Ultrastructure, Immunofluorescence, Western Blot, and PCR Analysis of Eight Isolates of *Encephalitozoon (Septata) intestinalis* Established in Culture from Sputum and Urine Samples and Duodenal Aspirates of Five Patients with AIDS

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Microsporidia are ancient, intracellular, eukaryotic protozoan parasites that form spores and that lack mitochondria. Currently, as many as eight species included under six genera are known to infect humans, mostly patients with AIDS. Among these, *Enterocytozoon bieneusi*, the agent of gastrointestinal (GI) disease, is the most frequently identified microsporidium in clinical laboratories in the United States. *Encephalitozoon (Septata) intestinalis*, the agent that causes a disseminated infection including infection of the GI tract, is the second most frequently identified microsporidian parasite. In spite of this, not many isolates of *E. intestinalis* have been established in culture. We describe here the continuous cultivation of eight isolates of *E. intestinalis* obtained from different samples including the urine, sputum, and duodenal aspirate or biopsy specimens from five AIDS patients originating from California, Colorado, and Georgia. The specific identification was made on the bases of ultrastructural, antigenic, and PCR analyses.

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**MATERIALS AND METHODS**

**Patient specimens.** Urine, sputum, duodenal aspirate or biopsy, and serum specimens were obtained from five patients from California, Colorado, and Georgia, as indicated in Table 1. A fecal sample was also obtained from one patient (Table 1). Smears were made from all of these samples (except the serum) as well as the fecal sample that we had obtained previously from a patient infected with this parasite (30) and were stained with chromotrope by the procedure of Weber et al. (34) and with the quick-hot Gram-chromotrope by the procedure of Moura et al. (20).

The urine, sputum, and duodenal aspirate or biopsy specimens were also processed for culture as described previously (29, 30), with slight modifications. Briefly, urine samples were washed once with Hanks’ balanced salt solution (HBSS) and were inoculated into cell cultures. Sputum samples were treated with spautolysin and were processed as described previously (30). A duodenal aspirate was inoculated directly into cell culture, whereas the biopsy specimen was broken into small pieces by trituration, inoculated into cell cultures, and incubated at 37°C as described previously (31). After 24 h the culture medium from each flask was decanted into a centrifuge tube and mixed with an equal volume of HBSS, and the mixture was centrifuged at 2,000 × g at 4°C. The supernatants were aspirated and the sediment was inoculated back into the respective culture flasks containing fresh growth medium. Thereafter, the me-
TABLE 1. Patient data, specimens obtained, and isolates established in culture

<table>
<thead>
<tr>
<th>Patient no.</th>
<th>Age (yr)</th>
<th>Location</th>
<th>Samples obtained</th>
<th>Isolate name</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (control)</td>
<td>26</td>
<td>California</td>
<td>Urine, feces, saliva, nasal mucosa, serum</td>
<td>CDC:V297</td>
</tr>
<tr>
<td>2</td>
<td>32</td>
<td>Georgia</td>
<td>Duodenal aspirate and biopsy specimen</td>
<td>CDC:V307</td>
</tr>
<tr>
<td>3</td>
<td>34</td>
<td>Colorado</td>
<td>Urine, sputum, serum</td>
<td>CDC:V308&lt;sup&gt;a&lt;/sup&gt; and CDC:V309&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>4</td>
<td>30</td>
<td>California</td>
<td>Urine, sputum, serum</td>
<td>CDC:V314&lt;sup&gt;c&lt;/sup&gt; and CDC:V315&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>5</td>
<td>25</td>
<td>Colorado</td>
<td>Urine, sputum, serum</td>
<td>CDC:V324&lt;sup&gt;e&lt;/sup&gt; and CDC:V325&lt;sup&gt;e&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> All patients were males.
<sup>b</sup> Data for this patient are from a previous report (30).
<sup>c</sup> Isolated from urine.
<sup>d</sup> Isolated from sputum.

Nucleic acid from each sample was resuspended in 50 μl of distilled water and was amplified by PCR.

PCR. PCR was performed by using three different diagnostic primer pairs, as follows: (i) primer pairs SINTF-SINTR, in which the forward diagnostic primer was SINTF (5'-TTTCGATGGTAAAGGAGTTGA-3'), which was designed on the basis of the sequence from positions 362 to 382, and the reverse diagnostic primer was SINTR (5'-CGCTCTCTGGTGTCCTGCGG-3'), which was based on the sequence of the nucleotides from positions 861 to 881 of the E. intestinalis small-subunit (SSU) rRNA sequence that amplifies an E. intestinalis-specific fragment of 520 bp encoding a region for E. intestinalis SSU RNA, as described previously (b); (ii) primer pair EHEL-F–EHEL-R, which amplifies an E. hellem-specific fragment of 546 bp encoding a region for E. hellem SSU RNA (32); and (iii) primer pair ECUN-F–ECUN-R, which amplifies a fragment of 549 bp encoding a region for E. cuniculi SSU RNA (32). PCR was performed with the GenAmp kit (Perkin-Elmer Cetus, Norwalk, Conn.) according to the manufacturer's instructions. A total of 35 PCR cycles were performed with denaturation, annealing, and elongation temperatures of 94°C, 55°C, and 72°C, respectively. The products of the amplification were resolved on a 2% agarose gel (SeaKem GTG; FMC BioProducts, Rockland, Maine) and were visualized for analysis by staining with ethidium bromide (6).

**RESULTS**

Clinical specimens. Chromotrope- and quick-hot Gram-chromotrope-stained smears of fecal, urine, and sputum samples from the patients being studied showed spores with the morphological features characteristic of microsporidia such as a vacuole, belt-like stripe, and gram-positive granules (20). Abundant spores were seen in all samples; the spores measured 1.2 to 2.4 μm. The spores stained pinkish red and dark violet with the chromotrope and the quick-hot Gram-chromotrope stains, respectively (Fig. 1A). When smears made from these samples were reacted with the anti-E. intestinalis serum at a dilution of 1:400, the spores exhibited an apple-green fluorescence (Fig. 1B).

Parasite growth and ultrastructure. The E6 and HLF cell cultures inoculated with the samples from the patients showed no overt bacterial contamination, and foci of infected cells were seen after 4 to 8 weeks. Thereafter, the spores were continually released into the cultures. In the initial stages of culture and growth, the sedimented spores were reincubated into the original culture flasks. By 2 to 4 months of continuous culture, the parasites had adapted well to culture conditions so that amplification of the cultures and regular harvest of spores for use in subsequent assays was possible. In all cultures large numbers of spores were found lying free in the supernatant. Additionally, many cells were found to be distended with spores (Fig. 1C and D). The spores appeared to be birefringent when examined with a microscope equipped with phase-contrast optics.

Scanning electron microscopic images of cells distended with spores appeared as though the spores were covered with a muslin-like cloth (Fig. 1E). By transmission electron microscopy all of the developing stages and spores from all isolates were found to be present within chambers of a septated, hon-
eycomb-like parasitophorous vacuole, a feature typical of *E. intestinalis* (Fig. 2A). Developing stages consisted of meronts, some with two nuclei, which were always found attached to the membranous vacuolar space. Sporogonial stages consisted of mostly disporoblastic and a few tetrasporoblastic stages. Mature spores measured 1.2 to 2.4 μm by 0.9 to 1.2 μm and were smooth walled. The extruded polar tubules seen in many of the spores measured 15 to 22 μm in length. The spores had a thin electron-dense exospore, a thick electron-lucent endospore, and a thin cell membrane surrounding the spore contents. Although the spore contents were dense, the lamellar polaroplast could be discerned at one end and a vacuole could be

FIG. 1. Optical and scanning electron microscopic images of microsporidia spores after treatment by various procedures. (A) Stool smear stained by the quick-hot Gram-chromotrope technique. Bar, 10 μm. (B) Stool smear from the same patient whose stool smear was used in panel A reacted with the anti-*E. intestinalis* serum. Bar, 10 μm. (C) Growth of *E. intestinalis* in cell culture. Note the host cells filled with spores (at the arrows). Differential interference contrast optics were used. Bar, 5 μm. (D) Smear of the culture supernatant from the same flask used for panel C but stained by the quick-hot Gram-chromotrope technique. Note the cell filled with darkly staining spores. Bar, 10 μm. (E) Scanning electron microscopic appearance of *E. intestinalis* from cell culture. Note the delicate thread-like polar tubules at the arrowheads. Bar, 10 μm.
discerned at the opposite end in some spores (Fig. 2B). Cross sections of four to seven coils of the polar tube were also seen in some spores from all five isolates (Fig. 2C). These morphological features were consistent with those of *E. intestinalis*.

**IIF assay.** The spores present in the urine, sputum, and fecal smears, as well as those generated from cell cultures infected with the patient samples described above, reacted with the anti-*E. intestinalis* serum (30) to the same extent (1:4,096) as the spores of strain CDC:V297, the positive control strain used for comparison, and produced a bright apple-green fluorescence. When spores from all cultures under study were treated with anti-*E. cuniculi* (7) and anti-*E. hellem* (29, 32) sera, only slight reactions were observed at a 1:200 dilution and none were observed at a 1:400 dilution, whereas the homologous reaction was recorded at a titer >1:4,096.

**SDS-PAGE and immunoblotting.** When the electrophoretically separated proteins extracted from the reference strains *E. hellem*, *E. cuniculi*, and *E. intestinalis* and the test isolates were stained with the silver reagent, they exhibited a complex pattern producing more than 50 bands ranging from 14 to 200 kDa. Even a cursory analysis of the patterns revealed, in spite of the complexity and a number of shared bands, characteristic

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**FIG. 2. Ultrastructure of *E. intestinalis* within host cells. (A) Infected E6 cell demonstrating the classical septated parasitophorous vacuole (PV), a characteristic feature of *E. intestinalis*, filled with spores (S). M, meront; SB, sporoblast; ST, sporont; N, host cell nucleus. Bar, 1 μm. (B) Spore demonstrating a lamellar polaroplast (PL), a polar tubule (PT) with the anchoring disk (AD), and a nucleus (Nu). PS, polar sac. Bar, 200 nm. (C) Spore with four to five turns of the polar tube (at the asterisk), EX, exospore; EN, endospore. Bar, 200 nm.
banding patterns that could be easily visualized and that differentiated the three species of *Encephalitozoon* represented by the reference strains (Fig. 3). The protein banding patterns of the test isolates were almost identical to those of the reference strain (strain CDC:V297) of *E. intestinalis* (Fig. 3).

After the proteins were transferred to PVDF membranes and after reaction with the three rabbit polyclonal serum samples against *E. intestinalis*, *E. hellem*, and *E. cuniculi*, respectively, Western blot analysis of the separated proteins indicated that the test isolates belonged to *E. intestinalis* (Fig. 4A to C). In the homologous reaction, proteins from the reference strain and the test isolates reacted extensively with the anti-*E. intestinalis* serum, the homologous reaction (*E. cuniculi* with the anti-*E. cuniculi* serum) was strong, with bands at 155, 62, 44, 41, 39, 33, 26, 21, 18, and 16 kDa (Fig. 4C). *E. intestinalis* and the test isolates, however, reacted moderately with both antisera and produced a number of bands in the region between 20 and 160 kDa. The homologous reactions were much stronger and could be easily distinguished from the heterologous ones (Fig. 4A to C). The serum from one of the patients (patient A) showed clear reactivity with extracts from all four isolates as well as with *E. intestinalis* (Fig. 5A). Darkly staining bands migrated at approximately 170, 95, 75, 58, 53, 48, 40, 37, 30, and 28 kDa. The serum from this patient reacted minimally with the proteins of *E. hellem* and *E. cuniculi*. When the membrane was treated with serum from patient B, a fainter reaction was observed with homologous proteins; however, clear bands were distinguished at approximately 170, 86, 84, 50, 48, 40, 37, 30, 28, and 25 kDa. A number of lightly staining bands in the region of 26 to 32 kDa were also seen in reactions with sera from both patients. Minimal reaction was observed with heterologous proteins (Fig. 5B). When patient C’s serum was assayed, it also reacted with protein extracts and again showed a common pattern for proteins from all four isolates and *E. intestinalis* proteins, with the most prominent bands migrating at 50, 39, 37, and 29 kDa. A faint reaction was observed with *E. hellem* and *E. cuniculi* proteins (data not shown). The protein extract from E6 cells reacted minimally with the antisera and produced a number of bands in the region between 20 and 160 kDa. Several darkly staining bands with approximate molecular masses of 78, 74, 64, 45, 42, 35, 32, 28, 25, 22, 19, and 16 kDa.
with all rabbit or human serum samples assayed, producing only a few, light bands in the region of 43 to 90 kDa (Fig. 4 and 5).

**PCR analysis.** The three species-specific PCR primers targeting SSU rRNA-coding sequences selectively amplified fragments diagnostic for *E. intestinalis*, *E. hellem*, and *E. cuniculi*, respectively, with no background from mammalian cellular DNA being found. For example, DNA isolated from cell cultures infected with the test isolates (isolates CDC:V307, CDC: V308, CDC:V309, CDC:V314, CDC:V315, CDC:V324, and CDC:V325) and isolate CDC:V297 reacted with *E. intestinalis*-specific primers only and a diagnostic band of 520 bp was detected in the agarose gel (Fig. 6). PCR primers specific for *E. hellem* amplified only the DNA isolated from cell cultures infected with *E. hellem* and a diagnostic band of 546 bp was detected in the agarose gel, whereas PCR primers specific for *E. cuniculi* amplified only the DNA isolated from cell cultures infected with *E. cuniculi*, resulting in a diagnostic band of 549 bp (data not shown).

**DISCUSSION**

*E. bieneusi* causes most of the microsporidial infections in patients with AIDS and is the most prevalent intestinal parasite of AIDS patients. It infects the small intestine and can spread into the hepatobiliary tree (4, 25, 33). However, recent reports indicate a gradual increase, albeit small, in the number of cases of disseminated microsporidiosis caused by *E. intestinalis* involving the kidneys, eyes, and the lungs, as well as the GI tracts, of people with AIDS. According to Schwartz and Bryan (25), infection with *E. intestinalis* is the second most prevalent microsporidial infection in AIDS patients. The two other species of *Encephalitozoon* (*E. cuniculi* and *E. hellem*) are also known to cause infections of the urogenital, respiratory, and ocular organs but not GI tract infections. Recent reports also indicate that *E. cuniculi* may also disseminate into the brain (19, 35).

Continuous in vitro cultivation of *E. bieneusi* has not yet been achieved, although several isolates of *E. bieneusi* have been cultured for short periods (31). However, a number of isolates of *E. cuniculi*, *E. hellem*, and *E. intestinalis* have been established in continuous culture in a variety of mammalian cell cultures, and the isolates have been compared with each other by antigenic and molecular biology-based analyses (8, 10–14, 16, 23, 24). On the basis of those studies, *E. cuniculi* isolates have been differentiated into three distinct types, e.g., murine, canine, and rabbit (12). On the other hand, *E. hellem* isolates cultured from samples from different sites of infected patients originating from different geographic locales have been shown to be similar on the basis of antigenic and molecular data (5, 23, 24). Several isolates of *E. intestinalis* have also been shown to be similar on the basis of antigenic and molecular data (13). Infection of animals and birds with *E. cuniculi* and *E. hellem*, respectively (1, 9, 16, 18), and infection of animals, including pigs and primates, with *E. bieneusi* (9, 17) have been reported. Recently, however, we have incontrovertible evidence that *E. intestinalis* also causes infection of animals, e.g., pigs, goats, and cows (1a). Many *E. cuniculi* isolates from animals have been established in culture, and efforts are under way to establish in culture other microsporidia from domestic animals.

Here we report the establishment in culture of eight isolates of *E. intestinalis* from five AIDS patients with disseminated microsporidiosis, including infection of the GI tract. All of these isolates readily adapted to culture conditions and grew rapidly and produced high yields of spores within 4 to 5 months. All of these isolates were identified as *E. intestinalis* on the basis of (i) their ultrastructural morphology, namely, the presence of the characteristic septated parasitophorous vacuole and di- and tetrasporous sporogony; (ii) their reactivity in the IIF assay with the rabbit anti- *E. intestinalis* CDC:V297

![Image](http://jcm.asm.org/)
sodium to the same extent as the type species; (iii) PCR analysis, when a 520-bp diagnostic fragment of the SSU rRNA-coding region of *E. intestinalis* was amplified from the DNA of the cell cultures infected with these isolates; and (iv) the similarity of the patterns of the proteins extracted from these isolates to the patterns exhibited by proteins extracted from the type species, as analyzed by SDS-PAGE and immunoblotting.

Although the strongest reactivity always occurred between the homologous systems when the antigenic profiles were analyzed in immunoblots with rabbit anti-*E. hellem*, anti-*E. cuniculi*, and anti-*E. intestinalis* sera, a certain amount of cross-reactivity between the three species also occurred, indicating that all these *Encephalitozoon* species share some common antigenic determinants. Although all isolates of *E. intestinalis* tested had very similar antigenic profiles by Western blotting, minor differences were noted in two of them (strains CDC:V307 and CDC:V308). It is possible that these differences are probably due to relative differences in the growth phases of the organisms, resulting in the expression of different proteins such as surface antigens or intracellular antigens. Since we had selected culture flasks containing parasites of approximately the same age and density, it is possible that these differences are perhaps indications of important variations between isolates of *E. intestinalis*. Further work with proteins derived from culture flasks of different ages as well as flasks with different parasite densities would provide answers to these questions.

When serum samples from three patients were reacted in the immunoblot assay, they clearly reacted very strongly with antigens of the CDC:V297 isolate and those of the test *E. intestinalis* isolates and produced identical profiles. For serum samples from all patients, specific bands were seen at about 50, 48, and 37 kDa. Slight cross-reactivity was also observed with sera from all patients, specific bands were seen at about 50, 48, and 37 kDa. Slight cross-reactivity was also observed with sera from all patients, specific bands were seen at about 50, 48, and 37 kDa. Slight cross-reactivity was also observed with sera from all patients, specific bands were seen at about 50, 48, and 37 kDa.

The patients, however, may have reacted more to the membrane antigens than to the cytosol antigens, hence the responses of our patients are in agreement with those described previously (10, 30).

As has been reported by other investigators (8, 13, 14, 16, 21, 23, 30, 32), in vitro cultivation of microsporidial organisms that infect humans and animals is invaluable for several reasons. For example, it facilitates (i) understanding of the biology of the parasite and the host-parasite relationships, (ii) the development of immunologic and molecular reagents for use in clinical diagnosis, (iii) the development of assays for screening newer and promising therapeutic agents, and (iv) the development of antigenic and molecular markers for isolates that may be useful in molecular epidemiology, particularly in tracing the sources of the causal agent, and thus that will be helpful in formulating preventive strategies.

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REFERENCES


