The effects of storage conditions on viability of Clostridium difficile vegetative cells and spores and toxin activity in human faeces

J Freeman, M H Wilcox

Aims: Clostridium difficile is a common nosocomial pathogen and as such diagnostic and research methods may necessitate storage of faecal specimens for long periods, followed by subsequent re-examination. This study investigated the effects of storage conditions upon the viability of this organism and its toxin.

Methods: Three genotypically distinct strains of C difficile (two clinical isolates including the UK epidemic strain, and an environmental isolate) were grown anaerobically at 37°C for 72 hours in a pool of five faecal emulsions. Aliquots of each emulsion were stored at either −20°C (frozen) or 4°C (refrigerated). Emulsions were assayed for viable cells, spores, and cytotoxin titre before storage and at days 1, 3, 5, 7, 14, 28, and 56. An aliquot of each emulsion was also removed, assayed, and replaced in storage at each time point to investigate the effects of multiple freezing/refrigeration/thawing.

Results: Neither storage temperature nor multiple cycles of refrigeration/freeze-thawing had minimal effects upon the viability of C difficile or its spores. Storage at 4°C has no discernible effect on C difficile cytotoxin. However, storage at −20°C has a detrimental effect upon C difficile cytotoxin, and multiple cycles of freezing and thawing may further adversely affect toxin titres.

Conclusion: Storage temperature and multiple cycles of freezing/thawing had minimal effects upon the viability of C difficile or its spores. Storage at 4°C has no discernible effect on C difficile cytotoxin. However, storage at −20°C has a detrimental effect upon C difficile cytotoxin, and multiple cycles of freezing and thawing may further adversely affect toxin titres.

Methods

Faecal emulsions were prepared by pooling five faecal samples from healthy elderly individuals and diluting 1/20 (wt/vol) in pre-reduced phosphate buffered saline (PBS; pH 7.4). Emulsions were pre-reduced in an anaerobic cabinet (Don Whitley Scientific, Bradford, UK) for 24 hours before inoculation.

Pre-reduced faecal emulsions were each inoculated with 50 µl of a 24-hour Schaedler’s anaerobic broth culture containing either C difficile strains p24, B32 (clinical toxigenic isolates, polymerase chain reaction (PCR) ribotype 1 and 78, respectively), or E16 (environmental toxigenic isolate, PCR ribotype 44). Faecal emulsions were incubated in an anaerobic environment for 72 hours, after which they were removed. Each faecal emulsion was divided into 16 aliquots of 200 µl in Eppendorf tubes and two aliquots of 700 µl. Eight aliquots were placed in a −20°C freezer, along with a 700 µl aliquot. The remaining aliquots were placed in a 4°C refrigerator. On days 0, 1, 3, 5, 7, 14, 28, and 56 a single 200 µl aliquot was removed from each storage condition and assayed for vegetative cells, spores, and cytotoxin (using the Vero cell culture assay), as described below. These aliquots were then discarded. At the same time points, the 700 µl aliquots were removed from storage and assayed in the same way. These aliquots were then replaced in their respective storage conditions.

Clostridium difficile viable count

Each faecal emulsion was vortexed and serially diluted 10 fold in PBS. Three Brazier’s cycloserine-cefoxitin egg yolk agar plates containing 5 mg/litre lysozyme without egg yolk...
A10 was removed and serially diluted (1/10) to a dilution of 16 000 × g A 100 Clostridium difficile incubated and counted as described above. Inoculated on to CCEYL plates, which were anaerobically culture toxin assays were incubated at 37°C, in the presence of (Pro-lab Diagnostics, South Wirral, Cheshire, UK)). Cell toxin (prepared according to the manufacturer’s instructions procedure of Borriello was used to kill the vegetative cells. Each faecal emulsion was vortexed and a 200 l aliquot of each faecal emulsion was centrifuged at 6 000 × g for 10 minutes. After centrifugation, the supernatant was removed and serially diluted (1/10) to a dilution of 10^6 in PBS. Vero cell monolayers were inoculated with these l aliquot neat supernatant and 20 l un inoculated on to CCEYL plates, which were anaerobically incubated and counted as described above.

Clostridium difficile spore assay

Each faecal emulsion was vortexed and a 200 l aliquot removed to an Eppendorf tube. The bench alcohol shock procedure of Borriello et al was used to kill the vegetative cells. Briefly, an equal volume of absolute ethanol was added to the tube and the mixture was vortexed. The faecal emulsion/ethanol mix was left at room temperature for one hour. The mixture was then serially diluted (1/10) in PBS and 20 l inoculated on to CCEYL plates, which were anaerobically incubated and counted as described above.

Clostridium difficile toxin assay

A 100 l aliquot of each faecal emulsion was centrifuged at 16 000 × g for 10 minutes. After centrifugation, the supernatant was removed and serially diluted (1/10) to a dilution of 10^6 in PBS. Vero cell monolayers were inoculated with these serial 10 fold dilutions. Specific confirmation of the presence of C difficile toxin was achieved by inoculating a duplicate well with 20 l neat supernatant and 20 l Clostridium sordelli antitoxin (prepared according to the manufacturer’s instructions (Pro-lab Diagnostics, South Wirral, Cheshire, UK)). Cell culture toxin assays were incubated at 37°C, in the presence of 5% CO₂ and examined at 24 and 48 hours under an inverted microscope. A positive reaction was indicated by cell rounding. The toxin titre was defined as the dilution adhering to the following criteria:

- Cell rounding of approximately > 50% of the monolayer.
- Cell rounding of approximately all of the monolayer in the preceding well (containing less dilute sample).
- No evidence of cell rounding in the subsequent well (containing more dilute sample).

The assay was considered valid if cell rounding was completely prevented by the presence of C sordelli antitoxin.

RESULTS

Viable counts and spore counts of all strains of C difficile (p24, B32, and E16) remained approximately constant throughout the course of the experiment (56 days). Neither storage temperature nor multiple cycles of refrigeration/freezing and thawing adversely affected the viability of C difficile vegetative cells or its spores.

Single and multiple exposures of samples to 4°C had little effect upon C difficile toxin titre. Toxin titres, although fluctuating slightly during the course of the experiment, generally remained within one or two logarithmic concentrations of the initial titre.

A detrimental effect was seen upon toxin titre at −20°C, in both singly and multiply frozen and thawed samples. Strain p24 decreased from an initial titre of 10⁶ toxin units to 10⁴ toxin units (single freeze/thaw cycle) and 10³ toxin units (multiple freeze/thaw cycle) by day 56 of the study. Similar decreases were seen in strains B32 and E16. The most dramatic decrease was seen when strain B32 was exposed to multiple freeze/thaw cycles. Toxin titres decreased from 10⁶ toxin units to undetectable values by day 3 of the experiments. Figures 1–3 show the decreases in toxin titres relative to the initial titre. Analysis of variance was performed upon each time point. Error bars represent the minimum significant difference relating to data obtained for that time point; comparisons are not made between time points. Toxin titres of multiply frozen and thawed faeces became significantly lower than those stored at 4°C (p < 0.01) by day 5 of the experiment in two of the three strains, and in all strains by day 28. Toxin titres of singly frozen and thawed faeces became significantly lower than those stored at 4°C (p < 0.01) by day 56 of the experiment in two of the three strains.

DISCUSSION

Our study showed that C difficile remains viable both as vegetative cells and spores, with little quantitative variation, after single or multiple exposure to either 4°C or −20°C for at least 56 days. Bowman and Riley reported recovery of C difficile in faecal specimens for up to 10 days, depending on the isolation medium used, and Weese et al reported recovery of C difficile in 25 of 26 C difficile positive equine faecal samples after
anaerobic storage at 4°C for 30 days. The above studies described the survival of *C. difficile* in specimens as either “positive” or “negative” with no reference to numbers or the presence or absence of *C. difficile* spores. Our present study used lysozyme supplemented media to optimise *C. difficile* recovery. Culture showed little fluctuation in the numbers of either total viable counts or spore counts, indicating that the organism does not sporulate in response to storage at either 4°C or −20°C. It is interesting to note that Weese et al found that samples stored aerobically at 4°C yielded decreasing counts with time. This was not seen in our study, although all specimens were stored aerobically, and some were periodically exposed to air during multiple freezing/refrigeration and thawing processes. This discrepancy in observations may have arisen as a result of differences in methodology. In our study, faeces were emulsified in PBS, providing a buffered environment for the organism after its removal from an anaerobic environment. It is possible that exposure to air may influence the pH of faeces and this may affect *C. difficile*. Continued fermentation of unabsorbed sugars to acids by bacteria outside the buffered environment of the large bowel may lower faecal pH. Some studies have reported a decrease in faecal pH associated with decreased numbers of *C. difficile*. In addition, Futter and Richardson investigated the effect of the pH value of solid medium on the recovery of spores of *C. histolyticum*, *C. septicum*, *C. perfringens*, and *C. wheelei* (*perfringens*), and found that the optimum pH for spore recovery lay between 7.5 and 8.8.

Richardson investigated the effect of the pH value of solid broth cultures stored at 70°C, 20°C, and 4°C, but no titrations were performed to detect reductions in toxin titre. It is possible that the buffering of faecal specimens would be a feasible method of preserving *C. difficile* viability during transport or storage. This requires further investigation.

Although the viability of *C. difficile* is unaffected by storage at either 4°C or −20°C, the activity of cytotoxin was considerably reduced when samples were stored at −20°C. This is in agreement with previous observations that cytotoxin degrades at −20°C. Conversely, Weese et al reported detectable toxin in broth cultures stored at −70°C, −20°C, and 4°C, but no titrations were performed to detect reductions in toxin titre. In addition, toxin was detected using an enzyme linked immunosorbent assay system, whereas our study used a cell culture cytotoxicity assay. Therefore, it is possible that storage at −20°C results in the loss of cytotoxic, but not immunological, activity. Bowman and Riley found that 1.7 log reduction in average toxin titre after two days of storage at 25°C and a similar, although less pronounced reduction (1.4 log), after two days of storage at 5°C. However, we found no such decrease after storage at 4°C, when samples were subjected to either single or multiple cycles of freeze/refrigeration and thawing. We cannot account for this discrepancy.

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The differences in the reduction of toxin titres between *C. difficile* strain p24 (PCR ribotype 1) and strain B32 (PCR ribotype 78) were interesting. *Clostridium difficile* PCR ribotype 1 is the UK epidemic strain, and accounts for 60% of UK *C. difficile* isolates. Virulence assays performed in our laboratory have shown several differences between *C. difficile* strain p24 and other strains, namely: (1) *C. difficile* strain p24 produced significantly more spores than other non-prevalent strains; (2) *C. difficile* strain p24 produced significantly more spores than other strains in response to ampicillin treatment (including *C. difficile* strains B32 and E16) (Freeman J, Wilcox MH). Does antibiotic exposure affect sporation in an epidemic *Clostridium difficile* strain? Poster presented at 40th Interscience Conference on Antimicrobial Agents and Chemotherapy; and (3) *C. difficile* strain p24 germinated to a significantly greater degree than strains B32 and E16 (Freeman and Wilcox, unpublished data, 2000). Thus, the cytotoxin of the UK epidemic strain may be more robust than those of other *C. difficile* strains, a hypothesis that warrants further study.

It is also possible that the structure of the toxin is damaged by ice crystals, formed during freezing, which may disrupt protein structure. This may explain why repeated freezing and thawing caused the greatest reductions in toxin titres. If this is the case, the fact that we emulsified the faecal samples in PBS may have contributed to toxin degradation because more water molecules would have been present in samples. This could be alleviated by the use of a cytoprotective storage medium. Samples stored as solid faeces may provide a more protective environment for the toxin.

On the basis of our results, we recommend that specimens should be stored at 4°C instead of −20°C to minimise toxin degradation. The high recovery rates of *C. difficile* in PBS emulsified faeces stored at either 4°C or −20°C throughout the duration of the experiment may indicate that buffering of faeces allows *C. difficile* to survive for prolonged periods of time. This warrants further investigation, but may provide a suitable means of long term storage of faecal specimens containing *C. difficile*.

**Authors’ affiliations**

Freeman, M.H. Wilcox, Department of Microbiology, University of Leeds and The General Infirmary, Leeds LS2 9JT, UK

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