Continuous culture of Cryptosporidium parvum using hollow fiber technology

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A B S T R A C T

Diarrheal disease is a leading cause of pediatric death in economically low resource countries. Cryptosporidium spp. are the second largest member of this group and the only member for which no treatment exists. One of the handicaps to developing chemotherapy is the lack of a reproducible long-term culture method permitting in vitro drug screening beyond 48 h. We have adapted the well-established hollow fiber technology to provide an environment that mimics the gut by delivering nutrients and oxygen from the basal layer upwards while allowing separate redox and nutrient control of the lumen for parasite development. Using this technique, oocyst production was maintained for >6 months, producing approximately 1 x 10⁷ oocysts ml⁻¹ day⁻¹, compared with 48 h with a yield of 1 x 10⁴ - 10⁶ oocysts ml⁻¹ in two-dimensional cultures. Oocysts, after 4 and 20 weeks in culture, produced a chronic infection in a TCR⁻ deficient mouse model. In vivo infectivity of oocysts was confirmed using oocysts from a 6 week culture in a dexamethasone immunosuppressed mouse model.

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1. Introduction

Cryptosporidium spp. are the second (after rotavirus) leading cause of diarrheal disease and death in infants in economically low resource countries (Liu et al., 2012), and remains the only member of the major diarrheal diseases for which no consistently effective therapy is available. In parts of Asia and Africa as many as 31.5% of all children under 2 years of age are infected with the parasite, posing a significant health risk (Kotloff et al., 2013; Checkley et al., 2014). There are five species responsible for infections in humans, Cryptosporidium hominis, Cryptosporidium parvum, Cryptosporidium meleagridis, Cryptosporidium canis and Cryptosporidium felis, of which C. hominis or C. parvum are the most common (Xiao, 2010). Infection of the epithelial cells of the gastrointestinal tract by this parasite results in an acute watery diarrhea, which can become life threatening in children, the elderly and immune-compromised individuals (Mathur et al., 2013). Malnourished children are at a greater risk of death from the infection and those that survive are at greater risk of stunted growth and wasting (Banwat et al., 2003; Hamedi et al., 2005; Checkley et al., 2014). The underlying cause of disease transmission is poor hygiene and contaminated water sources (Mollov et al., 2011; Melbak et al., 2013). In some countries it is endemic due to the poor sanitation and lack of appropriate health practices. Despite the significant morbidity and mortality associated with cryptosporidiosis, drug discovery lags behind that of many other infectious diseases (Striepen, 2013). Nitazoxamide, the only FDA-approved drug for treatment of cryptosporidiosis, is not effective in immune-compromised individuals and is not routinely used in standard care in countries such as India (Abubakar et al., 2007).

The parasite has a multistage life cycle producing thick walled oocysts that are shed from infected individuals and act as the infective stage; other stages include motile sporozoites, merozoites, macro- and microgametocytes, as well as thin-walled oocysts that are responsible for autoinfection. In common with other members of the Apicomplexa, Cryptosporidium spp. require a host cell to complete the life cycle. The parasite is capable of parasitising a number of in vitro cultured cell lines (Upton et al., 1994) producing thin walled oocysts that are not infective to the primary host.
Significant advances have been reported, (reviewed by Arrowood (2008)). More recent is the use of an HCT-8 organoid model to replicate the intestine (Alcantara Warren et al., 2008), and the use of primary cultured intestinal epithelial cells (Castellanos-Gonzalez et al., 2013) which have extended the in vitro infection to >5 days. Recently Varughese et al. (2014) described a model using small intestinal epithelial cells, FHS 74 Int, demonstrating improved infection kinetics that has great potential for host-parasite studies. It was the goal of this study to extend the in vitro culture of *C. parvum* to >1 month, permitting long-term drug studies to be performed. We considered the following issues as critical to the development of a long-term culture system. The presence of two controlled environments (biphasic) would provide host cells with oxygen and nutrients from the basal layer, while allowing a low oxygen nutrient rich environment to be developed on the apical surface where the parasite would be present, thus mimicking the situation in situ. The development of such a system provides the added advantage of allowing the development of a parasite-specific growth medium, which is not possible in current two-dimensional (2D) cultures where emphasis is directed to maintaining host cell growth to support the parasite burden. The parasite completes its life cycle within the host intestine, a low oxygen environment. In addition, a microaerophilic parasite metabolism is inferred from genome sequence information (Abrahamson et al., 2004), and is supported by several studies demonstrating the loss of typical mitochondria (Hieinz and Lithgow, 2013), and dependence upon a low oxygen metabolism, reviewed by Zhu (2008). These observations prompted us to modify the standard medium developed by Upton et al. (1994) to include reducing agents and a mixture of fatty acids, which have been shown to be important in the growth of various anaerobic protozoa (Yichoy et al., 2011).

2. Materials and methods

2.1. Culture of *HCT-8* host cells

Human ileocecal colorectal adenocarcinoma cells (*HCT-8* (HRT-18), ATCC CCL-244) were used to provide a host cell layer on the extra-capillary surface of the fibers. MEM plus supplements and serum (MEMSS) (Upton et al., 1994) circulated through the hollow fibers and glucose levels were recorded using an Accu-Chek Active monitor (Hoffmann-La Roche Ltd., Indianapolis, IN, USA). The pH was monitored by sampling 5 mL of the recirculating medium using a Corning 440 pH meter (Corning, NY, USA).

2.2. Real-time quantitative reverse transcription-PCR (*qRT-PCR*) analysis of parasite 18S rRNA

Total RNA was isolated from 5 mL of *C. parvum* and host cells collected from the cartridge. The sample was centrifuged at 6449g for 5 min in a swing-out rotor (Beckman–Coulter, Indianapolis, IN, USA), then the pellet was resuspended in 200 μL of PBS and washed three times; 50 μL was removed and the remaining centrifuged, resuspended in lysis buffer (iScript RT–qPCR sample preparation reagent, Bio-Rad Laboratories, Hercules, CA, USA) and subjected to six cycles of freezing-thawing using liquid nitrogen and a 65 °C heating block. Total RNA was isolated using RNeasy (Qiagen Inc., Valencia, CA, USA) and the total RNA was determined using a Qubit 3.0 fluorometer (Life Technologies, Thermo-Fisher Scientific Inc., Waltham, MA, USA). The qRT-PCR analysis was also performed for the Iowa isolate (Bunch Grass Farms, Deary, ID, USA), diluted to give 10^3, 10^4 and 10^5 oocysts. The ΔCt for parasite 18S rRNA was determined by subtraction of the Ct for h18S rRNA from the Ct for parasite C176 18S rRNA.

2.3. Immunocompromised and immunosuppressed mouse models

Oocysts collected from the culture system were tested for the ability to infect two well established mouse models. Oocysts were collected at specified times from the culture system, stored in 2.5% potassium dichromate at 4 °C for less than 1 month, washed with distilled water three times to remove the potassium dichromate, and enumerated using a hemacytometer prior to use.

The first model employed nine 6 week old female TCR-α−/− deficient mice on a C57BL/6 background (Jackson Laboratories, Bar Harbor, ME, USA); mice were segregated into groups of three; three were infected per os with 10^4 oocysts in 200 μL of PBS collected after 4 and 20 weeks in culture; three were infected with 10^4 oocysts of the Iowa isolate used to establish the culture; three were administered 200 μL of PBS as negative controls. All mice were housed in metabolic chambers (VWR, Atlanta, GA, USA) and every 24 h feces were removed, weighed and placed into potassium dichromate for storage. After 7 days, mice were euthanised by CO2 asphyxiation as recommended by the American Veterinary Medical Association (AVMA) (Pace University, NY, USA) and intestinal sections from the distal ileum–cecum and proximal colon removed and fixed in 10% neutral buffered formalin (3.7–4.0% formaldehyde in 33.3 mM monosodium phosphate and 45.8 mM disodium phosphate buffer).

The second model employed 15 female CD-1® Igcs mice (Charles River Laboratory, Wilmington, MA, USA) aged 3–5 weeks, weighing 17–22 g each, which were immunosuppressed with dexamethasone 21-phosphate (D1159, Sigma, USA) administered ad libidum in drinking water (16 μg/mL) from 4 days prior to inoculation to the end of the study. Mice were separated into the following three groups of five per group, and infected per os with 200 μL of sterile water containing either 10^6 oocysts from 6 weeks in culture (group 1), the parent Iowa isolate used to initiate the culture (group 2) or were an uninfected control group (group 3). Mice were monitored twice daily following the infection and weighed three times per week; fecal samples were collected daily for 17 days and fecal smears stained using the modified Kinyoun acid fast method (Ma and Soave, 1983). After 17 days, mice were euthanised by CO2 asphyxiation as recommended in the AVMA (Tufts University, MA, USA) and gastrointestinal sections removed for histopathological examination. The ileum, cecum and proximal colon were fixed in 10% natural buffered formalin and processed for H&E stains.

Animal studies were performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health, USA under the Animal Welfare Assurance numbers A3112-01 (Pace University, Haskins Laboratories, NYC, USA) and A4059-01 (Tufts University, Cummings School of Veterinary Medicine, USA). Animal experiments were performed in accordance with the procedures approved by
the Institutional Animal Care and Use Committees of Pace University, NYC and Tufts University, Cummings School Veterinary Medicine, MA, USA.

2.4. Staining with fluorescently labeled antibodies

Samples (5–10 mL) were removed from the extra capillary space of the cartridge by displacement using two 10 mL sterile syringes. Samples were centrifuged at 16,162g for 1 min (Sorvall Biofuge Fresco, Thermo Fisher Scientific) and the pellet resuspended in Dulbecco's PBS. Oocysts were stained using 50 µL of FITC-labeled mouse monoclonal antibody to C. parvum oocyst surface proteins (Crypt-a-Glo®. Waterborne Inc., New Orleans, LA, USA) and incubated in the dark at room temperature for 30 min, after which they were washed twice with PBS and examined using a fluorescence microscope (Nikon Optiphot, Melville, NY, USA) with an excitation λ 410–485 nm, and an emission λ 515 nm. Samples were counter-stained with a fluorescent-labeled polyclonal antibody specific for sporozoites and other motile intracellular stages (Sporo-Glo®, Waterborne Inc.), and examined using an excitation λ 535–550 nm, and an emission λ 580 nm.

2.5. Preparation of oocysts and excystation

Cryptosporidium parvum oocysts were purified from the other stages (empty oocyst shells, host cells, etc) using isopycnic percoll gradient centrifugation (Arowood and Sterling, 1987) consisting of 3 mL of 1.04 g mL⁻¹ percoll overlaid onto 5 mL of 1.08 g mL⁻¹ percoll. One millilitre samples from the cartridge were layered onto these gradients and centrifuged at 400g for 30 min. The oocysts formed a light colored band approximately 1/3 of the distance from the bottom of the tube and were then aspirated and washed with PBS by centrifugation at 1000g for 10 min. Sporozoites were obtained by excysting oocysts in MEM containing 0.25% trypsin, 0.75% taurodeoxycholate at 37 °C in a shaking water bath.

2.6. Polyamine analysis

Polyamines were separated and identified by reverse phase HPLC using a Perkin Elmer LC410 system coupled to a C18 10 µm column (4.5 × 250 mm) at a flow rate of 1 mL min⁻¹. The method employed a 40 min discontinuous gradient starting with 90% (v/v) buffer A (0.1 M NaH₂PO₄, 8 mM octanesulfonic acid, and 0.05 mM EDTA) and 10% (v/v) buffer B (acetonitrile). The initial conditions were held for 5 min before changing to 20% buffer B for 15 min, 40% buffer B for a further 10 min, and then returning to 10% buffer B for 10 min (Morada et al., 2013). Samples and standards were mixed 1:1 with 1.5 mM l-threonine, MA, USA.

2.7. Enzyme assays

Spermidine/spermine N⁴-acetyltransferase (SSAT) was determined using 100 µM bicine buffer (pH 8.0) containing 17 µM [1-¹⁴C]Acetyl-CoA (60 µCi/mmol) and supplemented with 50 µM unlabeled acetyl-CoA, 500 µM spermine and 25 µg of protein. The reaction was stopped after 30 min with ice-cold 50 mM hydroxylamine, placed in a boiling water bath for 3 min, cooled and centrifuged at 9000g for 1 min to remove precipitated protein. The supernatant (50 µL) was spotted onto filter discs, dried and washed with 6 × 200 mL changes of distilled water to remove unreacted [1-¹⁴C]-acetyl-CoA, with a final wash with 200 mL of methanol. The dried discs were placed in 10 mL of Omni Fluor and the radioactivity present as [¹⁴C]-acetylperoxime was counted using a Beckman Tri-Carb 1600CA liquid scintillation counter (Perkin Elmer Life Sciences, Waltham, MA, USA). Blanks containing [1-¹⁴C] acetyl-CoA and spermine without protein were also analyzed and subtracted from the experimental results (Morada et al., 2013).

Spermidine acetyl transferase (SAT) was assayed using the same method for SSAT but using 500 µM spermidine in place of spermine (Keithly et al., 1997).

S-adenosyl-l-methionine decarboxylase (putrescine stimulated) was determined by measuring the ¹⁴CO₂ produced from 0.5 µCi [¹⁴C-carboxyl]-S-adenosylmethionine (55 mCi/mmol) incubations containing 10 mM Tris–HCl pH 6.5 and 1 mM putrescine (Keithly et al., 1997). The ¹⁴CO₂ released in 60 min was trapped on benzenthionium hydroxide soaked filter paper and counted by scintillation using a Beckman Tri-Carb 1600CA liquid scintillation counter (Perkin Elmer Life Sciences).

2.8. Electron microscopy

For Scanning Electron Microscopy (SEM), samples were fixed in 2.5% glutaraldehyde, 0.1 M sodium cacodylate pH 7.4, dehydrated through a graded series of ethanol, critical point dried using liquid carbon dioxide in a Tousimis Samdri 790 Critical Point Drier (Rockville, MD, USA) and sputter coated with gold–palladium in a Denton Vacuum Desk-2 Sputter Coater (Cherry Hill, NJ, USA). The samples were examined in a JEOL JSM6400 Scanning Electron Microscope (Peabody, MA, USA), using an accelerating voltage of 10 kV. Images were recorded with AnalySIS, Soft Imaging Systems (Munster, Germany).

2.9. Statistics

The statistical comparison of the differences among experimental groups was made by Mann–Whitney U test using GraphPad Prism 5 software (GraphPad Software, San Diego, CA, USA). Results were considered to be statistically significant at P < 0.05.

3. Results

We describe a method for the long-term in vitro culture of C. parvum in a simulated gut-like environment using hollow fiber technology. Human ileocecal adenocarcinoma (HCT-8; ATCC CCL 244) cells were selected as the cell line to support growth of C. parvum because previous studies found this cell line supported a greater infection than alternative ones (Upton et al., 1994; Meloni and Thompson, 1996), a feature that does not diminish with age of the cells (Sifuentes and DiGiovanni, 2007). HCT-8 cells have also been shown to differentiate into organoids that express microvilli and desmosomes characteristic of normal intestinal tissue (Carvalho et al., 2005). We considered these features ideal for a long-term in vitro culture system. A 20 ml hollow fiber cartridge (CS2011; FiberCell® Systems, Inc., Frederick, MD, USA), conditioned according to the manufacturer’s instructions, was inoculated with 10⁴ HCT-8 cells via the side port, and a 125 mL reservoir containing MEMSS (Upton et al., 1994) circulated through the hollow fibers at a flow rate of 51 mL min⁻¹ (setting five; Fig. 1). The fiber surface area (2100 cm²) provides efficient nutrient, metabolite and gas exchange between the HCT-8 cell layer and the re-circulating medium. The cartridge, pump and growth medium were kept inside a

Fig. 1. Diagram of the hollow fiber system. The cartridge is composed of 200 μm diameter polysulfone hollow fibers with a 20 kDa molecular weight cut-off. The growth medium (MEM plus supplements and serum, MEMSS) was pumped through the fibers to supply nutrients and oxygen to, and remove waste products from, the host cells (HCT-8) which colonize the outside of the hollow fibers. The extra capillary space was inoculated with Cryptosporidium parvum and the MEMSS medium in this environment was modified to include additives that promote parasite growth - lipids, redox buffers and vitamins (as described in Section 3). Additions can be made and parasites removed from the extra capillary space using the side port shown on the top of the cartridge.

Fig. 2. Daily glucose and cell counts from the hollow fiber culture system. (A) The cartridge was inoculated with 10⁸ HCT-8 cells and allowed to achieve confluence for 11 days. On day 12 the cartridge was inoculated with 10⁶ Cryptosporidium parvum oocysts and the glucose levels monitored daily. Twenty-four hours p.i. there was an initial increase in medium glucose levels associated with infection of host cells by C. parvum. The increase in glucose levels may be due to disruption of the host cell monolayer, resulting in lysis and release of stored glycogen. (B) The pattern of medium glucose levels (bars) cycled for approximately 20 days and exhibited a dampened oscillation, indicative of loss of synchrony of the parasitemia of host cells by C. parvum. The line at 350 mg dL⁻¹ is the glucose concentration of the MEM plus supplements and serum (MEMSS) growth media circulated through the microfibers. (C) Seven days p.i., half of the extra capillary volume (7.5 mL) was removed for enumeration of oocysts and sporozoites. This was repeated at 3–4 day intervals; asterisks indicate removal of 50% of the extra capillary volume and replacement with fresh medium. Oocysts were stained with a fluorescent-labeled mouse anti-C. parvum oocyst wall monoclonal antibody and counted using a hemacytometer by fluorescence microscopy with an excitation λ of 410 nm–485 nm and an emission λ of 515 nm. (D) Sporozoites were counted using a hemacytometer after staining with a polyclonal antibody to intracellular and motile stages (Sporo-Glo®; Waterborne Inc., New Orleans, LA, USA) using an excitation λ 535 nm–550 nm and an emission wavelength of 580 nm.
5% CO₂ incubator maintained at 37 °C. At 24 h intervals the glucose (Fig. 2A) and pH levels (data not shown) were recorded. When the glucose fell below 50% of the fresh MEMSS, the reservoir volume was doubled to 250 mL, and this process was continued until a reservoir volume of 1 L was achieved (8–11 days); when the HCT-8 cells were at full confluent density the pump rate was increased to 105 ml/min⁻¹ (setting 10).

3.1. Redox and nutrient requirements

The provision of oxygen and nutrients for the host cells from the basal layer upwards provided us with the opportunity to develop a C. parvum culture medium that more closely approximates the intestinal lumen on the apical epithelial surface. Cryptosporidium parvum parasitises the lumen of the gut that has sub-micromolar oxygen tensions and the biochemistry of the parasite supports the presence of an anaerobic metabolism; the C. parvum genome reveals a lack of a Krebs cycle and oxidative phosphorylation and a striking increase in the number of amino acid and fatty acid transporters (Abrahamsen et al., 2004; Zhu, 2008). Addressing these nutritional and environmental requirements is necessary to developing a long-term culture system. To this end we developed a redox buffer with the aim of creating a low redox environment developing a long-term culture system. To this end we developed a redox buffer with the aim of creating a low redox environment that would mimic that present in the gut (Circu and Aw, 2011).

The redox buffer contained 20 mg mL⁻¹ each of glutathione, taurine, betaine and cysteine prepared in nitrogen gassed distilled water. This mix resulted in a higher oocyst production than the addition of a mix containing 20 mg mL⁻¹ each of DTT/cysteine or 20 mg mL⁻¹ each of mercaptoethanol/cysteine. Many parasites have specific lipid requirements that may not be provided by the addition of serum alone, and there is some evidence in the literature that C. parvum also has a selective lipid requirement (Mazumdar and Striepen, 2007; Bushkin et al., 2013) and to this end we focused on developing a lipid mix composed primarily of omega-3 lipids. The composition and final amount was determined by titration of individual components of the mix that produced the maximum number of oocysts. The lipid mix was composed of 1.5% deoxycholate, 6.7 mg mL⁻¹ oleic acid, 10 mg mL⁻¹ phosphocholine, 1.6 mg mL⁻¹ α-linolenic acid, 6.8 mg mL⁻¹ eicosapentaenoic acid, 2 mg mL⁻¹ docosahexaenoic acid, 18 mg mL⁻¹ cholesterol (dissolved in ethanol and mixed 1:1 with Tween 80). The extra capillary space of the cartridge (Fig. 1) was flushed with 50 mL of MEMSS (Upton et al., 1994) containing 1 μg mL⁻¹ resazurin (7-hydroxy-3H-phenoxazin-3-one-10 oxide) as a redox indicator, in place of phenol red, and containing the redox buffer and lipid mix as additives.

The cartridge was incubated for 2 h at 37 °C, during which time the resazurin indicator changed from a pink color (redox potential $\geq -51$ mV) to colorless, indicating a redox potential of $\leq -110$ mV. The redox potential was determined from a standard graph of the difference in the OD₅₇₀ nm to that at OD₆₀₀ nm versus the buffer redox potential in mV.

The cartridge was inoculated with $10^6$ C. parvum oocysts (Iowa isolate), and 3 mL samples removed daily by displacement using 3 mL of the MEMSS plus redox and lipid additives. Glucose consumption, pH, redox conditions and microscopic examination were recorded to determine the integrity of the host cell feeder layer and parasite development. The medium glucose levels present in the host cell medium exhibited a dampened oscillation over the first 10 days of the infection (Days 12–22, Fig. 2A) that may be the result of initial synchrony of disruption of the host cell monolayer caused by the infection. By day 23 the glucose consumption was approximately 50% of the added glucose in 24 h (Fig. 2B). The glucose concentration of the medium in the hollow fibers, providing nutrients to the host cells varied from 50 to 250 mg/dL, whereas the glucose concentration in the cartridge space (providing nutrients to the parasites) varied between 270 and 334 mg/dL over the duration of the study. The pH changed by $-0.81$ pH units over 48 h in the host cell medium, whereas it showed minor variation in the cartridge space (pH 6.99–7.09). These measurements indicate that there is a difference between the environments inside the cartridge (parasite environment) and that inside the hollow fibers (host cell environment). Samples inside the cartridge (5–10 mL) were removed weekly and examined under light microscopy (Fig. 3A) after duel staining using fluorescein-labeled polyclonal antibody for intracellular and motile stages (Sporo-Glo®; Fig. 3B), and FITC-labeled monoclonal antibody for oocysts (Crypt-a-Glo®; Fig. 3C). Parasites achieved a density of $1 \pm 0.1 \times 10^6$ oocysts per mL (Fig. 2C), and $8 \pm 2 \times 10^7$ intracellular motile stages per mL (Fig. 2D), in the hollow fiber culture system compared with $0.8 \times 10^6$ per mL oocysts in 25 cm² flasks, and continued growing for >6 months (at time of submission) compared with 48 h when grown in 2D tissue culture.

3.2. Parasite enumeration by qRT-PCR

A second culture system was established to determine the effects of flow rate on parasite numbers and host cell maintenance (Fig. 4A). Parasite and sloughed host cell numbers were evaluated by determining the Ct of C18S rRNA and h18S rRNA for samples collected from the culture system. The h18S rRNA had minor variation over 15 weeks, and was therefore used to normalise the par...
asite \Delta C_T \text{ values. The } \Delta C_T (C_p 18S rRNA - h18S rRNA) \text{ values were compared with a standard graph prepared using } 10^5-10^8 \text{ oocysts, enabling determination of parasite numbers from the culture system (Fig. 4B). Using this method, parasite numbers varied from } 10^5 \text{ to } 5 \times 10^8 \text{ per mL over } 15 \text{ weeks (Fig. 4A). Parasite numbers showed a cyclical rhythm of 3–4 weeks. After 15 weeks the numbers of host cells in the sample collected increased, which may be indicative of a chronic infection resulting in significant damage to the host cell layer.}

3.3. Cryptosporidium parvum oocysts infect immunosuppressed and immunodeficient mice

TCR-α-deficient mice infected with } 10^4 \text{ oocysts (Fig. 5) harvested after 4 and 20 weeks in culture (Table 1); and dexamethasone immunosuppressed mice infected with } 10^6 \text{ C. parvum oocysts harvested after 6 weeks in culture (Table 2) resulted in oocyst shedding in the feces at comparable numbers to the control Iowa isolate used to initiate the culture system. Mice infected with oocysts obtained from the 6th week of the culture system and the

![Image of a graph showing parasite numbers over weeks](image1)

![Image of FITC-labeled monoclonal antibody stained Cryptosporidium parvum oocysts](image2)

**Table 1**

<table>
<thead>
<tr>
<th>Weeks since culture</th>
<th>Days p.i.</th>
<th>Sample</th>
<th>Uninfected control</th>
<th>Cultured oocysts</th>
<th>Iowa oocysts</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 weeks</td>
<td>4</td>
<td>0 ± 0</td>
<td>1.5 ± 0.7</td>
<td>2.5 ± 3</td>
<td>1 ± 1</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>0 ± 0</td>
<td>3 ± 2</td>
<td>7.7 ± 8.9</td>
<td>15.8 ± 6.3</td>
</tr>
<tr>
<td>6</td>
<td>6</td>
<td>0 ± 0</td>
<td>13.3 ± 8.6</td>
<td>29.8 ± 19.7</td>
<td>42.3 ± 26.4</td>
</tr>
<tr>
<td>20 weeks</td>
<td>12</td>
<td>0 ± 0</td>
<td>9.6 ± 8.2</td>
<td>69.8 ± 32.3</td>
<td>53.9 ± 39.2</td>
</tr>
<tr>
<td>14</td>
<td>14</td>
<td>0 ± 0</td>
<td>31.2 ± 12.4</td>
<td>19.7 ± 11.6</td>
<td>23.4 ± 16.6</td>
</tr>
<tr>
<td>17</td>
<td>17</td>
<td>0 ± 0</td>
<td>29.8 ± 19.7</td>
<td>42.3 ± 26.4</td>
<td>53.9 ± 39.2</td>
</tr>
<tr>
<td>19</td>
<td>19</td>
<td>0 ± 0</td>
<td>69.8 ± 32.3</td>
<td>19.7 ± 11.6</td>
<td>23.4 ± 16.6</td>
</tr>
<tr>
<td>21</td>
<td>21</td>
<td>0 ± 0</td>
<td>159.2 ± 41.8</td>
<td>87.5 ± 51.4</td>
<td>121.5 ± 62.5</td>
</tr>
</tbody>
</table>

* The colon content was analyzed.

**Table 2**

<table>
<thead>
<tr>
<th>Group</th>
<th>Mouse number</th>
<th>Days p.i.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not treated</td>
<td>m1</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>m2</td>
<td>14</td>
</tr>
<tr>
<td>Cultured oocysts</td>
<td>m1</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td>m2</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>m3</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>m4</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>m5</td>
<td>15</td>
</tr>
<tr>
<td>Iowa oocysts</td>
<td>m1</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>m2</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>m3</td>
<td>19</td>
</tr>
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<td></td>
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<td>19</td>
</tr>
<tr>
<td></td>
<td>m5</td>
<td>19</td>
</tr>
</tbody>
</table>

parent Iowa isolate demonstrated an average 17% weight loss compared with uninfected immunosuppressed control mice after day 10 (Fig. 6A). Post 10 days, oocyst shedding was detected in fecal samples from all of the mice infected with the culture system (Table 2). Oocyst shedding intensities were similar between the mice from both the culture-derived oocyst infection and the parent-derived oocyst infection. On day 14 the colon content was analyzed for mouse four from the culture-derived oocyst group (Table 2). Histological sections of the intestine (ileum) from mice 17 days p.i. demonstrated the presence of \textit{C. parvum} life cycle stages in the terminal ilea and proximal colon (Fig. 6B).

3.4. \textit{Cryptosporidium parvum} cultures have unchanged polyamine biosynthesis

\textit{Cryptosporidium parvum} oocysts collected from the hollow fiber cartridge were purified by percoll gradients (Arrowood and Sterling, 1987) and the sporozoite stage obtained as described in Section 2. Homogenates of sporozoites were subjected to various biochemical analyses for comparison with data obtained using oocysts supplied from calf infections. We previously characterised the intracellular polyamine levels and their respective biosynthetic enzymes from \textit{C. parvum} oocysts and sporozoites (Keithly et al., 1997; Morada et al., 2013). In many of the Apicomplexa the levels of putrescine and spermidine have been shown to increase dramatically during infection of the host cell (Niemand et al., 2012). These polycaitonic molecules are critical to parasite growth and development, and play a key role in the production of hypusine and therefore activation of eukaryotic initiation factor (eIF-5A), which is present in the parasite (Mittal et al., 2014). For this reason we evaluated parasites harvested from the cartridge for polyamine content and polyamine biosynthetic enzyme activity. The major polyamines putrescine, spermidine and spermine were determined by HPLC (Morada et al., 2013) to be $2.6 \pm 0.9$, $41.7 \pm 5.3$ and $23.5 \pm 2.1$ pmol per $10^6$ cells ($\pm$ S.D. of triplicate results), respectively, which compared with 2.0, 47.8 and 20.0 pmol per $10^6$ cells previously observed (Morada et al., 2013), and are within the S.D. of experimental variation. The activity of polyamine biosynthetic enzymes was also measured in parasites collected from the cartridge. SSAT, S-adenosyl-L-methionine decarboxylase (SAMdc, putrescine stimulated) and SAT had activities of $0.15 \pm 0.05$, $0.009 \pm 0.002$ and $0.21 \pm 0.04$ nmol min$^{-1}$ (mg of protein)$^{-1}$, respectively ($\pm$ S.D. of triplicate results), which compared favorably to published (Keithly et al., 1997) values of 0.12, 0.006 and 0.17 nmol min$^{-1}$ (mg of protein)$^{-1}$, respectively.

3.5. Electron microscopy of infected host cells grown on hollow fibers

The structure of the host cell layer was determined after 8 weeks of parasite growth. The cartridge case was opened and the hollow fibers cut into thin sections, fixed as described in Section 2, and examined by SEM (Fig. 7). After 8 weeks of growth on
hollow fibers, the intestinal epithelial cells had differentiated into crypt-villus units (Fig. 7). Developmental parasite stages were evident as sacs covering the epithelial cells, and at this early stage populated mainly the proximal end of the cartridge (Fig. 7A) where they were introduced. The epithelial layer has many craters and scars (Fig. 7A) characteristic of those previously observed in Cryptosporidium sp.-infected intestinal sections (Fayer, 2008). At the 8 week stage the distal end of the cartridge has a lower parasite load (Fig. 7B), which was mainly due to oocysts produced from the initial infection as the parasites move down the cartridge, mimicking the situation found in the gut of infected animals (Fayer, 2008). Oocysts collected from the cartridge were identified using a specific FITC labeled mouse anti C. parvum oocyst monoclonal antibody, and excysted (as described in Section 2) producing motile sporozoites. Oocysts (10^6) collected from the cartridge were used to infect a HCT-8 monolayer in 2D cultures using 75 cm² flasks containing 10 mL of MEM plus 10% horse serum. The medium was removed 3 h p.i. and replaced with fresh MEM plus 10% horse serum. After 24 h at 37 °C in a 5% CO₂ incubator, the flasks produced 2.6 × 10^6 oocysts compared with 3.2 × 10^6 oocysts ml⁻¹ in control flasks infected with 10^6 oocysts obtained from neonatal calves.

4. Discussion

Building on recent advances in the genomic, biochemical and in vitro culture methods of Cryptosporidium research, we have developed a method for the long-term in vitro culture of C. parvum using hollow fiber technology. In developing the method, we concentrated on a design that would permit control of two separate environments (biphasic), permitting an aerobic nutrient supply to the host cells, while also allowing a separate anaerobic nutrient rich medium to be established for the parasite environment. The ability to generate the intestinal redox conditions is a critical factor in the success of the method. It has been determined that under physiological conditions the Eh for the oxidised/reduced (GSH/GSSG) glutathione couple is between −260 and −200 mV (Kemp et al., 2008). Intestinal epithelial cells have a highly reduced cytosolic glutathione pool (Eh for GSH/GSSG of −260 mV) whereas the inter-membranous space is slightly more oxidised (Eh of −255 mV) and the endoplasmic reticulum matrix is highly oxidised (Eh −170 to −185 mV), ensuring correct folding of nascent proteins (Circu and Aw, 2011). The luminal glutathione pool is efficiently taken up by the intestinal epithelial cells and is important for normal growth and development of these cells. It has been shown that changes in the redox potential correlate with intestinal cell phenotypes. A reducing redox environment (Eh −260 mV to −240 mV) favors proliferation, whereas an oxidised one (Eh −220 mV to −200 mV) favors differentiation and under highly oxidised conditions growth arrest occurs (Eh −170 mV), which is followed by necrosis or apoptosis at an Eh −150 mV (Jonas et al., 2002). The luminal cysteine pool is also a key player in redox control as a significant proportion of the extracellular glutathione pool is hydrolysed to cysteine (Kemp et al., 2008). Redox control of extracellular surface proteins by luminal cysteine is believed to regulate signaling processes at the apical plasma membrane (Jonas et al., 2002). Glutamine was included as an additive because it has been shown that this amino acid contributed to a more reduced extracellular cysteine pool, resulting in enhanced CaCo-2 cell growth by activation of redox signaling at the plasma membrane (Jonas et al., 2002). In addition glutathione is essential for the intestinal elimination of luminal peroxidised lipids (Circu and Aw, 2011), which if not removed would deter parasite colonisation of the epithelial cells. In this regard C. parvum sporozoites infect host cells in the crypts of the villi and develop into meronts as the epithelial cell migrates to the apical tip where the merozoites escape from the parasitophorous vesicle.

Biochemical and genomic data agree that C. parvum lacks a typical mitochondrion, oxidative phosphorylation pathway and a Krebs cycle, relying upon glycolysis and fatty acid oxidation for energy production (Mazumdar and Striepen, 2007; Zhu, 2008). The low redox environment of the media employed in this study would therefore favor the growth and development of a parasite with a fermentative metabolism.

In support of this hypothesis, the yield of parasites obtained from the culture system was significantly improved by including a lipid supplement which included the omega-3 fatty acids, α-linolenic acid, eicosapentaenoic acid and docosahexaenoic acid. Recently it has been shown that C. parvum oocyst walls are acid-fast and contain a complex set of triglycerides rich in polyhydroxy and long fatty acyl chains that may be synthesized by a polyketide synthase (Zhu et al., 2010; Bushkin et al., 2013). This wax-like rigid bilayer is impermeable to disinfectants and environmental stress, and is essential for continued propagation of the parasite. Cryptosporidium parvum lacks fatty acid synthase II biosynthetic machinery, suggesting they are dependent upon fatty acid salvage from the host (Zhu et al., 2010). However the parasite does possess a fatty acid synthase I complex that resembles a polyketide synthase that is proposed to act as a fatty acyl elongase and is responsible for the production of the long acyl chain fatty acids required for the formation of the oocyst cell wall (Mazumdar and Striepen, 2007). Hence it is likely that the medium chain fatty acids added to the extra capillary space favor formation of the thick cell wall oocyst stage and propagate the infection. Thick walled oocysts are shed in the feces of infected individuals and serve to transmit the infection; once ingested these thick walled oocysts pass through the stomach and produce motile sporozoites in the small intestine that infect the host epithelial cells, resulting in an asexual life cycle that produces Type I meronts which reinfect epithelial cells; the problem with current culture conditions is that the merozoites produced at this stage fail to enter the sexual cycle, producing Type II meronts that ultimately produce zygotes which transform into thin walled oocysts, resulting in auto-infection or thick walled oocysts that are excreted and are responsible for disease transmission. That the oocysts from the culture system undergo a complete life cycle is demonstrated using the well described and validated mouse model that employs either the dexamethasone immunosuppressed mice, or the genetically modified TCR-/-immunocompromised mouse model.

Hollow fiber technology has successfully been used to produce a constant supply of large numbers of Plasmodium falciparum for pharmacokinetic analysis of potential chemotherapeutic compounds (Bakshi et al., 2013) which mimicked the dynamic fluctuations of the drug in vivo, providing clinically relevant data that could be used to select antimalarial lead compounds based upon total exposure, peak concentration or time above the minimum inhibitory concentration, providing preclinical pharmacokinetic data in the absence of animal models. We are currently adapting our method to perform a similar pharmacokinetic function that will allow us to obtain critical information needed for the development of chemotherapeutic agents to treat disease caused by this parasite.

Hollow fiber technology provides several unique features: (i) a large surface area for metabolite and gas exchange, which are needed for efficient growth of host cells; (ii) the creation of a biphasic medium providing an oxygen rich nutrient supply to the basal layer of the host cells, while permitting the provision of an anaerobic nutrient rich supply to the apical side mimicking the gut; (iii) the ability to obtain high numbers of in vitro cultured C. parvum oocysts for biochemical and molecular studies; (iv) study of the host-parasite relationship in a long-term in vitro infection;
(v) the ability to obtain in vitro preclinical pharmacokinetic data, providing a unique method for drug selection; and (vi) this method can also be used for analysis and preparative isolation of *Cryptosporidium* growth factors.

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