Newly defined conditions for the in vitro cultivation and cryopreservation of Dientamoeba fragilis: new techniques set to fast track molecular studies on this organism

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SUMMARY

Dientamoeba fragilis is a pathogen of the human gastrointestinal tract that is a common cause of diarrhoea. A paucity of knowledge on the in vitro cultivation and cryopreservation of Dientamoeba has meant that few studies have been conducted to investigate its biology. The objective of this study was to define, for the first time, in vitro culture conditions able to support the long-term in vitro growth of Dientamoeba. Also, we aimed to define a suitable method for cryopreserving viable Dientamoeba trophozoites. A modified BD medium, TYGM-9, Loeffler’s slope medium, Robinson’s medium, Medium 199, Trichosel and a Tritrichomonas fetus medium were compared, using cell counts, for their ability to support the growth of D. fragilis at various temperatures and atmospheric conditions. Loeffler’s slope medium supported significantly better growth compared to other media. A temperature of 42 °C and a microaerophilic atmosphere were also optimum for Dientamoeba growth. To our knowledge, this is the first study to describe and compare different culture media and conditions for the growth of clinical isolates of D. fragilis. This new technology will aid the development of diagnostics for dientamoebiasis as well as facilitate large-scale sequencing projects that will fast track molecular studies on D. fragilis.

Key words: Dientamoeba fragilis, in vitro, culture, cultivation, cryopreservation, culture media, optimization.

INTRODUCTION

Dientamoeba fragilis is a protozoan pathogen of the human gastrointestinal tract first described in the scientific literature by Jepps and Dobell (1918). Dientamoeba was originally classified as an amoeba (subphylum Sarcodina (Jepps and Dobell, 1918)) though a later study utilizing electron microscopy confirmed its relationship with trichomonads (Camp et al. 1974). Although Dientamoeba was described almost a century ago, comparatively little information exists with regards to its life cycle, genetics and proteome. Given the recent recognition of D. fragilis as a significant human pathogen by numerous authors (Dickinson et al. 2002; Girginkardesler et al. 2003; Johnson et al. 2004, Stark et al. 2005b, Lagace-Wiens et al. 2006, Stark et al. 2006, 2007, 2009, 2010), more research with regards to this organism is warranted. The ability to maintain clinical isolates of Dientamoeba in culture for extended periods of time at high cell densities and, the ability to cryogenically preserve viable Dientamoeba trophozoites, would facilitate further study and analysis of this emerging pathogen.

Previous authors have reported on the xenic cultivation of D. fragilis (Lamy, 1960; Robinson, 1968; Robinson and Ng, 1968; Sawangjaroen et al. 1993; Clark and Diamond, 2002; Windsor et al. 2003). However, no axenic cultivation system has been successfully developed for D. fragilis. Many xenic media are able to support the growth of D. fragilis. Dobell was the first to isolate and grow D. fragilis using a biphasic, xenic medium consisting of an inspissated horse serum slope overlaid with a liquid phase consisting of egg whites diluted in Ringer’s solution supplemented with rice starch (Dobell, 1940). Other biphasic media able to support the growth of D. fragilis include Cleveland and Collier’s medium (a Loeffler’s slope with a liquid overlay (Cleveland and Collier, 1930)), modified Boeck and Drbohlav’s (BD) medium (LE medium) and Robinson’s medium (Clark and Diamond, 2002; Windsor et al. 2003). Some monophasic media can also support the growth of D. fragilis. The American Type Culture Collection (ATCC) recommends a monophasic TYGM-9 broth ((ATCC medium 1171) as used in experiments by Chan et al. (1994)) for the growth of D. fragilis strain ATCC 30948 (strain Bi/Pa)—though isolates of D. fragilis are no longer available from the ATCC (at the time of writing). Balamuth’s medium is another monophasic liquid medium which can support the growth of D. fragilis.
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D. fragilis

also studied by others suggest that the addition of 2:5% glucose allows for

D. fragilis

at a final concentration of 5% DMSO. To our knowledge, no

studies exist that compare methods for the cryo-

preservation and in vitro cultivation of D. fragilis.

Also, the ability of commercial media such as M199,

Trichosel and Trichomonas foetus medium to support the growth of D. fragilis has never been explored. Furthermore, the effect of different atmospheric conditions and temperatures on Dientamoeba growth has not been fully investigated either. In

order to address this paucity of knowledge a modified BD medium, TYGM-9, a Loeffler’s slope medium

(Trichosel), M199, Trichosel and a commercial Trichomonas foetus medium were investigated for their ability to support the growth of D. fragilis. Experiments were also performed to evaluate the growth of D. fragilis under anaerobic, microaerophilic and aerophilic conditions and, to explore its growth at a range of different temperatures. Cell counts were performed on cultures to determine the most suitable in vitro growth conditions for D. fragilis cultivation. Also, variations of the cryopreservation methods described by Sawangjaroen et al. (1993) and Dwyer and Honigberg (1971) were compared for their ability to preserve viable D. fragilis trophozoites.

MATERIALS AND METHODS

Modified BD medium

One hundred ml of the liquid overlay was prepared by mixing 90 ml of PBS (Sigma P4417-100TAB), 9 ml of heat-inactivated horse serum (Sigma H1138-500 ml) and 1 ml of a 20% (w/v) bacteriological peptone solution (Oxoid, LP0037). Solid egg slopes were prepared as described previously (Sawangjaroen et al. 1993), and approximately 2 mg of rice starch (Sigma, S-7260) was placed in the bottom of each slope. Each slope was then overlaid with 5 ml of the liquid overlay.

Loeffler’s medium

Loeffler’s serum slopes were prepared to contain heat-inactivated horse serum (700 ml/L), glucose (2.5 g/L) and nutrient broth No.2 (6.25 g/L) in distilled water. Five ml of this were poured into a 14 ml vol. McCartney bottle, sloped and inspissated in an 80 °C drying oven until slopes solidified. Approximately 2 mg of rice starch was placed into the bottom of each slope. The slopes were then overlaid with 5 ml of PBS. It should be noted that during inspissation, if Loeffler’s slopes are left for too long at 80 °C, Dientamoeba will not grow in the resulting medium. This is probably due to the dam-

aging of some essential cofactor/s within the serum. During inspissation, slopes should be observed frequently and allowed to cool immediately after they solidify.

TYGM-9 broth

TYGM-9 broth (formulation: 2.8 g K₂HPO₄, 0.4 g KH₂PO₄, 2.0 g casein digest, 1.0 g yeast extract, 7.5 g NaCl, 2.0 g gastric mucin all dissolved in 970 ml of distilled water) was prepared and poured into Pyrex 99447 15.5 ml vol. glass culture tubes in 10 ml volumes. Each tube was supplemented with approxi-

mately 2 mg of rice starch.

Trichosel

Trichosel was purchased from BD Diagnostic sys-

tems (cat. No. 298323) and prepared as per the manufacturer’s instructions. Ten ml volumes were dispensed into Pyrex 99447 15.5 ml vol. glass culture tubes. Each tube was supplemented with approxi-

mately 2 mg of rice starch.

Robinson’s medium

This medium was prepared as described previously by Clark and Diamond (2002) with the following modification: the inclusion of approximately 2 mg of rice starch.

Trichomonas foetus medium

Three ml vials of T. foetus medium were commer-
cially obtained (Micromedia, Cat. No. 3274) and supplemented with approximately 2 mg of rice starch.

Medium 199

Medium 199 (Sigma Aldrich, Cat. No. M3769) was

prepared according to the manufacturer’s instruc-
tions. Heat-inactivated horse serum (Sigma Aldrich, Cat. No. H1138-500 ml) was added to Medium 199
at a concentration of 10%. Ten ml volumes were dispensed into Pyrex 99447 15.5 ml vol. glass culture tubes and approximately 2 mg of rice starch was added to each tube.

**Source of D. fragilis trophozoites**

Four *Dientamoeba* strains were used in these studies. They were isolated from the stools of symptomatic patients known to be infected with *D. fragilis*. Patients were initially diagnosed with dientamoebiasis based on the observation of *Dientamoeba* trophozoites in iron-haematoxylin stained faecal smears. Oil-immersion microscopy performed on sodium acetate acetic acid formalin (SAF) fixed, iron-haematoxylin stained faecal smears is routinely carried out in the Department of Microbiology at St Vincent’s Hospital, Sydney, for investigation of gastrointestinal complaints. Stool specimens fixed in SAF were stained with a modified iron-haematoxylin stain (Fronine, Australia) according to the manufacturer’s recommendations. All stained smears were examined by oil-immersion microscopy (1000× magnification). Approximately 250 fields of view were examined on each slide. Definitive diagnosis was based on the characteristic morphology of *D. fragilis* in the permanently stained smears (trophozoites measuring 5–15 μm in diameter, with 1 or 2 nuclei each with a large fragmented central karyosome without peripheral chromatin (Fig. 1A)).

Patients with a microscopy-confirmed *Dientamoeba* infection were requested to submit a fresh stool specimen for culture prior to their treatment. Immediately upon receipt of the fresh, unpreserved stool specimen a portion was cultured. Approximately 10 mg of the unpreserved stool specimen was placed directly into a bottle of modified BD medium. Cultures were incubated at 37 °C and, after 48 h, a drop of sediment from each culture was examined at 400× magnification under phase-contrast microscopy for the presence of *Dientamoeba* trophozoites. When *Dientamoeba* trophozoites were observed, a portion of the culture sediment was fixed in SAF (1:1) and stained with a modified iron-haematoxylin stain (Fronine, Australia) and examined by oil-immersion microscopy to confirm the isolation of *D. fragilis*. Another portion of the sediment was kept for DNA extraction followed by PCR.

**Comparison of different media**

*Dientamoeba fragilis* trophozoites were grown in Loeffler’s slope medium (at 37 °C under microaerophilic conditions) and quantitated by cell counts using Kova slides (Hycor Biomedical Inc). For all cell counts, the medium vessels containing *Dientamoeba* trophozoites to be counted were very gently shaken and inverted several times to achieve an even cell suspension. In order to completely break up the sediment on the bottom of the medium vessels it was sometimes necessary to agitate the cultures by very gently pipetting up and down with a transfer pipette. Once an even cell suspension was achieved, counts were performed using Kova slides. A small drop of the cell suspension was placed into the counting chamber of a Kova slide and raw cell counts were performed.

Trophozoites were added to 2 vials of each of the different types of media (modified BD medium, Loeffler’s slope medium, TYGM-9 broth, Trichosel, Robinson’s medium, *Trichomonas foetus* medium and Medium 199) to a final concentration of 50 trophozoites/μl. This was achieved by diluting the trophozoites into the culture media to be used and/or concentrating by centrifugation at 500 g for 5 min where required. Each culture was then incubated at 37 °C under microaerophilic conditions. Cell counts were made each day as described above, until no more trophozoites could be observed (or until 9 days passed—whichever occurred first). Immediately after counts were completed each day, cultures were reincubated as described above. Cell counts were plotted graphically in order to visualize the growth patterns of *Dientamoeba* in each medium. This was repeated for the 4 isolates of *D. fragilis*. 

![Fig. 1. A trophozoite of *Dientamoeba fragilis* in a fixed, iron-haematoxylin stained faecal smear viewed under oil-immersion microscopy (A) and, live trophozoites from modified BD cultures viewed under phase contrast microscopy (B, C and D). The typical fragmented nuclear structure of *Dientamoeba* in stained smears is observable in the binucleate trophozoite shown in inset A. Trophozoites in culture can exhibit different morphologies including a spherical, granular, vacuolated form (B) and motile forms with visible pseudopodia (C, D). In modified BD cultures, rice starch will often be observed in the trophozoite cytosol as highly refractive granules (C, D), though not always (B). Scale bars represent 10 μm.](image-url)
Effect of atmospheric composition on in vitro growth

This set of experiments was performed using Loeffler’s slope medium and 4 clinical isolates of D. fragilis. Dientamoeba trophozoite cultures were grown in Loeffler’s medium. These trophozoites were inoculated into 6 fresh vials of Loeffler’s medium to a total concentration of 108 trophozoites/μl. This was performed by concentration of trophozoites via centrifugation (500 g for 5 min), and/or dilution in PBS where required, and quantifying by cell counts. Two vials were each grown under aerophilic, microaerophilic and anaerobic conditions. These steps were repeated for each of the 4 clinical isolates of D. fragilis.

Aerophilic conditions

In order to expose D. fragilis cultures to as close to aerophilic conditions as could be achieved, 2 culture vials from each isolate were left in a 37 °C incubator with the caps completely loosened. Cell densities were counted each day until trophozoites were no longer observed.

Microaerophilic conditions

Microaerophilic conditions were achieved using an Anoxomat machine (Mart Microbiology B.V.). The lids of all culture vessels were completely loosened and cultures were placed in an appropriate Anoxomat jar and connected to the Anoxomat machine according to the manufacturer’s instructions. Gases were introduced into the sealed Anoxomat Jars to the following concentrations: 6% O₂, 7-2% CO₂, 3-6% H₂, 83-3% N₂. Jars containing culture vials were placed at 37 °C and cell counts were made each day until trophozoites were no longer observed. After counting each day, cultures were placed in Anoxomat jars and microaerophilic gases were re-introduced. The jar was re-incubated at 37 °C.

Anaerobic conditions

Anaerobic conditions were achieved using an Anoxomat machine. Gases were introduced into the sealed Anoxomat jars to the following concentrations: 0-2% O₂, 9-9% CO₂, 5% H₂, 84-9% N₂. Each jar containing cultures was placed at 37 °C and cell counts were made every day until trophozoites were no longer observed. After all counts were made, cultures were placed in Anoxomat jars and anaerobic gases were reintroduced to the jar as previously described. The jar was re-incubated at 37 °C.

Effect of temperature on in vitro growth

Dientamoeba trophozoites were added to Loeffler’s medium at a total concentration of 185 cells/μl. Two of these quantified cultures were grown at each of the different temperatures, including room temperature, 30 °C, 37 °C, 40 °C and 42 °C (a total of 10 bottles of medium). All cultures were grown under microaerophilic conditions as already described. This was repeated for each of the 4 clinical isolates of D. fragilis. Cultures were then incubated under microaerophilic conditions and cell counts were made each day until trophozoites were no longer observed. Numerical data obtained for counting experiments relating to various media formulations, atmospheric conditions and temperatures are shown in Table 4.

DNA extraction

DNA was extracted from a 2 mg portion of the sediment of modified BD cultures containing D. fragilis trophozoites. Extractions were performed using a QIAamp™ DNA stool minikit (Qiagen, Hilden, Germany) using the stool extraction protocol described by the manufacturer. This was performed for each of the 4 D. fragilis isolates obtained in this study.

PCR and sequencing

A conventional PCR assay was employed as described previously (Stark et al. 2005a) for the amplification of the SSU rDNA gene of D. fragilis from DNA extracted from modified BD culture sediments. Following PCR, all PCR products were subjected to agarose gel electrophoresis in a 1% gel. PCR products were visualized under UV light and excised from the gel using a fine scalpel. The PCR products were then extracted from the gel slice using a QIAquick gel extraction kit (QIAGEN) according to the manufacturer’s instructions. Samples for sequencing were prepared to contain 10 pmoles of the primers DF 400 or DF 1250 (Stark et al. 2005a) and approximately 25 ng of PCR product in a total volume of 16μl. Sequencing was performed at least once in both forward and reverse directions. All sequencing was performed by the service provider SUPAMAC. Sequences obtained from all isolates obtained in this study were aligned with other SSU rDNA genes of D. fragilis available on GenBank (Accession nos U37461 and AY730405) using a Clustal W program.

Identification of microbial flora in Dientamoeba cultures

After several passages in modified BD medium, under microaerophilic conditions at 37 °C, bacterial cultures were performed on the sediments of culture vessels from each isolate of Dientamoeba. Cultures were inoculated onto Columbia horse blood agar...
plates (Oxoid Cat. No. PP2001) and incubated under aerobic conditions at 37 °C, and anaerobic media plates (Oxoid cat. No. PP2039) incubated under anaerobic conditions at 37 °C. After 24–48 h incubation the microbial flora was identified using standard phenotypic laboratory techniques.

**Cryopreservation method 1**

This method is a modification of that described by Sawangjaroen et al. (1993) with the addition of glucose as suggested by Dwyer and Honigberg (1971) for a concentration of 5% DMSO. A solution of 5% (w/v) D-glucose (Chem-Supply Pty Ltd) in single-strength PBS was prepared. Solutions of 5%, 7.5%, 10%, 12.5% and 15% (v/v) DMSO (Sigma-Aldrich Cat. No. 154938-100 ml) in single-strength PBS were also prepared. *Dientamoeba fragilis* trophozoites from each isolate were grown in 4 vials of Loeffler’s slope medium. The contents of these vials were pooled by removing the entire liquid portion (including sediments) and placing into 10 ml vol. Falcon tubes. Tubes were centrifuged at 500 g for 5 min. The supernatant was discarded. One ml of 5% glucose (w/v) in PBS was added to the pellet and tubes were inverted several times to obtain an even cell suspension. Cell counts were then performed as described previously. The quantified cell suspensions were then diluted with 5% glucose (w/v) in PBS to a final concentration of 180 trophozoites/μl.

Five hundred μl of each of the 5 different DMSO solutions were placed into a single Microbank™ vial (Pro-Lab Diagnostics, USA) for cryofreezing (Microbank™ vials for this experiment were first prepared by pouring off the medium and beads within the tubes followed by rinsing several times with PBS and blotting dry on a paper towel). Five hundred μl of quantified cell suspension (180 trophozoites/μl) from a single isolate of *D. fragilis* were then added to each of the 5 tubes containing different concentrations of DMSO. This resulted in solutions of 2.5%, 3.75%, 5%, 6.25% and 7.5% DMSO (v/v) in a 2.5% glucose (w/v) solution in PBS and a final trophozoite concentration of 90 trophozoites/μl. This was repeated for each of 4 isolates of *D. fragilis*. These 20 tubes were then placed overnight in a −80 °C freezer. The following morning the tubes were then removed from the −80 °C freezer and placed in a liquid nitrogen freezer (CHART™, Model MVE TEC 3000) for 4 days. On the morning of the fifth day, each tube was removed from liquid nitrogen and placed directly in a 37 °C water bath to thaw. Once thawed, the total contents of each tube were used to inoculate vials of fresh Loeffler’s medium. Each of these vials was then incubated at 37 °C for 48 h. After this time, a drop of sediment from each of these cultures was examined at 400× magnification under phase-contrast microscopy for the presence of motile trophozoites of *D. fragilis* (Fig. 1C and D). When motile trophozoites were observed, cell counts were performed as described previously.

**Cryopreservation method 2**

This method is a modification of that described by Dwyer and Honigberg (1971). Solutions of 2%, 2.25%, 2.5%, 2.75%, 3% and 3.25% (v/v) DMSO in single-strength PBS were prepared. *Dientamoeba fragilis* trophozoites from each isolate were grown in 4 vials of Loeffler’s medium. The 4 culture vials from each isolate were pooled in 10 ml vol. Falcon tubes. Tubes were centrifuged at 500 g for 5 min and the supernatant was discarded. One ml of PBS was added to the pellet and tubes were inverted several times to obtain an even cell suspension. Cell counts were then performed as described previously. The quantified cell suspensions were then diluted in PBS to a final concentration of 106 trophozoites/μl.

Five hundred μl of each of the 6 different DMSO solutions were placed into a single Microbank™ vial (Microbank™ vials were first prepared by removal of the beads and media as described above). Five hundred μl of quantified cell suspension (106 trophozoites/μl) from a single isolate of *D. fragilis* was then added to each of the 6 tubes containing different concentrations of DMSO. This resulted in solutions of 1%, 1.25%, 1.25%, 1.375%, 1.5% and 1.625% (v/v) DMSO in PBS and a final trophozoite concentration of 53 trophozoites/μl. This was repeated for each *D. fragilis* isolate. Each Microbank™ vial was then frozen as described above. Each vial was thawed on the morning of day 5 post-freezing as described previously and inoculated into fresh Loeffler’s medium. After 48 h incubation, a drop of sediment from each of these cultures was examined at 400× magnification under phase-contrast microscopy for the presence of motile trophozoites of *D. fragilis*. Where motile trophozoites were observed cell counts were performed as described previously.

**Statistical analysis of growth data from all experiments**

In order to statistically compare the differences in cell density between the growth plots obtained in all experiments, cumulative cell counts were calculated for each day and analysed using a paired t-test.

**RESULTS**

**Comparison of all media**

Figure 2 shows the average growth pattern of all 4 clinical isolates of *D. fragilis* in Loeffler’s medium, modified BD medium and Robinsons’ medium at 37 °C under microaerophilic conditions. *Trichomonas foetus* medium and Trichosel, failed to support...
the growth of *D. fragilis* entirely and so were excluded from further study. Loeffler’s medium supported significantly higher cell densities of all clinical isolates when compared to all other media (Table 1). While the average growth of all *Dientamoeba* isolates in Robinson’s medium was slightly higher than that of modified BD medium (T value 0.552), the cell densities achieved for these media were not significantly different from each other (P value 0.595). TYGM-9 and M199 supported very poor growth of 3 of 4 isolates. Isolate 4 demonstrated visibly better growth in TYGM-9 and in particular, M199 when compared to Isolates 1, 2 and 3 and, as such the results for these two media were excluded from Fig. 2. Figure 3 shows the average growth of Isolate 4 in TYGM-9 and M199 compared to the average growth of Isolates 1, 2 and 3. This figure demonstrates the obvious phenotypic difference between the growth of Isolate 4 compared to Isolates 1, 2 and 3 in TYGM-9 and M199.

Figure 4 compares the growth of Isolate 4 in MBD medium, Loeffler’s slope medium, Robinsons’ medium, TYGM-9 and M199. This figure shows that M199 is a suitable medium for the growth of Isolate 4 only. Despite this observation, Loeffler’s medium followed by the modified BD medium and Robinson’s medium support (on average) the most efficient growth of *D. fragilis*.

### Table 1. P and T values obtained using a paired t-test to compare the growth of *Dientamoeba* under different conditions

<table>
<thead>
<tr>
<th>Paired variables</th>
<th>T value</th>
<th>P value</th>
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</thead>
<tbody>
<tr>
<td><strong>Comparison of different atmospheric conditions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average growth of all isolates under microaerophilic conditions vs aerobic conditions</td>
<td>6.085</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Average growth of all isolates under microaerophilic conditions vs anaerobic conditions</td>
<td>7.197</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Average growth of all isolates under aerobic conditions vs anaerobic conditions</td>
<td>-5.337</td>
<td>0.001</td>
</tr>
<tr>
<td><strong>Comparison of different temperatures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average growth of all isolates at 37°C vs 40°C</td>
<td>4.814</td>
<td>0.002</td>
</tr>
<tr>
<td>Average growth of all isolates at 37°C vs 42°C</td>
<td>-5.077</td>
<td>0.003</td>
</tr>
<tr>
<td>Average growth of all isolates at 40°C vs 42°C</td>
<td>-5.931</td>
<td>0.001</td>
</tr>
<tr>
<td><strong>Comparison of different media</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average growth of all isolates in Loeffler’s medium vs modified BD medium</td>
<td>3.381</td>
<td>0.008</td>
</tr>
<tr>
<td>Average growth of all isolates in Loeffler’s medium vs Robinson’s medium</td>
<td>4.903</td>
<td>0.001</td>
</tr>
<tr>
<td>Average growth of all isolates in Robinson’s medium vs modified BD medium</td>
<td>0.552</td>
<td>0.595</td>
</tr>
</tbody>
</table>

#### Effect of atmospheric composition on in vitro growth

The results shown in Fig. 5 indicate that concentrations of oxygen found in the terrestrial atmosphere are not favourable for the growth of *D. fragilis*. Concentrations of oxygen equal to or below 6% are conducive for the growth of *D. fragilis*. Interestingly, *D. fragilis* appears to prefer oxygen concentrations higher than those observable in anaerobic environments, demonstrating significantly better growth under microaerophilic conditions when compared to anaerobic conditions (P value 0.000).

#### Effect of temperature on in vitro growth

The data in Fig. 6 show high *Dientamoeba* trophozoite numbers in cultures grown between 37 and 42 °C. Temperatures between that of ambient room temperature and 30 °C failed to support any growth of the organism. Growth of *Dientamoeba* at 42 °C was significantly better when compared to the growth of *Dientamoeba* at lower temperatures (Table 1). Growth at 37 °C was found to be significantly better than growth at 40 °C although the calculated standard error was quite large for values obtained at 40 °C (Fig. 6).
PCR and sequencing

Despite the observable differences between the growth of some isolates of *D. fragilis* in certain media, no genetic differences were observed in a section of the SSU rDNA gene sequenced for all 4 isolates in this study. Clinical isolates from this study are of a similar SSU rDNA genotype to the more common genotype 1 (GenBank Accession no. AY730405), previously identified in patients from Sydney, Australia and dissimilar to SSU rDNA genotype 2 (the Bi/pa genotype (GenBank Accession no. U37461)).

Identification of microbial flora in *Dientamoeba* cultures

Several species of bacteria were identified in the cultures from each isolate of *D. fragilis* (Table 2). No eukaryotic species other than *D. fragilis* was identified in these cultures. The predominant bacterial species in cultures from all isolates was *Escherichia coli*.
Despite this, comparatively little information is available about the biology of this organism including genome and proteome studies. Also, the most effective drug treatment for dientamoebiasis is yet to be defined, although progress has been made (Stark et al. 2010). The ability to cryogenically preserve this organism and to culture it in vitro at high cell densities will facilitate the development of new diagnostics, the characterization of novel antigens, the study of this organism’s life cycle and biology and the development of in vitro methods to assess drug treatments.

Four of 7 different types of media were able to support the growth of all 4 clinical isolates of *D. fragilis*. These 4 media were TYGM-9 broth, Robinson’s medium, modified BD medium and Loeffler’s medium. *Trichomonas foetus* medium and Trichosel failed to support the growth of *D. fragilis*. On average, the growth of *D. fragilis* was significantly better in Loeffler’s slope medium compared to Robinson’s medium and modified BD medium. This is in contrast to the results of Sawangjaroen et al. (1993) who found that the modified BD medium was the only medium which supported the good long-term growth of

**Table 2. Microbial flora identified in the cultures of each isolate of *Dientamoeba fragilis***

<table>
<thead>
<tr>
<th>Isolate name</th>
<th>Bacterial* flora identified</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isolate 1</td>
<td><em>Escherichia coli</em>, <em>Citrobacter</em> spp., <em>Proteus mirabilis</em>, <em>Bacteroides fragilis</em> gp, <em>Veillonella</em> spp.</td>
</tr>
<tr>
<td>Isolate 2</td>
<td><em>Escherichia coli</em>, <em>Serratia</em> spp., <em>Proteus mirabilis</em>, <em>Enterococcus faecium</em>, <em>Prevotella intermedia</em>, <em>Bacteroides ureolytic</em> gp.</td>
</tr>
<tr>
<td>Isolate 4</td>
<td><em>Escherichia coli</em>, <em>Enterococcus faecium</em>, <em>Citrobacter</em> spp., <em>Proteus mirabilis</em>, <em>Fusobacterium</em> spp., <em>Bacteroides fragilis</em> gp.</td>
</tr>
</tbody>
</table>

* No eukaryotic organisms other than *D. fragilis* were identified in cultures.

**Table 3. The presence (+) or absence (−) of in vitro growth for each isolate of *Dientamoeba fragilis* after cryopreservation using different concentrations of DMSO followed by cultivation in Loeffler’s medium.**

<table>
<thead>
<tr>
<th>Isolate name</th>
<th>Final DMSO concentration (v/v) in PBS solution</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>1%</td>
</tr>
<tr>
<td>Isolate 1</td>
<td>−−−−−</td>
</tr>
<tr>
<td>Isolate 2</td>
<td>−−−−−</td>
</tr>
<tr>
<td>Isolate 3</td>
<td>−−−−−</td>
</tr>
<tr>
<td>Isolate 4</td>
<td>−−−−−</td>
</tr>
</tbody>
</table>

**DISCUSSION**

*Dientamoeba fragilis* is a protozoan parasite of the gastrointestinal tract of humans which has recently gained recognition as a significant human pathogen (Johnson et al. 2004; Stark et al. 2006, 2009, 2010). Despite this, comparatively little information is available about the biology of this organism including genome and proteome studies. Also, the most effective drug treatment for dientamoebiasis is yet to be defined, although progress has been made (Stark et al. 2010). The ability to cryogenically preserve this organism and to culture it in vitro at high cell densities will facilitate the development of new diagnostics, the characterization of novel antigens, the study of this organism’s life cycle and biology and the development of in vitro methods to assess drug treatments.

Four of 7 different types of media were able to support the growth of all 4 clinical isolates of *D. fragilis*. These 4 media were TYGM-9 broth, Robinson’s medium, modified BD medium and Loeffler’s medium. *Trichomonas foetus* medium and Trichosel failed to support the growth of *D. fragilis*. On average, the growth of *D. fragilis* was significantly better in Loeffler’s slope medium compared to Robinson’s medium and modified BD medium. This is in contrast to the results of Sawangjaroen et al. (1993) who found that the modified BD medium was the only medium which supported the good long-term growth of

![Growth plots comparing the growth of *Dientamoeba fragilis* at various temperatures](image)

Fig. 6. Growth plots comparing the growth of *Dientamoeba fragilis* at various temperatures [room temperature (●●●●), 30°C (---), 37°C (--), 40°C (— —) and 42°C (■■■■)] in Loeffler’s slope medium.初始细胞在各温度下的细胞数量。
Dientamoeba when they compared its growth in monophasic TYSGM-9 (TYGM-9 medium with the addition of serum), Cleveland and Collier’s medium and a modified BD medium. Sawangjaroen et al. (1993) found that the modified BD medium was the only medium which supported the good long-term growth of the organism (Sawangjaroen et al. 1993).

Dientamoeba was found to grow well under both microaerophilic (6% O₂, 7·2% CO₂, 3·6% H₂, 83·3% N₂) and anaerobic (0·2% O₂, 9·9% CO₂, 5% H₂, 84·9% N₂) conditions. However, *D. fragilis* does not appear to grow as well in the presence of atmospheric levels of oxygen. The trophozoite densities obtained under microaerophilic conditions were significantly greater when compared to the trophozoite densities obtained under both aerobic and anaerobic conditions. The comparatively poor growth of *Dientamoeba* at higher oxygen concentrations reflects the fragile nature of *D. fragilis* once it is passed from the body and comes into contact with the atmosphere. The results suggest that *Dientamoeba* is best grown at 42 °C. However, it appears that temperatures between 37 °C and 42 °C are acceptable for *D. fragilis* growth. Significantly higher cell densities are obtained at 42 °C compared to 40 °C and 37 °C. Dobell previously reported optimum growth of *D. fragilis* at 41 °C (Dobell, 1940) and the results of the current study support Dobell’s observations. The growth of *D. fragilis* at 42 °C is an interesting observation as one would expect an optimum *D. fragilis* growth temperature closer to the core body temperature of humans (≈37 °C) rather than that of birds (≈42 °C). All isolates demonstrated similar temperature affinities although it was often observed that the trophozoite densities from 2 culture vials of the same isolate, grown at the same temperature were quite different from each other. This phenomenon was particularly apparent when isolates were grown at 40 °C and is the reason for the large error bars observable in Fig. 6. Trophozoite densities obtained at 37 °C were significantly greater than those obtained at 40 °C. This seems paradoxical as the highest cell densities were achieved at 42 °C and one would expect greater cell densities at temperatures approaching 42 °C. All temperature experiments were repeated to exclude human error, though with a similar paradoxical outcome. The reason behind this phenomenon is unclear although we recommend that for cultivation of *Dientamoeba*, multiple culture vessels of a given isolate should always be maintained, and that subcultures should be made from the ‘healthiest’ of these cultures only in order to select protozoa that grow vigorously. However, it should also be taken into account that selecting certain subpopulations of trophozoites introduces bias and has the potential to reduce the genetic variability of the original clinical isolate. As such, this kind of selection will reduce the similarity between the original wild-type organisms and the cultured organisms. These points should be considered when undertaking this kind of selection.

### Table 4. Numeric data obtained from *Dientamoeba fragilis* counting experiments used to construct Figures 2, 5 and 6

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* S.D., Standard deviation.
During the course of these experiments, it also became apparent that 1 of the clinical isolates of *D. fragilis* was quite different from the other 3 in terms of its ability to grow comparatively well in 2 monophasic liquid media (TYGM-9 broth and M199) while all other isolates could not. The growth of this isolate (Isolate 4) in Robinson’s medium, modified BD medium and Loeffler’s medium followed a very similar growth pattern to all other isolates and so raw counts made for Isolate 4 were included when calculating the average cell densities shown in Fig. 1. However, due to the visibly different patterns of growth observed for Isolate 4 in TYGM-9 broth and M-199, a separate figure was created in order to compare the growth of Isolate 4 to Isolates 1–3 in these monophasic media. Fig. 3 clearly demonstrates the ability of Isolate 4 to grow well in TYGM-9 broth and M-199 when compared to all other isolates.

Sequencing of the small subunit gene of all 4 isolates demonstrated that they were all greater than 99% similar in the sequenced region of the SSU rDNA. All 4 isolates demonstrated almost identical sequences to 7 other sequences derived from a previous PCR study of *D. fragilis* performed in Sydney (GenBank Accession no. MY730405). Undoubtedly, some genetic differences must exist between Isolate 4 and all other isolates and future work may investigate these differences. Interestingly while M199 failed to support 3 isolates of *D. fragilis*, it seems that M199 was able to support high cell densities for Isolate 4 at 4 days of growth (higher cell densities than were achieved for this isolate in any other media). While this may be the case we do not consider M199 a good medium for the cultivation of *D. fragilis* for the obvious reason that growth in this media is isolate/strain dependent.

Given the obvious phenotypic differences observable between Isolate 4 and all other isolates, and the similarity between the SSU rDNA sequences obtained from all isolates, we conclude that the SSU rDNA gene is well conserved, even amongst very different isolates of *D. fragilis*, suggesting that the SSU rDNA gene is a poor genetic marker for distinguishing amongst different strains of *D. fragilis*. This point is supported by the findings of other studies (Stark et al. 2005a; Bart et al. 2008). Only 2 SSU rDNA genotypes of *D. fragilis* are known to exist (genotypes 1 and 2) and all isolates in this study are similar to SSU rDNA genotype 1, which is the first (and only) SSU rDNA genotype identified in Sydney to date (Stark et al. 2005a). Genotype 1 was also found to be more common than genotype 2 (the Bi/pa genotype) in previous studies (Johnson and Clark, 2000; Windsor et al. 2004).

In order to loosely explore the possibility that *D. fragilis* may exhibit better growth in the presence of certain bacterial species, the bacterial flora accompanying the cultures of each isolate of *Dientamoeba* was defined. As mentioned previously, the growth of all *Dientamoeba* isolates was similar in Loeffler’s medium, Robinson’s medium and modified BD medium. This was in spite of the slight variation in the bacterial flora accompanying each isolate in culture. *Escherichia coli* was the predominant bacterial species within all cultures. As such, we can conclude that *D. fragilis* will grow happily with a support flora consisting mostly of *E. coli*. We also suggest that slight variations in the species of prokaryotic support flora present within *D. fragilis* cultures, are unlikely to exhibit any significant effect on growth. It is possible that *D. fragilis* could exhibit better (or worse) growth if the predominant bacterial species within a culture is changed, although no conclusions regarding this hypothesis can be drawn from the present study. Differences in growth patterns exhibited by Isolate 4 compared to other isolates of *Dientamoeba* are unlikely to be attributable to the bacterial flora. Once again, *E. coli* was the predominant bacterium within cultures of Isolate 4 and nearly all other bacterial species present within Isolate 4 cultures were also present in cultures of other isolates. However, the presence of *Fusobacterium* spp. was a feature unique to Isolate 4. Despite this, *Fusobacterium* spp. were not a major constituent of the flora within cultures of Isolate 4 and we feel that the differences observable for Isolate 4 are inherent within the isolate itself rather than the support flora.

It is possible that the temperature dependence observed for all *Dientamoeba* isolates could be directly related to the effect of changes in temperature on the microbial support flora. It is possible that an increase in temperature may reduce the doubling time for bacteria and have little/less effect on *Dientamoeba*, thus increasing the availability of certain nutrients to *Dientamoeba*. It is also possible that an increase in temperature directly increases the doubling time of *Dientamoeba*. However, it should also be noted that some species of enteric bacteria (such as *E. faecium*) have an optimal growth temperature of 42–45 °C (Zanoni et al. 1993). Clearly, these hypotheses require further investigation. The development of cultures with defined bacterial species’ (monoxenic or dixenic cultures) or preferably, axenic cultures, would be useful to explore the effects of temperature on *Dientamoeba* growth.

No monoxenic, dixenic or axenic cultures of *D. fragilis* exist to our knowledge (at the time of writing). According to studies performed by Brug (1938) and Jacobs (1953), *Dientamoeba* trophozoites are heavily dependent on the presence of live bacteria as a food source, with dead bacteria being insufficient for *Dientamoeba* growth (Jacobs, 1953). Brug (1938) observed that if bacteria become scarce in cultures, *Dientamoeba* trophozoites eventually die off. This suggests that an axenic culture of *Dientamoeba* would be difficult to achieve (if this is achievable at all). However, the creation of monoxenic or dixenic
cultures of *Dientamoeba* seems more obtainable. According to Jacobs (1953), a monoxenic culture of *Dientamoeba* with *Clostridium perfringens* was obtained through the use of various antibiotics. Jacobs (1953) also noted that *Dientamoeba* was quite tolerant to fairly high concentrations of the drugs penicillin, streptomycin and sulfadiazine. As such, the use of various antibiotics may represent a simple means of obtaining monoxenic or dixenic cultures of *Dientamoeba*. Monoxenic or dixenic cultures of *Dientamoeba* would prove useful in genetic and proteomic studies as the defined support flora could then be used as a negative control. Experiments which test the *in vitro* susceptibility of *Dientamoeba* to various antimicrobial drugs would also benefit from the existence of such cultures. Experiments which aim to achieve monoxenic, dixenic and/or axenic cultures of *Dientamoeba* are currently underway.

A final concentration of 1.375% DMSO (as described in cryopreservation method 2) resulted in excellent recovery and growth of trophozoites when compared to other concentrations of DMSO used. This is in agreement with the study by Dwyer and Honigberg (1971) who found the highest percentage recoveries at a final DMSO concentration of 1.375%. Cryopreservation method 1 failed to preserve viable *D. fragilis* trophozoites entirely. Interestingly, Isolate 4 was the only isolate that could not be cryogenically preserved using method 2. We speculate that this is probably the result of genetic differences between isolates. Future efforts will be made to identify these genetic differences and to develop a method for cryopreservation of Isolate 4. Dwyer and Honigberg (1971) utilized a liquid nitrogen freezer with controlled cooling capabilities in their *D. fragilis* cryopreservation experiments. Phillips et al. (1984) noted higher recoveries of *Giardia* trophozoites when they were frozen using a liquid nitrogen freezer with controlled cooling capabilities compared to simply placing trophozoites into a −70°C freezer. This equipment was not available to us although we believe that the cryopreservation of *D. fragilis* trophozoites may be improved using a controlled cooling freezer. Final DMSO concentrations >1.5% failed to preserve viable *D. fragilis* trophozoites entirely. This is in contrast to the findings of Sawangjaroen *et al.* (1993), who reported the successful cryopreservation of viable *D. fragilis* trophozoites in a final concentration of 7.5% DMSO.

To conclude, our studies indicate that the best conditions for the growth of *D. fragilis* in the media tested in this study are as follows; microaerophilic conditions, at 42°C in Loeffler's slope medium (a modified Cleveland and Collier's medium). However, *D. fragilis* will also grow quite well in modified BD medium or Robinson's medium under microaerophilic conditions at 37°C and 40°C. It should also be noted that while M199 was able to support good growth of 1 isolate of *Dientamoeba* only, Loefflers' medium, modified BD medium and Robinson's medium were able to support good growth of all isolates in these experiments. Therefore these 3 media are recommended for those attempting to cultivate clinical isolates of *D. fragilis* from fresh stool specimens. Furthermore, effective cryopreservation of *D. fragilis* trophozoites is best achieved using method 2 of cryopreservation described herein at a final DMSO concentration of 1.375% (v/v) in single-strength PBS.

**ACKNOWLEDGMENTS**

We acknowledge the help of the staff at St Vincent's Hospital Microbiology Department in the collection and processing of stool samples positive for *D. fragilis*. We especially acknowledge the assistance of Tamalee Roberts in typing the bacterial species in our *D. fragilis* cultures.

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**REFERENCES**


