Bovine immunoglobulin concentrate—Clostridium difficile retains C difficile toxin neutralising activity after passage through the human stomach and small intestine

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Abstract
Background—Bovine immunoglobulin concentrate (BIC)—Clostridium difficile is prepared from the colostrum of cows immunised against C difficile toxins and contains high concentrations of neutralising IgG antitoxin.

Aims—To determine the proportion of BIC-C difficile which survives passage through the human stomach and small intestine.

Methods—Six volunteers with an end ileostomy took 5 g of BIC-C difficile containing 2.1 g of bovine IgG on four occasions: alone, with an antacid, during treatment with omeprazole, and within enteric coated capsules.

Results—When BIC-C difficile was taken alone, a mean (SEM) of 1033 (232) mg of bovine IgG was recovered in the ileal fluid representing 49% of the total ingested dose. Bovine IgG recovery was not significantly increased by antacid (636 (129) mg) or omeprazole (1052 (268) mg). The enteric capsules frequently remained intact or only partially opened in the ileal effluent and free bovine IgG levels were low in this treatment group (89 (101) mg). Bovine IgG recovery was higher in volunteers with shorter (less than two hours) mouth to ileum transit times (68% versus 36%, p<0.05). Specific bovine IgG against C difficile toxin A was detected in ileal fluid following oral BIC. Toxin neutralising activity was also present and correlated closely with bovine IgG levels (r=0.95, p<0.001).

Conclusion—BIC-C difficile resists digestion in the human upper gastrointestinal tract and specific anti-C difficile toxin A binding and neutralising activity was retained. Passive oral immunotherapy with anti-C difficile BIC may be a useful non-antibiotic approach to the prevention and treatment of C difficile antibiotic associated diarrhoea and colitis.

Keywords: pseudomembranous colitis; toxin; diarrhoea; IgG; immunotherapy; antibiotic; Clostridium difficile

Clostridium difficile infection is a common cause of diarrhoea in hospital and nursing home patients. Antibiotic therapy alters the normal colonic microflora and allows opportunistic infection by C difficile. Infection is more likely to occur in hospital than in outpatients receiving antibiotics because hospital patients and their environment frequently harbour C difficile and its spores. C difficile diarrhoea and colitis are caused by two protein exotoxins produced by pathogenic strains of the organism. Toxin A is a 308 kDa protein with cytotoxic, enterotoxic, and proinflammatory effects. Toxin B, a 280 kDa protein, is a more potent cytotoxin than toxin A, but is not enterotoxic for rodent intestine. However, toxin B does cause injury to human colonic mucosa in vitro. Toxin A and toxin B show considerable sequence homology and both share the same intracellular mechanism of cytotoxicity. They catalyse the monoglucoolyslation of small GTP binding rho proteins leading to disruption of the cytoskeleton, cell rounding, and cell death. C difficile colitis is currently treated by antimicrobial therapy using metronidazole or vancomycin. These agents are effective in the short term. However, when treatment is discontinued up to 20% of patients suffer a recurrence of diarrhoea and, in some instances, multiple recurrences occur requiring repeated courses of therapy. Treatment with metronidazole or vancomycin may predispose to recurrent C difficile diarrhoea by causing further disruption of the colonic microflora, a primary event in the pathogenesis of C difficile colitis. The use of antimicrobial agents to treat antibiotic associated diarrhoea has also led to concern regarding nosocomial bacterial antibiotic resistance; in particular vancomycin resistance in enterococcal species. These issues have led to a search for an effective, non-antibiotic agent, for prevention or treatment of nosocomial C difficile diarrhoea.

We previously reported the production of a hyperimmune bovine colostral antibody preparation against C difficile. Holstein cows are immunised against C difficile and its toxins during the gestation period and later an immunoglobulin concentrate is prepared from their colostrum. The bovine immunoglobulin concentrate—Clostridium difficile (BIC-C difficile) contains high levels of IgG against C difficile toxins, neutralises the biological effects of C difficile toxins in vitro and, in animal studies,
proteins against *C difficile* toxin induced enteroocolitis. BIC-*C difficile* is designed for use as an oral passive immunotherapy against *C difficile* and its toxins. To be therapeutically active, oral BIC-*C difficile* must exert toxin neutralising activity in the human colon. In an earlier study of healthy volunteers, we found detectable levels of bovine IgG as well as *C difficile* toxin neutralising activity in the stool after oral BIC-*C difficile* administration. In that study protection from acidic gastric secretions and short gastrointestinal transit times both appeared to increase faecal bovine IgG concentrations.

The survival of intact bovine IgG in the faeces is relevant to the therapeutic use of BIC-*C difficile*. However, of equal or even greater importance, is the delivery of intact bovine IgG to the human colon. In this study, volunteers with an end ileostomy but otherwise apparently normal gastrointestinal function took BIC-*C difficile* orally and their ileal fluid effluent was collected for analysis. The main study aims were to measure accurately the amount of bovine IgG surviving passage to the distal small intestine in humans and to determine whether specific *C difficile* toxin binding and neutralising activity was preserved. Additional aims were to determine whether reduced exposure to acidic gastric secretions or individual variations in upper gastrointestinal transit times would significantly alter bovine IgG survival.

**Methods**

**PREPARATION OF BOVINE IMMUNOGLOBULIN CONCENTRATE-CLOSTRIDIUM DIFFICILE**

BIC-*C difficile* was prepared from the colostral milk of Holstein cows, as previously described. For this study, cows were immunised with a toxoid of purified *Clostridium difficile* toxin A. The resulting BIC-*C difficile* contained 42 mg of bovine IgG per 100 mg of powdered concentrate. This preparation of BIC-*C difficile* showed high concentrations of anti-*C difficile* toxin A activity as demonstrated by: (1) ELISA; (2) inhibition of toxin A binding to its brush border membrane receptor; (3) neutralisation of the cytotoxic effects of toxin A in the tissue culture cytotoxicity assay; and (4) inhibition of the enterotoxic effects of toxin A in the rodent ileal loop assay.

**HUMAN STUDY PROTOCOL**

A single site, open, phase I study was performed in volunteers with an end ileostomy to examine the safety and bioavailability of a single oral dose of BIC-*C difficile*. The study protocol was approved by the Beth Israel Deaconess Medical Centre Committee on Clinical Investigations. An outline of the study protocol is illustrated in figure 1.

**Study subjects**

Six volunteers (four men, two women; mean age 49 years, range 35–69) with a surgically created end ileostomy were enrolled into the study. In all cases, end ileostomy was performed following colectomy. The indications for colectomy were ulcerative colitis in five patients and indeterminate colitis in one. At enrolment none of the patients had a history or any clinical evidence of upper gastrointestinal tract disease apart from ileostomy. All volunteers reported stable ileostomy function during the six month period prior to study entry.

Exclusion criteria included: a history of other gastrointestinal diseases or surgery which might be expected to affect normal gastrointestinal function, a history of clinically significant cow’s milk protein allergy or other allergy to milk products, lactose intolerance, evidence of ongoing systemic or infectious disease, or use of medication that might be expected to affect normal gastrointestinal function. All volunteers received monetary compensation for study participation.

**Administration of BIC-*C difficile**

BIC-*C difficile* 5 g, containing 2.1 g of bovine IgG, was administered orally. Volunteers adhered to a clear liquid diet from eight hours until four hours prior to BIC-*C difficile* administration at which time they consumed a standard meal (fig 1). They then fasted for four hours after which the BIC-*C difficile* was consumed, reconstituted in 250 ml of an isosmotic polyethylene glycol solution containing 421 mg of sodium bicarbonate. The polyethylene glycol was used as a non-absorbable transit marker. After a further two hours they were again allowed to take clear liquids until the end of the six hour ileostomy fluid collection period. Each volunteer was studied on four separate occasions with an intervening washout period of at least 72 hours. The BIC-*C difficile* was taken alone, with an antacid (30 ml of BICTRA), during therapy with the proton pump inhibitor omeprazole (Prilosec 20 mg, twice daily for four days prior to and on the day of BIC ingestion), or within enteric coated capsules. Treatments were administered in random order except omeprazole which was given last to avoid any carryover effect. The enteric coated capsules were designed to increase colonic delivery of BIC-*C difficile* and were prepared by the Pharmaceutical Service Division, College of Pharmacy, University of Iowa, as described previously.

Twelve hard gelatin capsules were consumed, each containing 420 mg of BIC and coated with two layers of cellulose acetate phthalate.

**Collection of ileostomy fluid**

Fluid was completely drained from the volunteers’ ileostomy appliance immediately prior to...
**BIC-C difficile** ingestion and, at 30 minute intervals, for six hours thereafter. The fluid volume was recorded and an aliquot taken which was filter sterilised and used for cytotoxicity neutralisation experiments. Protease inhibitors were added to the remaining fluid which was immediately frozen and stored at −20°C. The final concentrations of protease inhibitors were as follows: aprotinin 10 µg/ml, leupeptin 10 µg/ml, phenylmethylsulphonyl fluoride 10 µg/ml, N-p-tosyl-l-lysine chloromethyl ketone 10 µg/ml, and 1-1-tosylamido-2-phenylethyl chloromethyl ketone 10 µg/ml.

**SPECIAL LABORATORY STUDIES**

Total bovine IgG levels in the ileostomy fluid were measured by single radial immunodiffusion. Specific anti-*C difficile* toxin A bovine IgG levels were measured by enzyme linked immunosorbent assay (ELISA) as previously described. Cytotoxicity was determined by rounding of Chinese hamster ovary (CHO) cells (American Type Culture Collection, Rockville, Maryland, USA) in monolayer culture after exposure to purified *C difficile* toxin A. The minimum 50% cytotoxic dose of toxin A was defined as the minimum dose resulting in 50% cell rounding at 24 hours (40 ng/ml in these experiments). Inhibition of cytotoxicity was quantified by adding serial twofold dilutions of the ileostomy fluid to purified *C difficile* toxin A to achieve an end concentration of twice the minimum 50% cytotoxic dose of toxin A. After 30 minutes the mixture was added to the cell monolayers and cell rounding was assessed after 24 hours as previously described.

**STATISTICAL ANALYSES**

Statistical analyses were performed using SigmaStat for Windows version 1.00 (Jandel Scientific Software, San Rafael, California, USA). Unless stated otherwise, analysis of variance was used for intergroup comparisons and the Spearman rank order test was used to evaluate correlations.

**Results**

**STUDY SUBJECTS AND SAMPLE COLLECTION**

Six volunteers participated in the study and all completed the study protocol. However, during the study one volunteer developed symptoms of ileostomy dysfunction and, two weeks after study termination, required hospital admission because of small bowel obstruction. This resolved without surgical intervention. Ileostomy samples from this volunteer were excluded from the data analysis. No other significant adverse experiences were encountered by the study volunteers.

**ILEAL RECOVERY OF BOVINE IgG AFTER ORAL BIC-C DIFFICILE**

Figure 2 shows the total free bovine IgG content of the six hour ileostomy fluid collections. When the BIC-C difficile was taken alone the mean (SEM) recovery of bovine IgG from the terminal ileum was 1033 (232) mg. This represented 49% of the total ingested dose of 2100 mg of bovine IgG. The total ileal recovery of bovine IgG was 636 (129) mg (30% of the oral dose) when additional acid buffering capacity was provided in the form of an oral antacid. The omeprazole group had the highest ileal bovine IgG content at 1052 (268) mg (50% of the oral dose). However, analysis of variance did not reveal any significant difference between these three treatment groups (p=0.13).

The mean recovery of free bovine IgG from the ileal fluid after ingestion of BIC-C difficile in enteric coated capsules was 89 (45) mg (4% of the oral dose). This was significantly less than for the BIC-C difficile alone and omeprazole groups (p<0.05 for both). However, many of the capsules were recovered intact or only partially opened. Furthermore, when free bovine IgG was detected it was only found at the latter end of the six hour collection period (table 1). For all other treatment groups peak bovine IgG levels were detected much earlier (table 1) and negligible amounts of bovine IgG were found at six hours (fig 3). Since the release of bovine IgG was delayed and a substantial amount remained encapsulated the measurement of free bovine IgG in the ileal fluid is an incomplete measure of BIC-C difficile survival in this treatment group.

**Table 1 Time of peak bovine IgG recovery from ileal fluid following administration of oral bovine immunoglobulin concentrate-C difficile**

<table>
<thead>
<tr>
<th>Subject</th>
<th>Alone</th>
<th>Antacid</th>
<th>Omeprazole</th>
<th>Capsules</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>60 min (1632 mg)</td>
<td>90 min (963 mg)</td>
<td>90 min (901 mg)</td>
<td>360 min (118 mg)</td>
</tr>
<tr>
<td>2</td>
<td>60 min (1558 mg)</td>
<td>60 min (1396 mg)</td>
<td>300 min (57 mg)</td>
<td>300 min (57 mg)</td>
</tr>
<tr>
<td>3</td>
<td>150 min (611 mg)</td>
<td>150 min (677 mg)</td>
<td>210 min (1275 mg)</td>
<td>360 min (244 mg)</td>
</tr>
<tr>
<td>4</td>
<td>150 min (596 mg)</td>
<td>180 min (195 mg)</td>
<td>150 min (622 mg)</td>
<td>240 min (26 mg)</td>
</tr>
<tr>
<td>5</td>
<td>150 min (767 mg)</td>
<td>120 min (534 mg)</td>
<td>180 min (1067 mg)</td>
<td>0 mg</td>
</tr>
</tbody>
</table>

The collection time point at which ileal fluid contained the greatest amount of bovine IgG is shown for each subject and each study arm. The total amounts of free bovine IgG recovered during each six hour collection period are also shown (in parentheses).
The highest bovine IgG concentrations were observed between 60 and 180 minutes. Very little IgG was recovered after 300 minutes. In Figure 3, a biphasic pattern is evident with two peaks of bovine IgG concentrations: the first at 60 to 90 minutes and the second at 150 to 180 minutes after ingestion. In fact, this reflects two groups of subjects with different upper gastrointestinal tract transit times rather than a biphasic delivery of IgG to the distal small intestine. As shown in Table 1, two subjects (1 and 2) had consistently shorter transit times causing the early peak at 60 to 90 minutes. The remaining three subjects had slower transit times and these accounted for the second peak in mean bovine IgG concentration at 150 to 180 minutes. Release of free bovine IgG from enteric coated capsules was limited and delayed.

When the three groups receiving reconstituted BIC were examined, the average recovery of bovine IgG was twofold higher in the two subjects (1 and 2) with short transit times (1421 mg; 68% of the total dose) when compared with the three subjects with longer transit times (705 mg; 36% of the total dose; p=0.047 by paired t test).

The ability of ileal fluid samples to neutralise the biological activity of C difficile toxin A was then determined using the tissue culture cytotoxicity assay. Ileal fluid samples containing high levels of bovine IgG showed toxin neutralising effect whereas those with no measurable bovine IgG had little if any toxin neutralising activity. Negative control ileostomy fluid, obtained from study subjects immediately prior to BIC-C difficile ingestion, showed no detectable toxin neutralising activity. When the toxin A neutralising titre of a range of ileal fluid samples was determined, a positive correlation between neutralising titre and ileal fluid bovine IgG level was confirmed (r=0.95, p<0.001, n=24) (fig 5).

**Discussion**

The main finding of this study is that almost half of the bovine IgG in an oral dose of BIC-C difficile can be retrieved intact from the distal small intestine in humans. The surviving IgG
retains its specific antigen binding activity, in this instance binding to and neutralising C difficile toxin A. This finding supports the feasibility of using BIC as oral passive immunotherapy against enteric pathogens. Oral administration of BIC results in substantial levels of bovine IgG within the lumen of the human small intestine. Thus, infectious diarrhoea caused by small bowel pathogens such as Cryptosporidium parvum, microsporidial species, or Vibrio cholerae may be amenable to prophylaxis or therapy using appropriate hyperimmune bovine collostral antibody preparations. Our study findings also suggest that BIC may be effective in colonic infectious diarrhoea since a substantial portion of the oral dose of bovine IgG is delivered intact to the caecum.

Roos et al. administered a {superscript}15N-labelled preparation of bovine immunoglobulin concentrate to volunteers and sampled their ileal content using a nasointestinal tube. They calculated a 19% recovery rate for intact {superscript}15N-IgG in the ileum, substantially lower than the 49% recovered in this study. In an earlier study, we found that exposure of BIC-difficile to acidic human gastric secretions (pH<4) resulted in rapid loss of bovine IgG activity. In this study, the BIC-difficile was administered in a solution containing sodium bicarbonate. The additional buffering capacity of this solution may have improved bovine IgG survival in the stomach, thereby increasing ileal recovery of intact IgG.

The use of additional antacid or a proton pump inhibitor did not result in any further significant increase in bovine IgG survival. Thus, it appears unnecessary to use these extra measures to protect oral BIC-difficile from gastric acid degradation.

In our earlier study, we also found that faecal bovine IgG levels are notably increased when oral BIC-difficile is administered within enteric coated capsules. Our previous interpretation of this finding was that the enteric capsules protected BIC from gastric acid degradation. The findings of this study suggest an additional explanation. The capsules frequently passed through the stomach and small intestine intact. When bovine IgG was released from the capsules, this typically occurred at the latter end of the six hour ileo-

Figure 5 Comparison of specific anti-C difficile toxin A neutralising activity and total bovine IgG levels in ileal fluid after oral BIC-difficile. The greatest dilution of ileal fluid which neutralised toxin A is shown together with the bovine IgG level of that ileal fluid sample.

This interpretation is further supported by our finding that volunteers with slower mouth to ileum transit times had significantly lower bovine IgG recovery than those with more rapid transit. For example, when BIC-difficile was taken alone 76% of the oral dose was recovered from the distal ileum in volunteers with transit times of less than two hours whereas only 31% was recovered from those with transit times of more than two hours. A prolonged gastrointestinal transit time was also found to reduce bovine IgG recovery from the stool. Thus, it appears that enteric encapsulation raises faecal bovine IgG levels by delaying the release of BIC until the distal small intestine or colon. Formulation of BIC-difficile for targeted release in the colon may increase its efficacy in neutralising C difficile toxins within the colonic lumen and at the same time allow a reduction in the oral dose.

Animals immunised against C difficile toxins are protected against challenge with toxigenic C difficile whereas non-immunised animals develop fatal enterocolitis. Some, but not all, clinical studies indicate that patients with severe or recurrent C difficile colitis have an inadequate antibody response to C difficile and its toxins. The first animal study describing passive immunotherapy for C difficile enterocolitis was reported in 1979; hamsters were treated with parenteral Clostridium sordellii antitoxin, which cross reacts with C difficile toxins A and B, and protected against clindamycin induced colitis. Protective oral immunisation of hamsters using breast milk was reported in 1987. There are also limited reports of passive immunotherapy against C difficile diarrhoea and colitis in humans. A number of patients with recurrent, or refractory, C difficile diarrhoea have been treated using pooled normal human IgG administered intravenously. This treatment increases serum anti-C difficile toxin antibody concentrations and was effective in controlling symptoms in the small number of cases reported. In the present study, we showed that oral administration of BIC-difficile provides C difficile toxin binding and neutralising activity in the distal ileum in human volunteers. A randomised, controlled, clinical trial examining the efficacy of BIC-difficile in treating mild to moderately severe C difficile diarrhoea is now underway.

**References**

Bovine immunoglobulin concentrate—C difficile