based in part on its requirement of an intermediate host (the daggerblade grass shrimp *Palaemonetes pugio*). In the present study, grass shrimp fed livers of Gulf killifish (*Fundulus grandis*) infected with sporulated oocysts of *C. funduli* exhibited numerous sporozoites suspended in the intestinal contents when fresh squash preparations were examined by light microscopy. Using this method, sporozoites were not seen in intestinal epithelial cells of the grass shrimp or in any other cell type. Ultrastructural examination, however, revealed sporozoites in the cytoplasm of the gut basal cells. Cross-sections of 1–13 sporozoites were seen within a single cell, and those sporozoites each appeared to be situated in individual membrane-bound vesicles, rather than in a single parasitophorous vacuole. These ultrastructural observations indicate that in the grass shrimp intermediate host, sporozoites that develop into an infective stage probably undergo that development in gut mucosal-bound vesicles, rather than in a single parasitophorous vacuole.

*Solanogi and Overstreet (1980), and postulated for some species of Goussia by Overstreet (1982). However, the first heteroxenous life cycle to be fully demonstrated experimentally for fish coccidia was that of *C. funduli* in killifish (Fournie and Overstreet, 1993). The role of an invertebrate intermediate host in the transmission of fish coccidia was first suggested by Landau et al. (1975), demonstrated for *C. funduli* by Solangi and Overstreet (1980), and postulated for some species of *Goussia* by Overstreet (1981) and Paterson and Desser (1982). However, the first heteroxenous life cycle to be fully demonstrated experimentally for fish coccidia was that of *C. funduli* in killifish (Fournie and Overstreet, 1983). They showed that excysted sporozoites in the intestine of palaemonid shrimps changed morphologically before they became infective to fishes; however, at the light microscopic level, sporozoites were not seen in host cells. Subsequently, Steinhagen and Körting (1990) confirmed that tubificid oligochaetes could serve at least as a paratenic host or vector for *Goussia carpelli*, and that the sporozoites were located in the cytoplasm of the intestinal epithelial cells of the oligochaete. However, Steinhagen and Körtíng (1988) previously showed that the oligochaete was not necessary for transmission and that direct infections could occur by fecal contamination from fish to fish.

We report here details regarding the life cycle of *C. funduli* in its intermediate host and clarify the taxonomic status of this coccidium. Specific information is provided regarding intracellular residence of sporozoites in the alimentary tract of the grass shrimp intermediate host and the route by which these infective sporozoites reach the liver of the killifish definitive host.

**MATERIALS AND METHODS**

Materials for this study consisted of livers infected with sporulated oocysts of *Calyptospora funduli* from the Gulf killifish (*F. grandis*) caught in minnow traps from Halstead Bayou, Ocean Springs, Mississippi. Daggerblade grass shrimp (*Palaemonetes pugio*) to be used for feeding studies were caught by dipnet from the same locality. Adult fish (>80 mm total length) to be used for feeding studies were uninfected longnose killifish (*F. similis*) caught by minnow traps from Horn Island, Mississippi. These fish and shrimp were maintained in separate cages in 75-L glass aquaria with 12–15 ppt salt (Instant Ocean®) at approximately 23°C. Both were fed daily with a varied diet, including frozen brine shrimp, TetraMin®, and Kordon Maintenance Flakes®.

To determine where in the shrimp alimentary tract the sporozoites reside, grass shrimp were experimentally infected by feeding them liver containing large numbers of sporulated oocysts from wild specimens of *F. grandis* once a day for 3 days. Shrimp were not examined for natural infections prior to feeding with infected fish livers. The intestines were then dissected from the shrimp after 7 days and either examined by light microscopy as fresh squash preparations or processed for electron microscopy.

Grass shrimp intestines for electron microscopy were fixed for 2 hr in 3% (v/v) glutaraldehyde in 0.1 M sodium cacodylate buffer (pH 7.4), rinsed in 0.1 M sodium cacodylate buffer (pH 7.4), postfixed in cacodylate-buffered 1% (wt/v) osmium tetroxide for 2 hr, rinsed briefly in buffer, and dehydrated in a graded series of ethanol. Tissues were embedded in Spurr’s low viscosity resin. Semithin sections (1 µm) were cut using a Reichert Ultracut E microtome and stained with toluidine blue. Ultrathin sections (50–60 nm) were cut with a diamond knife, mounted on uncoated copper grids, stained with uranyl acetate and lead citrate (Hayat, 1981), and examined with a Zeiss EM 902A transmission electron microscope.

To determine the route by which sporozoites of *C. funduli* reached hepatocytes, 20 uninfected wild specimens of *F. similis* were fed experimentally infected grass shrimp, and 2 individuals each were examined at 4, 8, 12, 18, 24, 36, 48, 72, and 96 hr postinfection (PI). Before these fish were used as test subjects, livers from 20 were examined by squash preparations and found to be free of infection. Examination of fish administered infected shrimp involved preparations of peripheral blood smears, tissue impressions of liver and spleen, and histologic sections of liver and intestine. Blood smears were prepared using blood collected from the dorsal aorta after removal of the caudal buffer, and dehydrated in a graded series of ethanol. Tissues were embedded in Spurr’s low viscosity resin. Semithin sections (1 µm) were cut using a Reichert Ultracut E microtome and stained with toluidine blue. Ultrathin sections (50–60 nm) were cut with a diamond knife, mounted on uncoated copper grids, stained with uranyl acetate and lead citrate (Hayat, 1981), and examined with a Zeiss EM 902A transmission electron microscope.

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FIGURES 1–4. Infective sporozoites of *Calyptospora funduli* in the daggerblade grass shrimp, *Palaemonetes pugio*, 7 days after being fed liver containing sporulated oocysts. (1) Infective sporozoite from intestinal contents. Note the two prominent refractile bodies (arrowheads). Bar = 15 μm. (2) Low-magnification electron micrograph (EM) showing cross-sections of two sporozoites in basal cell (arrow) of intestinal mucosa. Note brush border (b), columnar epithelial cells (e), and basement membrane (m). Bar = 10 μm. (3) Higher magnification of infected basal cell.
Tissue impressions were prepared by touching the cut surface of the liver and spleen to glass microscope slides and allowing them to air dry. Blood smears and tissue impressions were stained with American Scientific Products Camco Quik Stain (available from Sigma, St. Louis, Missouri) or Fisher Scientific Giemsa stain (Pittsburgh, Pennsylvania). Smears stained with Camco Quik Stain were dipped in stain for 10 sec then in distilled water for 20 sec. Smears stained with Fisher Scientific Giemsa stain were fixed in methanol for 5–10 min, stained in a working solution of the stain for 25 min, then dipped in distilled water. The 2 remaining fish were examined 20 days PI to verify infection with C. funduli.

RESULTS

Light microscopic examination of squash preparations of intestine from experimentally infected grass shrimp 7 days PI showed large numbers of nonmotile sporozoites. The sporozoite had a distinct nucleus near the center of the body, with 2 relatively large refractile bodies, 1 being situated anterior to and the other posterior to the nucleus (Fig. 1). Sporozoites were suspended in intestinal contents and occurred alone or appeared to be situated in nests between intestinal cells. Sporozoites could not be seen in intestinal epithelial cells of the grass shrimp or in any other cell types by this method.

Semithin cross-sections of grass shrimp intestine revealed sporozoite-like stages in the cytoplasm of the basal cells, which are located proximal to the nuclei of the simple columnar epithelial cells (Fig. 2). Cross-sections of many infected cells exhibited 1–5 sporozoite-like stages within a single cell, but a few had up to 13 per cell (Fig. 3). Ultrastructurally, the intracellular stages were identified as sporozoites, and they contained the characteristic cytoplasmic organelles. These structures included refractile bodies, a pellicle, micronemes, rhoptries, amyllopectin granules, mitochondria, Golgi complexes, ribosomes, and a nucleus (Fig. 4). Additionally, individual sporozoites appeared to be situated in single membrane-bound vesicles rather than within a single parasitophorous vacuole (Figs. 3, 4). No evidence of replication was evident.

Sporozoites were found in the peripheral blood from 1 of 2 specimens of F. similis examined at 4 hr PI, and all sporozoites contained 2 distinctive refractile bodies (Figs. 5, 6). No sporozoite was observed in the peripheral blood of the other 16 killifish examined at the later time intervals. Additionally, eosinophilic inclusions resembling coccidian trophozoites and meronts were seen in hepatocytes from several killfish at 48, 72, and 96 hr PI. Sporozoites or sporozoitelike stages were not found in the intestinal lumen, mucosa, or submucosa nor in tissue impressions of liver and spleen of F. similis examined anywhere from 4 to 96 hr after ingesting experimentally infected grass shrimp. The 2 positive control fish had infections with developing oocysts at 20 days PI.

DISCUSSION

Using transmission electron microscopy, Fournie and Overstreet (1983) noted that in 1 instance, a few sporozoites were seen within a degenerated cell of a P. pugio intestine. This was the only evidence provided that indicated that sporozoites might infect shrimp intracellularly as previously suggested (Landau et al., 1975; Dyková and Lom, 1981; Upton and Duszynski, 1982). Whereas Fournie and Overstreet (1983) saw no indication of division of the parasite in the grass shrimp in fresh squash preparations, the present study clearly demonstrates that sporozoites can enter host cells. However, based on the lack of reproducing stages, intracellular asexual multiplication appears unlikely in the grass shrimp intermediate host. The fact that infected cells exhibited from 1 to 13 sporozoites within a basal cell and each of those occurred within its own parasitophorous vacuole provides evidence for multiple penetrations rather than for a form of asexual multiplication. Because the grass shrimp were not laboratory reared, some of the many sporozoites seen showing nine sporozoites (s), each within a distinct membrane-bound parasitophorous vacuole (v). Bar = 2 μm. (4) Higher magnification EM showing characteristic features of sporozoites. Note refractile body (Rb), micronemes (mn), amyllopectin granules (a), and membrane (arrows) surrounding individual sporozoites. Bar = 1 μm.
by electron microscopy could have been present from a natural infection. Sporozoites can apparently survive in the grass shrimp at least 201 days (Fournie and Overstreet, 1983).

The ultrastructure of the sporozoite of *C. funduli* within basal cells of the grass shrimp intestine was consistent with fine structural features of other piscine coccidian sporozoites (Paterson and Desser, 1984; Morrison and Poynton, 1989; Steinhagen, 1991) and eimerian-type sporozoites in general (Chobotar and Scholtyscek, 1982). All have the distinctive refractile bodies, micronemes, rhoptries, and amylopectin granules. Sporozoites of *G. carpelli* invaded intestinal epithelial cells, and sporozoites of *G. subepithelialis* were found in epithelial and phagocytic cells of the alimentary canal of tubificid oligochaetes (Steinhagen, 1991). In both cases, the sporozoites occurred singly in membrane-bound parasitophorous vacuoles. The infection of *C. funduli* is different because single or multiple sporozoites, which undergo development, usually occurred within individual membrane-bound vesicles in basal cells of the grass shrimp intestine.

Once an infected grass shrimp is ingested by a fish host, infective sporozoites of *C. funduli* apparently gain access to the fish’s hepatocytes through its circulatory system, as evidenced by the occurrence of sporozoites in the peripheral blood of killifish 4 hr after ingesting infected grass shrimp. A few authors (Odense and Logan, 1976; Paterson and Desser, 1982) have suggested that sporozoites of some extraintestinal forms reach their target organs through the circulatory system. Odense and Logan (1976) reported sporocysts of *G. gadi* within and around a major hepatic vein of the haddock but no sporozoites in peripheral blood smears. Paterson and Desser (1982) reported finding “tiny apicomplexan-like meronts” in the vascular endothelium of the liver and swim bladder of common shiners infected with *G. deguistii*. Even though sporozoites were not seen in the intestinal mucosa or submucosa of exposed *F. similis*, they presumably penetrate the intestinal mucosal epithelium, enter small blood vessels in the lamina propria, and subsequently pass through the hepatic portal system, either directly or possibly within a migratory host cell. The sporozoites observed in a peripheral blood smear obtained from the severed caudal peduncle of 1 fish at 4 hr PI indicated that they reached the liver through the circulatory system. This route would account for the short time necessary for the parasite to reach the liver, and it would also explain the reported but rarely observed extrahepatic sites (Solangi and Overstreet, 1980). Hawkins et al. (1984) have demonstrated that the parasite reaches the liver quickly because the first of 2 generations of merozoites are well developed by day 4 PI in *F. similis*.

The eosinophilic inclusions seen in hepatocytes of killifish at 48 to 96 hr PI resembled eimerian trophozoites and meronts. This finding supports the hypothesis that the sporozoites reach the liver of killifish in a short period of time, which in turn provides further support for the sporozoites using the circulatory system to reach the liver. These inclusions morphologically resemble the eosinophilic bodies illustrated by Upton and Duszynski (1982) from experimentally infected *F. heteroclitus* at 5 and 6 days PI. Sporozoites of *C. funduli* may require more time to become established in the liver of *F. heteroclitus* than in the liver of *F. similis*.

Even though Molnár (1995) considered the grass shrimp to be a paratenic host for *C. funduli*, in which the sporozoites do not undergo development, Solangi and Overstreet (1980), Fournie and Overstreet (1983), and the present study clearly show that the grass shrimp serves as a necessary intermediate host in the life cycle of *C. funduli*, rather than a paratenic host in which the parasite undergoes no further development. *Goussia carpelli* and probably *G. subepithelialis*, in studies by Steinhagen (1991) and Steinhagen and Korting (1988, 1990) dealing with those intestinal coccidians and 2 oligochaete paratenic hosts, differ from ours because no development was noted and the oligochaete worm was not necessary to complete the life cycle. Whether sporozoites of *C. funduli* resulted from asexual multiplication or represent maturing individuals is not known. Nevertheless, without a developmental period of about 5 days in the grass shrimp, the sporozoite is not infective to killifishes (Fournie and Overstreet, 1983). Consequently, the crustacean invertebrate host is necessary to complete the life cycle.

Lom and Dyková (1992) stated that the definition of the Eimeriidae by Levine (1985) adequately covers *Calyptospora*. Levine’s definition included those coccidians that were homoxenous or at least without asexual multiplication in the non-definitive host. Those authors indicated that they preferred to leave *Calyptospora* in the Eimeriidae unless asexual multiplication can be demonstrated in the intermediate host. However, because *C. funduli* uses the crustacean as more than a paratenic host and has distinguishing morphological structures, features that clearly distinguish related coccidians from the “typical” eimeriids, we believe that *C. funduli* belongs in the Calypsoспорidiae as described by Overstreet et al. (1984), rather than in Eimeriidae. Even though life cycles of the other species in the genus *Calyptospora* have not been demonstrated experimentally, it is likely that they are similar. Subsequently, Levine (1988) did accept the *Calyptosporidae* in his revision of the Apicomplexa.

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**LITERATURE CITED**


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