Description of *Dientamoeba fragilis* Cyst and Precystic Forms from Human Samples

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*Dientamoeba fragilis* is a common enteropathogen of humans. Recently a cyst stage of the parasite was described in an animal model; however, no cyst stage has been described in detail from clinical samples. We describe both cyst and precystic forms from human clinical samples.

*Dientamoeba fragilis* is an amoeboid trichomonad parasite described in all continents with human habitation (1). The prevalence of *D. fragilis* varies widely; however, with the advent of molecular diagnostics, rates of infection are reportedly higher than those of *Giardia intestinalis* (2). These statistics are consistent with other studies using nonmolecular methods (3–6).

The pathogenic potential of *Dientamoeba* has been suggested by numerous researchers (7–9). Confirmation of virulence and the potential mechanisms of pathogenicity are yet to be determined (10). However, infection of mice with *D. fragilis* resulted in statistically significant weight loss, gastrointestinal disturbance, and unformed stools compared to the results for controls (11). Levels of calprotectin, a marker of inflammatory disease in the lower gastrointestinal tract (12), in infected animals were found to be twice those in control mice, which is indicative of intestinal inflammation (11).

The development of this animal model also led to the discovery and first description of the cyst form of *D. fragilis* (11). The discovery of a protozoan cyst in *D. fragilis*-infected mice was made initially by light microscopy and was confirmed by transmission electron microscopy. The mice were confirmed to be protozoan free by screening for several days by PCR (for *D. fragilis* DNA) prior to infection and by examination of iron hematoxylin-stained fecal smears for protozoa. The cysts possessed a distinct thick cyst wall, with a membranous, irregular inner cyst wall located directly adjacent to the encysted parasite surrounded by a distinctive peritrophic space. Furthermore, cysts of *D. fragilis* possessed a characteristic *D. fragilis* nucleus (11) (Fig. 1). While humans are thought to be the preferred host of *D. fragilis* (13), a number of animal hosts have been reported, including nonhuman primates, rodents, pigs, and a sheep (11, 14–16). While there have been sporadic reports of both precystic stages, pseudocysts, and true cysts of *D. fragilis* in human samples, there is a lack of conclusive evidence for these stages (17). In a recent report, Clark et al. (18) raise the point that the existence of a *D. fragilis* cyst stage would be greatly substantiated if these forms could be identified in human stool specimens also containing *D. fragilis* trophozoites. This study was carried out to determine if cysts (or precystic stages) identical to those reported in mice could be identified in human clinical specimens.

A retrospective study was performed on all permanently stained smears positive for *D. fragilis* performed at the Microbiology Department at St. Vincent’s Hospital Sydney. All positive slides collected over a 2-year period from 5 July 2012 to 12 December 2014 were included in the study. The slides were stained as previously described (19). A total of 500 slides were examined by oil immersion (1,000×) light microscopy, with approximately 500 fields of view examined for each slide. Special care was taken to identify forms bearing a strong resemblance (or considered identical) to the cysts described by Munasinghe et al. (11). Care was also taken to make note of forms bearing a morphology between that of *D. fragilis* trophozoites and cysts as these were likely to represent the precyst or pseudocyst forms previously described (and illustrated) by pioneering investigators (20–22). At the same time in the United States, a total of 47 Wheatley’s trichrome permanently stained smears positive for *D. fragilis* trophozoites were reexamined for forms consistent with the prior description of the cyst stage (11). These smears represented 47 different patients.

A total of 5 true cysts (as per reference 11) were detected from 4 patient samples (Fig. 2), giving a prevalence of 0.01% cyst per patient sample. The cysts were detected independently in 2 different laboratories from different locations (Australia and the United States).

Putative precyst or pseudocyst forms (as per the studies by Wenrich [21], Kofoid [20], and Kudo [22]) were found in a total of 163 of the 500 slides, giving a prevalence of 32.6% per patient sample.

In this article, we describe *D. fragilis* cysts and what appear to be precystic forms of *D. fragilis* from human clinical samples (Fig. 2 and 3). The role these forms play in the transmission of human dientamoebiasis is yet to be confirmed. However, the cyst forms described here are morphologically indistinguishable from the *D. fragilis* cysts recently described in detail by Munasinghe et al. (11) in rodents (Fig. 1). The cysts are comprised of a distinct cyst wall (~5 μm in diameter) with a zone of clearance around the cyst. A peritrophic space is present between the cyst wall and the amoebic parasite enclosed within. The nuclear structure is characteristically identical to what is found in *D. fragilis* trophozoites. All cysts

Received 20 March 2014 Returned for modification 23 April 2014 Accepted 2 May 2014 Published ahead of print 7 May 2014

Editor: P. H. Gilligan

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doi:10.1128/JCM.00813-14
seen were binucleate, with each nucleus containing a large central karyosome with a delicate nuclear membrane. No chromatin is visible on the nuclear membrane, and the nucleus is often fragmented into distinct granules of chromatin, often referred to as “chromatin packets.” According to these findings, these “true” cysts are rarely encountered in clinical samples, which probably accounts for the limited number of reports describing these structures. In contrast, the precystic forms of *D. fragilis* were more frequently encountered, with a prevalence of up to 5% in clinical samples. This precystic stage is characterized by a compact spherical shape with a reduction in size of up to 50% from “normal” trophozoites. These forms range in size from 4 to 5 μm. The cytoplasm is darkly staining, indicating a denser structure than what is found in normal trophozoites. The cytoplasm is homogeneous and rarely contains any inclusions (Fig. 3).

Despite recent reports that suggest that a cyst has never been reported in humans previously (18), this is not the case. The present study is one of several to report a *D. fragilis* cyst in clinical specimens. Charles Atwood Kofoid was the first protozoologist to describe an encysted stage of *D. fragilis* in 1923. He also described

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**FIG 1** *Dientamoeba fragilis* cysts from a mouse animal model (A) and a human clinical sample (B).

**FIG 2** *Dientamoeba* trophozoites (A to C), precysts (D to F), and cysts (G to I) stained with a modified iron hematoxylin stain at ×1,000 magnification.
a precystic form at the same time (20). Kudo some 3 years later while unable to find any cyst-like structures did, however, discover forms approximately 4 μm in diameter that he described as “small spherical amoeba without food particles,” which he concluded were precystic forms of the parasite (22). Cyst forms of *D. fragilis* were also described from Argentina in 1928 (23). Another study conducted in 1936 on the morphological observations of the parasite from 70 infected patients reported the absence of cysts (21). While no true cyst forms were described in this study, the author did describe what he thought was a precystic or pseudocyst stage of *Dientamoeba*. These precystic forms were small (3.5 to 5 μm) and both mononuclear and binucleate, in which the cytoplasm becomes finely and uniformly granular and exhibits intense staining. Wenrich ruled out these being a degenerate form of the parasite as the nuclear structure was “fairly normal,” a condition that is not usually seen in true degenerated trophozoites. It is also noted in a later study that these forms “appear no more common in 24-hour old feces than in fresher material,” once again not supporting the theory of these being a degenerated form (24). Piekarski reported a number of *D. fragilis* cysts from several patients; however, this work has been underrepresented in the peer-reviewed literature to date and should be appropriately considered (25).

In contrast to these findings, some researchers have dismissed the notion of *D. fragilis* having either a true cyst form or a pseudocyst. Wenrich dismissed the report of cysts by Kofoid by attributing the forms as “animals with thin clear ectosarcs” (24). The most vocal opponent of cyst formation in *D. fragilis* was Clifford Dobell, the parasitologist who first described *Dientamoeba* in the scientific literature. He dismissed these reports of both cyst and pseudocyst formation as a misinterpretation of degenerate individuals (16). Dobell also stated that it was inconceivable that *D. fragilis* would have a cyst stage while *Histomonas*—its closest relative—also has no cysts.” However, recent reports describe precystic and cyst forms of *Histomonas meleagridis* (26, 27). These stages are completely spherical compact structures with a size range of 4 to 7 μm in diameter, have been demonstrated to survive adverse conditions, including extreme acid conditions (pH 2) (27), and are morphologically similar to the precystic form of *D. fragilis* described herein. A recent study by our group utilizing cultured *D. fragilis* has demonstrated that *Dientamoeba* is not as fragile as first proposed as it can survive extreme acid conditions of pH 1 to 2 for approximately 4 h (unpublished data). It is probable that the *D. fragilis* precystic/cyst stages facilitate its survival under these extreme conditions. It should also be noted that Dobell described small forms that are identical to the forms that other researchers and our group believe are a precystic form of the parasite (16). In his report (16), Dobell describes “dwarfs, with diameters of only 3–4 μm are also to be found in cultures. They appear to be formed by rapid division, without intermediate growth, of normal individuals. Even the smallest often contain 2 nuclei. Most of these very small organisms also degenerate and die. They appear to be unable to feed, as their cytoplasm is usually free from all food-inclusions when their nuclei are breaking up—unlike giants,
which remain gorged with ingesta long after their nuclei have disintegrated.”

Further definitive studies are required to fully elucidate the role of *D. fragilis* cysts in transmission. *In situ* hybridization studies should be performed to allow visual correlation with the cyst structures identified by light microscopy. Studies utilizing both rodent and pig animal models should also be undertaken, and given the ease of establishing short-term *Dientamoeba* cultures, cyst formation experiments could also be undertaken on trophozoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. Next-generation sequencing of zoite cultures to establish what environmental conditions are needed to induce encystations. 

Moreover, we report the finding of these structures in two different diagnostic laboratories in geographically distant locations. What roles these distinct structures play in transmission is yet to be determined, and further study is needed on the life cycle of this peculiar parasite.

REFERENCES


