Pathogenesis of postantibiotic diarrhoea caused by Clostridium difficile: an in vitro study in the rabbit intestine

S GUANDALINI, A FASANO, M MIGLIAVACCA, M C VERGA, P MASTRANTONIO GIANFRILLI, A FERRARA, M ALESSIO, B MALAMISURA, P GALATI, A PANTOSTI, AND A RUBINO

From the Department of Pediatrics, 2nd School of Medicine, University of Naples, Italy, Istituto Superiore di Sanita, Rome, Italy, and Institute of Pathology, Veterinary School, University of Naples, Italy

SUMMARY To elucidate the pathophysiological changes leading to postantibiotic diarrhoea caused by Clostridium difficile and its cytotoxin, oral ampicillin was given to rabbits, and jejunal, ileal, and caecal segments of those that developed diarrhoea were investigated in vitro. The rabbits that, in response to treatment, harboured Clostridium difficile in their colonic lumen were studied, and the results expressed according to the presence or absence of Clostridium difficile and/or its cytotoxin. Thus, we refer to either CD+ or CD− segments. The influx of glucose, phenylalanine, glycylphenylalanine, and lysine across the brush border of jejunum and ileum of CD+ segments was severely impaired, while only slightly blunted in CD−. No significant change was detected in the influx of glutamic acid in the jejunum of all treated animals and in the CD− ilea. Morphologic damage in ileum and caecum of CD+ was also more evident than in CD−. Transepithelial ion transport across short circuited ileal mucosa (CD+ and CD−) revealed secretory changes in Cl net transport that were more marked in CD−. We conclude that: (1) Clostridium difficile may also colonize the upper intestinal tract, where it induces morphological and functional damage, severely impairing nutrient absorption; and (2) the ileum contributes to the diarrhoea caused by CD even when the micro-organism is confined to the more distal gut by showing moderate impairment of nutrient absorption and marked electrolyte secretion.

The administration of antibiotics, particularly by the oral route, may be accompanied by gastrointestinal complaints, the most severe of which is diarrhoea. It is well known that Clostridium difficile (CD) may mediate such a complication by inducing a colitis of variable severity; the most widely recognised form being pseudomembranous colitis, an entity occurring both in adults and in children. Although it is well known that CD releases toxic factors (two of which have been extensively investigated – toxin A or enterotoxin and toxin B or cytotoxin), the pathophysiological changes actually ensuing after CD development caused by antibiotic treatment and leading to diarrhoea have not been well characterised.

The aim of our study was therefore to investigate the changes in structure and/or function arising in several intestinal segments in animals receiving antibiotic treatment that developed CD and diarrhoea.

Methods

ANIMALS Male New Zealand white rabbits fed ad libitum were used throughout the study. They received oral ampicillin (40 mg/kg body weight/day) in a single daily dose for 10 days or until diarrhoea developed. Of the 25 rabbits so treated, three did not develop
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diarrhoea. Of the remaining 22 (in which diarrhoea occurred after a mean of 5-2 days), eight died quite soon after its onset and could therefore not be studied.

The remaining 14, as well as six untreated, control rabbits, were killed and underwent the following investigations.

Microbiology and Identification of Clostridium difficile and its Cytotoxin

Cultures for enterobacteria and Clostridia (CD and Clostridium spiroforme) were performed on the stools of all animals, including controls, before treatment and on the intestinal contents of jejunum, ileum and caecum after death.

Cultures for Clostridia were carried out in an anaerobic cabinet using Columbia blood agar after ethanolic shock and CCFA (Oxoid) for CD. The identification was done by biochemical reactions and gas chromatographic analysis. All samples were also examined for cytotoxic effects on monolayers of CHO, Hep2, and Vero Cells using 96 well microtitre plates. Clostridium difficile antitoxin (Virginia Polytechnic Institute) and Clostridium perfringens iota antitoxin were used to neutralise respectively CD and Clostridium spiroforme cytotoxins. Although we did not search for toxin A, the detection of toxin B can be assumed to signal the simultaneous presence of toxin A, as no strain of CD has yet been isolated that would produce either toxin alone.

Histological Studies

Fragments of different intestinal segments were taken after death and fixed in 10% formol saline. The samples were then embedded in paraffin wax, sectioned and stained with haematoxylin and eosin for examination by light microscopy.

Nutrient Influx Studies

These were done according to a published procedure. Briefly, 10 cm segments of jejunal or ileal mucosa were preincubated in lucite influx chambers, allowing four adjacent portions of the mucosal side (each having a surface area of 1-12 cm²) to be bathed on the luminal side with a bicarbonate Ringer's solution having the following composition (in mmol/l): Na+, 141; K+ 5; Ca++ 1.25; Mg++ 1.1; Cl− 61; SO4− 30.5; HCO3− 25; H2PO4− 0.3; HPO4− 1.65; and mannitol, 30.5; and gassed with 5% CO2 in O2 yielding a pH of 7.4, at 37°C for 20 minutes. Incubations then followed by substituting a solution of identical composition but with the presence of the substrate being tested, labelled with 14C, and 3H-Inulin as a marker of the extracellular space. After 45-50 sec, incubation was stopped by quickly removing the test solution and adding 0.3 M mannitol. Each piece was then punched out, blotted on filter paper, homogenised in 10% trichloroacetic acid, and centrifuged. Aliquots of supernatants were assayed for radioactivity in a liquid scintillation counter. Calculations were done according to the described method.

Ion Transport Studies

These were done as previously described, using Ringer solution. Two pieces of stripped ileum and two pieces of stripped caecum from each animal were mounted in Ussing chambers. Transepithelial electrical potential difference (PD), total electrical conductance (Gt), and short-circuit current (Isc) were measured as previously described. Oppositely directed, unidirectional transepithelial fluxes of sodium and chloride from mucosa to serosa (m-s) and from serosa to mucosa (s-m) were measured under short circuit conditions using 22Na and 36Cl as previously described. Ji net=Jm-s−Ji s-m; Ji net different from 0 represents active transport of the ion species i. JR net is calculated as: JR net=Jsc−JNa net +JCl net.

Cyclic Nucleotide Studies

These determinations were carried out according to the method of Harper and Brooker, using kits by Amersham. Immediately after death, samples from ileum and caecum, stripped from muscle and serosal layers, were frozen for subsequent determinations. After thawing, they were processed according to the previously described procedure. Results are expressed as picomoles of cAMP or cGMP/mg mucosal protein (protein content was determined by the method of Lowry et al.).

Statistical Analysis

All results are expressed as mean (SE). The significance of the differences has been calculated using the Student's t test for either paired or unpaired variates.

Results

Isolation of Clostridium difficile

Clostridium difficile was not detected in any of the stool samples collected before the beginning of therapy from every rabbit, or from the intestinal contents of any of the six control animals. Among the animals receiving ampicillin, CD and/or its cytotoxin were found in the colon of 13 animals (in the colon of the 14th rabbit another Clostridial strain – C. spiroforme, was detected. Results of studies on this animal are not included here). In three of such animals. CD and/or its cytotoxin were found throughout the gut; in five rabbits, they were detected only in ileum and colon, and in the remain-
Table 1  Detection of clostridia in rabbits developing diarrhoea after treatment with ampicillin

<table>
<thead>
<tr>
<th>Animal no</th>
<th>Species</th>
<th>Jejunum</th>
<th>Ileum</th>
<th>Colon</th>
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<tr>
<td>1</td>
<td>C. difficile</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>C. difficile</td>
<td>+</td>
<td>+</td>
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</tr>
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<td>+</td>
</tr>
<tr>
<td>14</td>
<td>C. spiroforme</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

The + and − signs mean the detection of clostridia and/or their cytotoxin (+) or lack of them (−) in the indicated intestinal segment.

Fig. 1  Nutrient influx in the jejunal mucosa of control (□), CD− (■), and CD+ (□□□) segments. Bars represent influx of different nutrients at concentration shown in the figure. Number of animals studied is given in parentheses. Brackets indicate 1 (SE). *p<0.05; **p<0.005.

Nutrient influx measurements

Results showing the rates of transport of glucose, phenylalanine, glycyl-phenylalanine, lysine and glutamic acid from the mucosal medium into the epithelium of either jejunal or ileum of control, CD+ and CD− segments are reported in Figs 1 and 2, respectively. It appears that in the jejunal, with the exception of glucose which is moderately inhibited, the absorptive transport of all tested nutrients is, at the concentrations tested, well preserved in CD− areas, while it appears clearly blunted in CD+ segments: only the influx of glutamic acid is in fact not significantly inhibited.

In the ileum, most nutrients are less effectively taken up by the mucosa of all diarrhoeic ampicillin treated rabbits. The uptake of all tested nutrients but glycyl-phenylalanine and glutamic acid was in fact significantly blunted in CD− segments; while that of all tested nutrients except glycyl-phenylalanine was impaired in areas where CD was detected. The rather higher degree of mucosal injury seen in CD+ ileum provides a reasonable morphologic support for such observation (see below).

Transepithelial ion fluxes

Ileum

The transport of Na and Cl across stripped, short circuited mucosa of control, CD− and CD+ ilea was also measured. Results are reported in Table 2. In controls, the m-s flux for both ions is bigger by about 2 μEq than the s-m corresponding flux, thus resulting in net absorption of both Na and Cl.

In contrast, in CD− ileum, while there was no statistically significant change in Na transport, JCl m-s was significantly reduced, leading to the development of a Cl net secretory state.

The changes observed in the CD+ ileal segments differ from the latter: first of all, a marked, significant

Table 2  Transepithelial fluxes in the ileum

<table>
<thead>
<tr>
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<td>F^10ms</td>
<td>15.02 (1-62)</td>
<td>12.75 (1-17)</td>
<td>2.27 (1-19)</td>
<td>7.95 (1-16)</td>
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<td>F^1ms</td>
<td>5.77 (1-08)</td>
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<td>1.28 (0-9)</td>
<td>1.19 (1-0)</td>
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<tr>
<td>F^3ms</td>
<td>0.86 (0-36)</td>
<td>0.34 (1-35)</td>
<td>34.90 (3-56)</td>
<td>33.24 (2-5)</td>
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<tr>
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<tr>
<td>F^4ms</td>
<td>10.06 (1-67)</td>
<td>11.07 (2-66)</td>
<td>12.05 (1-02)</td>
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<td>&lt;0-05</td>
<td>&lt;0-05</td>
<td>&lt;0-05</td>
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</tr>
<tr>
<td>ISC</td>
<td>1.97 (1-05)</td>
<td>10.77 (1-67)</td>
<td>11.07 (2-66)</td>
<td>11.07 (2-66)</td>
</tr>
<tr>
<td>JCl</td>
<td>1.28 (0-9)</td>
<td>1.19 (1-0)</td>
<td>33.24 (2-5)</td>
<td>33.24 (2-5)</td>
</tr>
</tbody>
</table>

Values are mean (SE) for the number of animals reported in square brackets. Fluxes and ISC are in μEq/h/cm²; Gt is in mmol/cm².

*significance of differences calculated comparing CD− with control rabbits; †significance of differences calculated comparing CD+ with control rabbits; ‡significance of differences calculated comparing CD+ with CD− rabbits.
increase in Gt (reflecting the enhanced mucosal permeability to electrolytes) is seen. As a result, all measured unidirectional fluxes are also enhanced. The modifications in net electrolyte transport are minor, as no net secretion developed: JNa net is unaffected, JCl net is abolished, because of an increment in its s-m component. Thus, in CD+ as compared with CD− ilea, an increase in tissue ion permeability and a lesser secretory shift in Na and Cl net transport is seen.

**Caecum**

Table 3 reports the results obtained when measuring the steady state transepithelial fluxes of Na and Cl across the short circuited mucosa of the caecum. Here too, in controls, Na and Cl are actively absorbed, to a similar extent. The magnitude of unidirectional fluxes is consistently lower than in the ileum (as pointed out by the lower Gt). In the CD+ caecal mucosa, Na and Cl transport rates are profoundly affected, with their m-s fluxes being impaired. Such decrease (which is larger for Cl) makes net Na absorption no longer significant and results in the development of a net secretory state for Cl transport. A sizeable rise in Isc accompanies such changes.

**HISTOLOGY**

Sections from CD+ and CD− ileum show morphologic damage, with the more severe changes (mucosal atrophy with severe villous blunting, crypt hyperplasia and mononuclear infiltration) being seen in the intestinal segments harbouring CD and/or its cytoxin. In CD+ caecum, severe dysepsielisation with disruption of gland architecture, mucus depletion, oedema, haemorrhagic effusions, and inflammatory cell infiltration is also evident.

**CYCLIC NUCLEOTIDES**

As the observed secretory changes in electrolyte transport could be theoretically attributed to increases in tissue concentrations of cAMP and cGMP, we measured their concentrations in ileal and caecal mucosa. We found no increase in either cyclic nucleotide in any of the tested conditions: in the ileum, cAMP averaged 95 (5)% of the control value in CD− segments and 112 (15)% in CD+; cGMP averaged 108 (7)% of control, while CD+ 90 (10)%.

In the caecum, cAMP in CD+ was 69 (20)% of control, and cGMP was 98 (4)%.

**Discussion**

Diarrhoea not uncommonly follows an antibiotic course: in its most serious presentation, this complication may be because of a colitis bearing the typical pseudomembrane formation – that is, pseudomembranous colitis. Such long known complication of antibiotic therapy has been shown in recent years to be due to the Gram positive anaerobe CD. Although it is known that CD releases at least four toxic factors, the pathophysiological changes that they mediate and that ultimately lead to diarrhoea (particularly where no pseudomembranous colitis is found) are not well defined. In this regard, the present study is the first report investigating, in an animal model *in vitro*, the intestinal alterations occurring after the induction of CD− mediated diarrhoea *in vivo*.
The major conclusions that can be drawn can be summarised as follows: (1) The administration of antibiotics may be followed by CD colonisation not confined to the large bowel, but extending to the upper gut. Such data are consistent with already reported scanty observations of CD detection in the duodenal juice in man and improves our understanding of the pathogenesis of diarrhoea caused by CD: in fact, the absorptive transport of nutrients was found to be severely affected in all the small bowel segments where CD was detected. Also, the CD ileum showed a highly significant increase in tissue permeability and the abolition of Cl absorption. (2) Small intestinal segments devoid of CD also show diarrhoeagenic pathophysiological changes: in fact, while morphology and nutrient transport appear relatively spared, net Cl secretion develops. (3) The presence of CD in the caecum induces secretory changes in Cl transport and a rise in Isc.

The changes that we observed can be conceivably traced to the known virulence factors of CD: namely, toxin A (‘enterotoxin’), toxic B (‘cytotoxin’) (both reviewed in ref 11), a heat labile toxin, and a ‘motility altering factor’. Although the aim of our work was not focused on determining which factor caused which change, we may, however, speculate that toxin A (which was recently shown to be the most active one) is responsible for the morphologic injury throughout the intestine and for the enhanced permeability in the ileum; on the other hand, the secretory changes seen in electrolyte transport in ileum and caecum, may be ascribed again to toxin A, as well as to the more recently described heat labile toxic factor. As for the secretory changes seen in GI segments proximal to sites of CD colonisation, they might be attributed to local mediators of inflammation, known to have a role in intestinal secretion, to the possible adsorption of CD toxin to more proximal tracts, or to the development of other enterotoxigenic bacteria. The latter possibility is, however, less likely, as we did not detect known enterotoxigenic bacteria (data not shown) in any of the sampled GI tracts, nor did we find increases in cyclic nucleotide levels that might have expected to occur as a result of stimulation by the known heat stable or heat labile enterotoxins.

References

1 Bartlett JG. The pseudomembranous enterocolitides.


*Since the first submission of this manuscript, the paper by Mitchell et al. *The effects of Clostridium difficile toxins on stripped rabbit ileal mucosa in Ussing chambers* appeared in J Med Microbiol 1987; 23: 199–204. The paper describes, in the same animal model used by us, the effects of CD toxins on ion transport, with findings that are in many ways very similar to those reported by us. These results therefore: (1) add further support to our observations which unlike those of Mitchell et al., were obtained in the intestine of animals that had previously been exposed, in vivo, to the development of the infection and; (2) indicate that toxin A, as we speculated in the discussion is responsible for the observed changes in CD+ segments.
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