A Most-Probable-Number Assay for Enumeration of Infectious Cryptosporidium parvum Oocysts

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Cryptosporidium is globally established as a contaminant of drinking and recreational waters. A previously described cell culture infectivity assay capable of detecting infectious oocysts was adapted to quantify viable oocysts through sporozoite invasion and clustering of foci. Eight experiments were performed by using oocysts less than 4 months of age to inoculate host HCT-8 cell monolayers. Oocysts were diluted in a standard 5- or 10-fold multiple dilution format, levels of infection and clustering were determined, and the most probable number (MPN) of infectious oocysts in the stock suspension was calculated. The MPN was compared to the initial oocyst inoculum to determine the level of correlation. For oocysts less than 30 days of age, the correlation coefficient (r) was 0.9726 (0.9306 to 0.9893; n = 20). A two-tailed P value (alpha = 0.05) indicated that P was less than 0.0001. This strong correlation suggests that the MPN can be used to effectively enumerate infectious oocysts in a cell culture system. Age affected the degree of oocyst infectivity. Oocyst infectivity was tested by the focus detection method (FDM)-MPN assay and in BALB/c mice before and after treatment with pulsed white light (PureBrite). The FDM-MPN assay and animal infectivity assays both demonstrated more than a 4-log10 inactivation. Municipal water systems and a host of other water testing organizations could utilize the FDM-MPN assay for routine survival and disinfection studies.

Cryptosporidium parvum was first associated with human illness in the 1970s (5, 6). The organism's environmentally stable oocysts are transmitted by the fecal-oral route, often through contaminated drinking water. Outbreaks have occurred as a result of oocyst resistance to disinfectants commonly used in drinking water treatment. Within the last 12 years, 19 documented outbreaks in unrelated geographic areas have occurred in the United States, Canada, United Kingdom, and Japan, affecting more than an estimated 427,000 individuals (4, 31). Consequently, this organism has become a major concern to public health and the drinking water industry.

Cell culture technology has developed into a tool that can be used to study C. parvum in an environment most similar to the situation in vivo without using animal models (13, 29, 30, 32-34, 36). The research described here uses the previously described focus detection method (FDM) set up in a multiple dilution format (27-29). The autoinfectious nature of the Cryptosporidium life cycle was observed to begin 12 h after incubation, and by 48 h, an average ratio of 17.9 foci to each oocyst was observed (29). Presumably, nonsynchronous excystation and infection were occurring during this complex life cycle, and the parasites produced clusters of reproductive stages in the host monolayer after an extended incubation period. At first, this confounded the ability to directly count the number of infected foci in the host cells. Rather than counting the individual foci, a presence or absence observation was made after 48 h of incubation and was used to detect one infectious oocyst or fewer (based upon dilution) per well in vitro.

The most-probable-number (MPN) method of enumerating microorganisms (specifically, bacteria) was adopted as a method to quantify the number of infectious oocysts. The method entails making a dilution series, plating in replicate, incubating the samples for an adequate period, and scoring the presence of growth. In the cell culture system, cluster presence (and absence) can easily be detected with the FDM and used to determine the number of viable organisms (in this case, infectious oocysts) per milliliter in a sample by the MPN method. Tables made by Halvorson and Ziegler (15) have traditionally been consulted for the MPNs; however, computer programs have enabled researchers to obtain a more accurate number and allow nonstandard volumes and dilutions to be used (11, 21). The objective of this study was to standardize and test statistically the reproducibility of the MPN method by using the cell culture FDM for infectious oocysts.

MATERIALS AND METHODS

C. parvum oocysts less than 4 months old were obtained from Pleasant Hill Farm, Iowa. The original strain was obtained from a naturally infected cow. The oocysts were perpetuated in newborn calves, purified by an ethyl ether and Percoll-sucrose gradient procedure (according to a protocol that can be obtained from Pleasant Hill Farm), and stored at 4°C in phosphate-buffered saline (PBS) with antibiotics (penicillin and streptomycin). Upon delivery, each oocyst lot number was counted with a hemacytometer and concentrations were recorded. All lot numbers were in concentrations greater than 105 oocysts per ml. Aliquots used for cell culture infection were processed as described below. Cell culture infectivity testing was performed simultaneously with in vitro excystation and vital dye exclusion to compare the viabilities determined by the three assays (3, 25).

Eight oocyst lots (from different calves) were evaluated, with three to eight replicate experiments/lot, in a total of 34 experiments in order to determine the variability of lot numbers and the FDM-MPN method. Each experiment was set up in a multiple dilution format, and 10- or 5-fold dilutions with three to six replicates per dilution were used.

In vitro cell culture. The cell culture system has previously been described (27, 28). A brief description (with changes) is presented here. Human ileocecal adenocarcinoma cells (HCT-8 cells) were maintained in 75-cm2 tissue culture flasks and passed every 2 or 3 days. LabTech II (Nalgene Nunc, Naperville, IL) eight-well chamber slides were seeded with 5 × 104 cells per well and grown to approximately 60 to 80% confluency in a 5% CO2 atmosphere at 37°C for 24 to 72 h. HCT-8 maintenance medium contained RPMI 1640 supplemented with 5% fetal bovine serum (Atlanta Biologicals), 2% 1 M HEPES, and 1% 200 mM l-glutamine. No antibiotics were used.

Oocyst preparation. Once the monolayer was established, C. parvum oocysts were prepared. Oocysts were pretreated with 10% (vol/vol) bleach (5.2% sodium hypochlorite) (4°C) and washed by centrifugation after 5 to 8 min in the bleach solution. Stock microscopic oocyst counts of the oocyst suspensions were determined by a direct hemacytometer count, and 10- or 5-fold oocyst dilutions in
growth medium were prepared in sterile polycarbonate microcentrifuge tubes. Each dilution was pipetted onto cell monolayers in three or six replicate wells. The entire volume of each diluted oocyst suspension was inoculated. The cultures were incubated in a 5% CO₂ atmosphere at 37°C for 48 h (the 90-min washing step previously described [29] was excluded because no deleterious effects to the host cells from the toxicity caused by the oocysts excysting have been noted in the diluted oocyst suspensions). Controls assayed concurrently for each experiment included uninoculated cell monolayers in two wells on each slide as negative controls. Uninoculated wells also served to determine if the chamber slides were leaking or if any transfer between wells occurred.

Antibody labeling with the FDM. Well chamber slides were fixed with 100% methanol and labeled by an indirect antibody procedure as previously described [29]. Rat anti-sporozoite polyclonal antibody, generously provided by Steve Upson (Kansas State University, Manhattan, Kans.) and Waterborne, Inc. (New Orleans, La.), was used as the primary antibody, while anti-rat fluorescein isothiocyanate-conjugated antibody (Sigma) was used as the secondary antibody.

Enumeration. Fixed and labeled slides were examined by epifluorescence microscopy at ×200 magnification, and each well was scored as either positive or negative. Positive infection was determined by the presence of visible sporozoite invasion that produced an infection focus, and clustering, a result of secondary infection, was determined when numerous life stages were present. When cells appeared positive for infection, a confirmation of Cryptosporidium life stages (meronts and microgametes, etc.) was made at ×400 magnification and/or under oil at ×1,000 magnification. Both invasion and replication in vitro were required to score the well as positive. When both invasion and replication were not present, the well was scored as negative.

The MPN of infectious oocysts was determined by using the information collection rule (ICR) general purpose Most Probable Number Calculator, version 1.00 (20). The ICR program was modified from a previously published ICR program specifically for the application of total cultivable viruses (MPNv) for samples processed under the ICR (19). The program enables the user to enter the number of replicates and dilutions, volumes used, and dilution factors to generate an MPN with confidence intervals. The program is easy to use and is preferred to referring to tables. The MPN data were statistically analyzed with the GraphPad Prism software (San Diego, Calif.) and Excel.

Percent infectivity was determined by the following formula: percent infectivity = (MPN per ml/microscopic oocyst count per milliliter of the stock) × 100.

In vitro viability assays. Excystation and vital dye exclusion methods were also used to evaluate viability (3, 25). The inclusion and/or exclusion of the fluorogenic vital dyes DAPI (4′,6-diamidino-2-phenylindole) and propidium iodide (PI) within oocysts was examined microscopically with an Olympus model BH2 microscope equipped with a UV filter block (350-nm excitation and 450-nm emission) for DAPI and a green filter block (500-nm excitation and 630-nm emission) for PI. Proportions of ruptured (ghost), PI (+), DAPI (+)-PI (−), and DAPI (+)-PI (+) oocysts were quantified (3).

Excystation procedures included the use of excysting fluid (0.5% trypsin and 1.5% sodium taurocholate in tissue culture with PBS) to incubate oocysts for 90 min at 37°C (25). Sporozoites, intact oocysts, and oocyst shells were counted under ×400 magnification by Nomarski differential interference contrast microscopy. A total of 300 oocysts were counted per experiment. As an additional experimental control, the sporozoite yield viability was determined by dividing the sporozoite yield by 4 (the theoretical number of sporozoites per oocyst) and multiplying the resulting factor by 100.

Animal infectivity. Oocyst infectivity was evaluated by modified neonatal BALB/c mouse infectivity assay in addition to cell culture infectivity (1). Inactivation experiments were performed by using a broad-spectrum pulsed white light water treatment device (PureBrite, San Diego, Calif.) (35). Pregnant BALB/c mice were shipped overnight from Jackson Labs (Bar Harbor, Maine) to Michael Arrowood (Centers for Disease Control and Prevention, Atlanta, Ga.). Oocyst samples were packed on ice and shipped by overnight express to Michael Ar-
each dilution series exhibited little or no variability among the MPNs. In many cases, the same MPN was produced. For example, lot 4 oocysts (18 days of age) were bleach treated, and three separate sets of dilutions were made from the same bleached stock. The three experiments showed $1.19 \times 10^4$, $1.01 \times 10^4$, and $1.01 \times 10^3$ MPNs of infectious oocysts/ml and the proportions of infective oocysts were 47.59, 40.47, and 40.47%, respectively.

Several lots were assayed repeatedly as the oocysts aged (at 4°C in PBS). The MPNs produced revealed differences among each experiment, and subtle changes in infectivity were observed. To determine the source of variability, all experiments were plotted to show the relationship of each lot number with the percent infectious oocysts and age (Fig. 1). Figure 1A shows the percent infection plotted against age for all lots tested in the cell culture system. Figure 1B shows the regression comparison of the percent infectious oocysts with age for four oocyst lot numbers. Both lot number and age were factors associated with the variation observed with infectivity.

Figure 2 shows the relationship between the hemacytometer count (number of oocysts/ml) and the MPN/ml for oocysts less than or greater than 30 days of age. Correlation coefficients ($r$) were 0.9726 and 0.9241 for oocysts less than and greater than 30 days of age, respectively ($P < 0.0001$). The $r$ value for all experiments (all ages) combined was 0.4081 (95% confidence interval [CI], 0.0811 to 0.6558; $P = 0.0166$). Stock oocyst counts for oocysts less than 30 days of age were statistically equivalent to the MPNs calculated; however, the relationship between the oocyst inoculum and the MPN diverged as the oocysts aged.

The log$_{10}$ of the ratio between the stock oocyst count/milliliter and the MPN/milliliter were compared with age for each experiment (Fig. 3). As the oocyst age increased, the ratio of the inoculum/milliliter to the MPN/milliliter generally became more negative. Oocysts beyond 30 days of age had a difference in ratios $>-1.0$ log$_{10}$ in 92% (12 in 13) and $>-1.5$ log$_{10}$ in 46% (6 in 13) of the experiments. Oocysts 30 days of age or less had only 19% (3 in 21) of ratios that were $>-1.0$ log$_{10}$ different.

Table 2 compares the average percent infectivity and viability of various oocyst lot numbers. The average infectivity (by the FDM) and viability (by excystation, sporozoite yield, and DAPI-PI) varied considerably for each lot number. Correlation values ($r$) of the FDM with excystation, sporozoite yield, and DAPI-PI were 0.1340, $-0.2249$, and $-0.8295$, respectively, which demonstrates a significant difference between oocyst in vitro infectivity and surrogate viability. Correlation ($r$) between excystation and DAPI-PI was 0.0335 for this study.

Figure 4 presents the comparison between the infectivity and viability assays for oocysts less than or greater than 30 days of age ($n = 8$). A statistical difference in infectivity by the FDM and viability by excystation was observed for oocysts less than 30 days of age when compared to that of oocysts greater than 30 days of age. No statistical difference in viability by DAPI-PI was observed for oocysts less than or greater than 30 days of age. It has generally been accepted that oocysts less than 90 days of age and stored in PBS at 4°C are viable as long as they exhibit ample excystation and vital dye exclusion. This study found that the percent infectious oocysts (by the FDM) decreased as the oocysts aged after 30 days and infectivity was significantly different from viability by the surrogate microscopic viability assays.

Table 3 shows the comparison between cell culture and BALB/c mouse infectivity. For the same lot of oocysts, the result of the FMD-MPN method was slightly higher than the mouse MPN of infectious oocysts (15 to 9.6 times greater) in untreated water samples. Cell culture and animal infectivity both showed similar reductions in the MPN/milliliter for infectious oocysts after treatment by pulsed white light (broad spectrum) (PureBrite) (16). After treatment, MPNs were 31.00 and 7.41 compared to $<5.64$ and 20.17/ml for cell culture and

![FIG. 2. Linear regression comparison of the number of oocysts/milliliter inoculated and the MPN/milliliter calculated for oocysts less than or greater than 30 days of age.](image)

![FIG. 3. Comparison of the log of the ratio of stock oocyst count/milliliter and MPN/milliliter with age for all experiments.](image)

| TABLE 2. Infectivity and viability of various oocyst lot numbers |
|-------------------|-------------------|-------------------|-------------------|
| Oocyst lot no. | FDM | Excystation | Sporozoite yield | DAPI-PI |
| 1 | 37.7 (7.3) | 58.2 (3.9) | 65.1 (3.5) | 66.7 (12.1) |
| 2 | 63.5 (7.8) | 71.2 (7.6) | 58.5 (12.0) | 76.0 (9.3) |
| 3 | 8.5 (11.8) | 33.2 (7.8) | 76.5 (17.6) | 82.6 (4.5) |
| 4 | 42.7 (4.1) | ND | ND | ND |
| 5 | 4.7 (12.2) | 42.6 (16.6) | 48.0 (18.5) | 86.7 (8.8) |
| 6 | 8.7 (4.7) | 58.9 (5.7) | ND | ND |
| 7 | 3.1 (7.0) | 89.0 (2.7) | 84.0 (1.0) | 92.3 (3.1) |
| 8 | 5.4 (2.7) | ND | ND | ND |

* The percent infectivity and viability is the geometric mean of the individual percents calculated for each lot number ($n = 2$ to 8). ND, no data recorded.
Antibody staining of infected cells provides an advantage over molecular detection methods such as PCR and in situ hybridization because the clustering of foci and specific life stages can be seen. This is a clear indication that reproduction has occurred. Sporozoites have been observed to excyst, initiate invasion of the host cells, but then not go further past that stage even after 48 h of incubation (16, 28). In this study, oocysts after pulsed broad-spectrum light treatment were observed to excyst and initiate infection (minimal) and they did form single foci. These oocysts could not proceed past the sporozoite invasion stage after 48 h of incubation, and thus, no infection clusters were observed. Molecular assays may detect the DNA or RNA produced in the invasion stage and may overestimate the total number of infectious oocysts.

The percent viable and infectious oocysts for each lot number varied considerably, indicating that there was significant lot-to-lot variability. Several freshly isolated oocyst lot numbers were less than 10% infectious. Oocyst processing to purify the oocysts from feces may contribute to the variability (e.g., defecating feces during process using ethyl ether, etc.). Future work will evaluate different oocyst purification assays to determine the effects on the oocysts.

The infectivity of all oocyst lot numbers decreased as the oocysts aged. Belosevic et al. showed similar findings using vital dyes for oocysts isolated from different calves and stored at 4°C (2). This variation in the viability and infectivity of different oocyst lot numbers and ages suggests that the oocysts are more affected by aging than previously thought, and each lot number should be analyzed before use for survival and disinfection studies.

This is the first study to report, quantitatively, oocyst viability over time by an infectivity assay. Several previous studies on oocyst infectivity or inactivation did not account for oocyst age prior to use. This study suggests that 30 days may be the maximum storage time, prior to experimentation, for maintaining a high proportion of infectious oocysts and that age may be an important quality control issue.

In our laboratory, the FDM-MPN method has now been used in disinfection studies for the water and food industry to determine oocyst inactivation after exposure to various water activities, pulsed white light, UV light, chlorine dioxide, ozonation and ultrahigh pressure. By using the MPNs, log_{10} and percent reductions can be calculated (16, 28, 30). The assay has been used for environmental (secondary sewage effluent [27]) and turbid water samples, which can be a problem in other viability assays such as microscopy, PCR, and in situ hybridization due to inhibition by the particulates. Future work will test the limits of particulates in the cell culture system.

Infectious oocysts can be detected repeatedly at very low concentrations (as low as 1 oocyst per ml determined by the FDM), indicating the high-level sensitivity of this assay. For each experiment, the oocysts were diluted to less than 1 oocyst per ml. Clusters of foci were detected at these highly diluted

### TABLE 3. Cell culture FDM and BALB/c mouse infectivity by *C. parvum* oocysts

<table>
<thead>
<tr>
<th>Sample</th>
<th>Cell culture (MPN/ml (CL)) for:</th>
<th>BALB/c mouse (MPN/ml) for:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Influent</td>
<td>$3.11 \times 10^6$ ($6.89 \times 10^5$–$9.03 \times 10^6$)</td>
<td>$1.97 \times 10^5$ ($6.69 \times 10^5$–$5.12 \times 10^5$)</td>
</tr>
<tr>
<td></td>
<td>$2.40 \times 10^5$ ($4.80 \times 10^5$–$9.70 \times 10^5$)</td>
<td>$2.49 \times 10^5$ ($8.95 \times 10^5$–$5.78 \times 10^5$)</td>
</tr>
<tr>
<td>Effluent*</td>
<td>$31.09$ ($6.90$–$90.30$)</td>
<td>$&lt;5.64^a$ ($0.92$–$19.05$)</td>
</tr>
<tr>
<td></td>
<td>$7.41$ ($1.70$–$21.50$)</td>
<td>$20.17$ ($3.94$–$42.72$)</td>
</tr>
</tbody>
</table>

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*a After treatment by pulsed white light (broad spectrum) (PureBrite).

*b No infection detected.
subsamples and were obvious against the dark host monolayer. The polyclonal antibody labels all the life stages present, thus enabling detection at low concentrations. Several monoclonal antibodies were tested; however, these did not have the same specificity or ability to pick up all the life stages as the polyclonal ones (data not presented). Since the FDM-MPN assay is both cell culture and antibody based, it may be very specific to infection only by *C. parvum*; however, the experiments to test specificity with all isolates and *Cryptosporidium* species have not been performed. *Cryptosporidium muris* was tested, and life stages past the initial trophozoite stage and clumping were not detected. Future efforts will focus on natural isolates from humans and animals, including various genotypes (24).

Vital dye and excystation methods have been routinely used for survival and disinfection experiments (9, 17, 18, 25). Frequently, researchers use animal models to validate the surrogate in vitro viability assays. When the FDM-MPN method was an excellent and reproducible assay for quantifying oocyst infectivity from different lot numbers may be a contributor of oocysts (8, 10, 22) and in humans was 132 oocysts (7). A high degree of scatter in the maximum likelihood analysis of the cell culture data was shown. The factors contributing to this are unknown; however, the variability of oocysts from different lot numbers may be a contributing factor.

In conclusion, the FDM-MPN method has been shown to be an excellent and reproducible assay for quantifying oocyst infectivity in vitro. Oocyst age is an important issue that must be addressed before proceeding with disinfection projects. It is also notable that the cell culture assay is more sensitive than surrogate in vitro viability assays. When the FDM-MPN method was compared to animal models, a similar level of infectivity was determined, suggesting that the FDM can be used as an alternative to animal assays.

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REFERENCES


