Role of bile in non-specific defence mechanisms of the gut

T Kalambaheti, G N Cooper, G D F Jackson

Abstract
The effect of depriving the intestine of bile for 48 hours was studied to determine any influence on various parameters of innate immunity in the gastrointestinal tract. Groups of rats were prepared by bile duct cannulation (with or without fluid replacement) or bile duct ligation. Normal and sham operated animals were used for comparing the thickness of the mucus layer and the cells contained therein, enumeration of goblet cells, and measurement of villus size. Histological examination indicated that the intestinal tissues of treated and control rats were similar. Though villus size and numbers of goblet cells were unaffected, a significant reduction occurred in the thickness of the mucus blanket in the duodenal regions of rats deprived of bile, and there were significantly lower numbers of mucus associated enterocytes and lymphocytes, suggesting a lower turnover rate of the epithelium. The balance of the bacterial populations in the caecum and intestine was altered by bile deprivation – increased numbers of coliform organisms were found in both regions. The range of factors, including antibodies and other known constituents, present in bile may contribute to the maintenance of tissue integrity and influence the balance in indigenous bacterial populations in the intestine. Disturbance of the host’s biliary system and concomitant effects on the microbial flora may weaken the overall processes of defence in the intestine.

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In recent times considerable attention has been given to the secretory adaptive immune system in mediating protection in the gastrointestinal tract. The induction and function of secretory IgA antibodies has been given high priority by those interested in exploiting immune mechanisms to control important bacterial and virus diseases of man. The finding that the hepatobiliary system carries immunoglobulins raises the possibility that it may also contribute to immunity through delivery of protective antibodies to the gut fluids, but its role in innate defence against microbial infection is unclear. The increased incidence of infections with organisms normally regarded as constituents of the gut bacterial ecosystem in a variety of conditions that affect the production and/or flow of bile might suggest that the absence of bile from intestinal fluid has disturbed, and presumably reduced, the efficacy of the natural defence mechanisms.

The present studies were undertaken in an attempt to determine whether bile has any impact on the natural intestinal defence mechanisms. The rat was chosen because it does not possess a gall bladder or cystic duct, thus making it possible to deprive the intestine of bile (but not pancreatic juices) through ligation or cannulation. The obvious approach of challenging animals treated in such a way with specific microbial infections was clearly not appropriate, however, as any increased susceptibility could easily be attributed to the trauma resulting from the surgical procedures. The alternative has been to examine the effects of bile deprivation on several physical and biological parameters believed to be involved in innate resistance. Since bile flows into the lumen of the intestine, it is reasonable to assume that its effects, if any, will be associated with resistance mechanisms expressed at or associated with the epithelial surfaces; several such factors have been investigated herein.

Methods

ANIMALS
Male Wistar rats of approximately the same age and weighing between 250 and 350 g were used. Groups designated as normal rats (N) were housed in plastic boxes on sawdust bedding, fed with a commercial fluorine and urea free diet (Rat and Mouse Feeds, Allied Feeds, Sydney), and given free access to water.

SURGICAL METHODS
Groups of rats were anaesthetised with ether and treated as follows:

(i) Bile duct cannulation
The method described by Lambert was used. Briefly, polythene tubing (internal diameter 28 mm and outside diameter 0.68 mm) was inserted into the bile duct at a point further than 10 mm from its entry into the duodenum. After surgery the animals were housed in restraining cases and were given food and free access to water over that period.

(ii) Bile duct cannulation with fluid replacement
The bile ducts were cannulated as described in (i). A small opening was made in the abdomen close to the point at which the duodenal papillae connects with the abdominal wall. Polythene tubing was inserted through this
opening and then into the duodenum in the direction of the ileum and fixed with two purse string sutures of nylon silk and one of surgical cat gut. The tubing was connected to a peristaltic pump (Microperpex, LKB) which maintained a flow of sterile normal saline of 0·7 ml/hr into the duodenum. This rate was chosen as it approximated to the flow of bile through the cannula. The animals were kept in restraining cages and given food and free access to water.

(iii) Bile duct ligation
The common bile ducts of each animal were ligated in two places with silk sutures; rats were kept in groups of four in plastic cages and given food and free access to water.

(iv) Sham operation
Groups of rats were anaesthetised with ether and a 25–30 mm incision was made through the abdominal wall. After five minutes, the incision was sutured and the animals were kept in the restraining cages used for the first two groups of rats.

Unless stated otherwise, treated rats were killed by spinal dislocation under ether anaesthesia 48 hours after surgery.

MEASUREMENT OF MUCUS GEL THICKNESS
A modification of the method described by Keress et al.\(^\text{10}\) was used. Briefly, 1–2 cm long sections of intestinal tissue were opened longitudinally and washed gently in saline. They were mounted on a Millipore filter with the luminal surface uppermost and 1·0–1·5 mm sections were cut with a razor blade placed at right angles to the mucosal surface. Clear adhesive tape was placed on glass microscope slides to support the sections, which were mounted transversely and viewed with an inverted microscope (×10 magnification). The mucus gel appeared as a translucent layer above the black of the mucosa and paper layers. Gel thickness was measured with an eyepiece graticule calibrated against a stage micrometer. Four sections of each of the tissue regions were examined and four separate measurements were taken for each section: mean (SD) values were determined from the 16 individual measurements. Unless otherwise stated, sections from the duodenum and the jejunum were taken 2–5 cm and 20–25 cm respectively from the pylorus, ileal sections were taken 5–10 cm from the ileocecal junction, and colonic sections were from the mid-point of the colon.

PREPARATION OF CELLS FROM INTESTINAL MUCUS
Rats were fasted for 18 hours then killed by spinal dislocation under ether anaesthesia. The small intestines were removed and divided into three equal length segments, each of which was flushed out with phosphate buffered saline (PBS-pH 7·2). They were then tied at one end with cotton, filled with 10–15 ml of N-acetyl-l-cysteine solution (prepared by adding 1·25 g to 50 ml of PBS and adjusting to pH 7·2 with 5 N NaOH) then tied at the other end. Small intestines were incubated for five minutes at room temperature then opened to collect the fluid contents. Each segment was again flushed with PBS (10 ml) and the washings from each segment were combined with its fluid contents. Cells were collected after centrifugation at 100 g for 10 minutes. The cells were washed once in RPMI 1640 medium (CSL, Parkville, Victoria) and recovered by centrifuging as before. They were then separated by Percoll density gradient centrifugation. Ninety two volumes of Percoll (Pharmacia, Sweden) were diluted with eight volumes of 10× concentrated PBS. Different concentrations of the Percoll were then prepared by dilution in RPMI 1640 medium containing 10% fetal calf serum (CSL, Parkville, Victoria). The gradient was prepared by layering 2·5 ml volumes of 80%, 35%, 25%, and 15% Percoll in 15 ml conical glass tubes; the cells were resuspended in 3 ml of 7% Percoll containing 3 mg/ml dithiotreitol (Sigma, USA) and layered onto the gradient surface. They were centrifuged at 700 g in a Beckman J6R centrifuge with a TH-4 rotor for 20 minutes at 4°C. Cells banding over the upper three layers were predominately mononuclear and despite variations in form from a few typical epithelial cells to large numbers of vacuolated and multigranular cells (classified as enterocytes), approximately 95% viability was found when assessed by the trypan blue dye exclusion test. The numbers of cells present in each preparation were determined by counting in a Neuberg haemocytometer. Preliminary studies of stained sections of intestinal segments indicated that the mucosal surfaces were largely unaffected by this treatment with N-acetyl-l-cysteine.
Determination of Villus Size and Numbers of Goblet Cells

Sections of intestine were prepared and stained by haematoxylin and eosin. The villus size was estimated by the method of Altmann and Leblond,11 examining 10 villi per section, and was expressed as the mean (SD) number of cells per villus-crypt unit. The numbers of unstained goblet cells in the 10 villus sections were determined and expressed as the mean (SD) number per villus-crypt unit.

Enumeration of Bacterial Populations

Duodenal, jejunal, ileal, caecal, and colonic sections weighing 0.5 g were placed in 5 ml of nutrient broth containing glass beads then stirred vigorously in a Vortex mixer. Sections of rat small intestine treated with N-acetyl-L-cysteine (w/v) were homogenized (and dilutions of them) were then stirred vigorously in a Vortex mixer. The thickness of the sections was estimated by the method of Altmann by haematoxylin and was expressed as the mean (SD) number of epithelial cells, multigranulated, and vacuolated cells.

Table I: Recovery of cells from mucus eluates prepared from sections of rat small intestine after five minutes’ incubation in 2-5% (v/v) of N-acetyl-L-cysteine

<table>
<thead>
<tr>
<th>Intestinal section</th>
<th>Mean (SD) no of epithelial cells×10⁴</th>
<th>Mean (SD) cells with morphological characteristics of epithelial cells (%)</th>
<th>Mean (SD) no of G and V cells (%)</th>
<th>Mean (SD) no of lymphocytes (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duodenum†</td>
<td>1500 (544)</td>
<td>12 (3)</td>
<td>72 (4)</td>
<td>16 (2)</td>
</tr>
<tr>
<td>Jejunum</td>
<td>349 (576)</td>
<td>9 (2)</td>
<td>72 (8)</td>
<td>19 (4)</td>
</tr>
<tr>
<td>Ileum</td>
<td>40 (19)</td>
<td>34 (10)</td>
<td>61 (3)</td>
<td>5 (2)</td>
</tr>
</tbody>
</table>

*Mean (SD) determined from seven rats; † includes 10 cm of ileum; ‡ multigranulated (G) and vacuolated (V) cells.

Table II: Effect of bile duct cannulation on numbers of mononuclear cells in mucus eluates prepared from section of rat small intestine treated with 2-5% (v/v) N-acetyl-L-cysteine

<table>
<thead>
<tr>
<th>Intestinal section</th>
<th>Mean (SD) no of epithelial cells×10⁴</th>
<th>Mean (SD) no of lymphocytes (%)</th>
<th>Mean (SD) no of epithelial cells×10⁴</th>
<th>Mean (SD) no of lymphocytes (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duodenum</td>
<td>1161 (531)</td>
<td>447 (352)</td>
<td>57 (48)</td>
<td>27 (31)</td>
</tr>
<tr>
<td>Jejunum</td>
<td>43 (31)</td>
<td>43 (26)</td>
<td>53 (44)</td>
<td>33 (36)</td>
</tr>
<tr>
<td>Ileum</td>
<td>4 (2)</td>
<td>0</td>
<td>41 (53)</td>
<td>23 (32)</td>
</tr>
</tbody>
</table>

*Mean (SD) determined from three rats; animals examined two days after operation; † mean (SD) determined from 10 rats; animals examined two days after operation; ‡ includes multigranulated and vacuolated cells.

Results

The thickness of the mucus blanket in the duodenum, jejunal, ileal, and colonic regions of the intestines of normal, sham operated, and ‘bile duct modified’ rats is compared in Figure 1. In both normal and sham operated rats, it is greatest in the duodenum (circa 80–100 μm) and least in the colon (circa 30 μm). When bile was excluded from the intestine by cannulation, the thickness of the mucus layer was reduced by around 80% in the duodenum and by 50% in the jejunum, but was unaffected in the colon. When tested by one way analysis of variance (ANOVA) the only significant reduction in mucus thickness was found in the duodenum (p<0.05). When saline infusion was used to replace the 15–20 ml of bile fluid entering the duodenum daily, mucus thickness in this region was still around 50% less than that found in normal and sham operated animals. This was also the case in rats in which systemic dehydration through loss of bile was prevented by bile duct ligation. In both these cases, reductions in the thickness of the duodenal blanket were statistically significant when tested by ANOVA (p<0.05) in comparison with normal and sham operated animals. No significant changes in mucus thickness were found in other regions of the intestine. Figure 2 indicates that the changes in the duodenum after the interruption of bile flow were not accompanied by reductions in villus size or in the numbers of goblet cells lining the duodenal villi.

Previous studies have shown that the intestine’s mucus blanket can be eluted by short term (3–5 minutes) treatment with the mucolytic agent N-acetyl-L-cysteine without significant damage to the epithelial surface itself.14 Table I indicates that mucus eluates obtained by this treatment contain cells

Figure 2: Goblet cells (left hand columns) and villus size (right hand columns) in the duodenum of normal and bile duct perturbed rats; error bars indicate SDs.
with varying morphological characteristics; moreover, their numbers differ in eluates prepared from different regions of the small intestine. In sections taken from normal rats, most mucus associated cells occurred in eluates from the duodenal-jejunal region whereas the fewest were found in ileal eluates. In all instances, granular and vacuolated cells predominated (in excess of 60% of the total); the remaining cells in each preparation were readily identified as lymphocytes and epithelial cells.

Mucus associated cells obtained from sections of small intestine prepared from sham operated and bile duct cannulated rats are compared in Table II. Exclusion of bile from the small intestine led to reductions of 10–20 fold in cell numbers present in mucus eluates prepared from the duodenal-jejunal region, whereas the numbers of cells recovered from more distal intestinal regions of the two groups of animals were not significantly different.

Microbial populations of the duodenum, ileum, caecum, and colon of normal, sham operated, bile duct cannulated (some receiving intraduodenal saline infusion) and bile duct ligated rats were assessed by viable counting procedures 48 hours after surgery. In the case of the anaerobic populations, numbers were assessed in the duodenal and jejunal regions (circa \(10^8\) per g tissue) and greatest in the caecum and colon (circa \(10^{9–10}\) per g tissue). Numbers of aerobic organisms were usually fewer (ranging from around \(10^6\) per g in the duodenum to \(10^{5–6}\) per g in the colon). In no instance were the numbers of these groups of organisms affected by bile duct cannulation or ligation. In contrast, as shown in Figure 3, increases in the numbers of organisms capable of growing on MCA (primarily coliforms) were found in most intestinal regions of rats which had undergone bile duct modification; the most significant and consistent changes were found in the caecum and colon.

Previous studies in mice have shown that increases of coliforms in the intestine were associated with quantitative and qualitative changes in the microbial flora of the caecum. The possibility that similar inter-relationships existed in bile deprived intestines of the rat was therefore investigated using an objective qualitative microscopic assessment method described by those authors. At the same time, coliform numbers in the caeca of the individual rats were determined by viable counts. The results are summarised in Table III. For simplicity, organisms present in the caecal contents were assigned to three distinct morphological groups. The first group clearly forms a large percentage of the caecal population in normal and sham operated animals but their numbers were affected when bile flow to the intestine was prevented over a two day period. For reasons which are not immediately obvious, changes in the populations were observed in only half of the bile duct cannulated animals. Importantly, however, in one of those subgroups, the percentage of fusiforms and tapered rods was greatly reduced while coliform numbers greatly exceeded those in the two control groups; in the other subgroup changes in neither of the two morphological groups were found. Findings in the group of rats which were bile duct cannulated and received intraintestinal saline infusion for two days also suggested that significant increases of coliform numbers were associated with decreases in the fusiform and tapered rod populations of the caecum.

### Table III: Effect of bile duct cannulation on rat caecal flora

<table>
<thead>
<tr>
<th>Rate treatment</th>
<th>No of rats</th>
<th>Fusiforms†</th>
<th>Curved rods‡</th>
<th>Others§</th>
<th>Mean (SD) no of coliforms (log10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>3</td>
<td>53</td>
<td>4</td>
<td>43</td>
<td>3-8 (0-4)</td>
</tr>
<tr>
<td>Sham operation</td>
<td>3</td>
<td>25</td>
<td>2</td>
<td>74</td>
<td>4-3 (2-0)</td>
</tr>
<tr>
<td>BDC-1**</td>
<td>3</td>
<td>3</td>
<td>2</td>
<td>99</td>
<td>5-4 (1-5)</td>
</tr>
<tr>
<td>BDC-2**</td>
<td>3</td>
<td>1</td>
<td>0</td>
<td>90</td>
<td>8-7 (0-9)‡‡</td>
</tr>
<tr>
<td>BDC (infused)††</td>
<td>3</td>
<td>14</td>
<td>0</td>
<td>86</td>
<td>7-9 (1-1)‡‡</td>
</tr>
</tbody>
</table>

* Determined by microscopic examination of slides prepared from caecal contents of rats two days after surgery; † includes Gram negative organisms with fusiform or tapered rod morphology; ‡ includes Gram negative curved rods and spiral shaped organisms; § includes Gram negative and Gram positive rods, cocci and cocco-bacilli; ** bile duct cannulated animals; †† bile duct ligated animals; sub-groups based on presence of normal or high numbers of coliforms determined by viable counting of caecal contents; ‡‡ statistically different from numbers found in normal and sham operated groups (p<0.05 by the ANOVA test).

**Discussion**

Mucosal surfaces, which are at continuous risk from environmental agents, are equipped with a full array of innate and adaptive immune processes in order to maintain tissue integrity. The magnitude of the surface area of the gastrointestinal tract and, under normal circumstances, the presence of ingested material together with the indigenous microbiota demand a particularly potent defence system, yet one which has great subtlety so that the prime function of food absorption is not compromised. Our interest in the hepatobiliary transfer of immunoglobulins and their contribution to immunity in the gut has led to this initial study into the possibility that biliary factors influence other aspects of natural host defence. In general terms, the results show that after a short period of bile deprivation, significant changes occur in...
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certain physical and biological properties of the intestine without any apparent changes to tissue structure or the cellular distribution within the epithelium.

Mucus provides a thick gel coating for the intestinal epithelium and is a natural physical barrier to potential pathogenic bacteria; only adaptable spiral organisms are likely to be present in it.15 The results reported here suggest that while there is a requirement for adequate hydration, the thickness of the mucus coat in the duodenal region is partly dependent upon the normal flow of bile. This reduction could enhance the susceptibilty to invading organisms by permitting reader access to the epithelial surface across this natural barrier. There is an indication of reduced synthesis or output from the mucus producing cells, a view supported by the finding that the numbers of goblet cells were similar in treated and control rats. One candidate molecule for this effect is histamine, which has been shown to influence mucus production16 and is known to be present in normal rat bile (Jackson and Storrer, unpublished observation).

The granular and vacuolated cells which predominate in mucus cluates carry the same surface markers (for example, asialo-GM1) and react similarly to specific cytochemical stains as cells with the characteristic morphology of epithelial cells. They are believed to be moribund forms of surface enterocytes which enter the mucus blanket after extrusion from the epithelial surface17 and have been grouped with epithelial cells in Table II. The numbers of these cells recoverable from duodenal mucus is greatly reduced in animals deprived of bile through cannulation of the bile duct. Possession of some, if not all, epithelial cell membrane characteristics suggests that these mucus associated cells may retain the ability to participate in bacterial adhesion reactions. Indeed, unpublished studies with Vibrioh cholerae in experimental intestinal infections clearly show that many of the organisms are firmly attached to mucus associated cells. As this is probably true for other gut associated pathogens which require attachment to the intestinal epithelium as part of their pathogenic process,18 gross reductions in the number of mucus associated enterocytes could increase the ease with which pathogens penetrate the mucus to reach their epithelial target cells. Greater susceptibility to infection might therefore be anticipated in individuals lacking regular bile flow into the duodenum. It is known that epidermal growth factor (EGF)19 is a constitutive molecule of bile; it is known to act in the lumen20 and is therefore a prime candidate for homeostatic regulation of epithelial cell turnover. Its absence through prevention of bile flow to the lumen could therefore account for the observed significant decrease in epithelial cells present in the duodenal mucus.

A significant role in innate resistance has been assigned to the resident bacterial flora of the gut provided they are maintained within their natural ecological niches and that the balance between species is undisturbed. The continual interplay between host and parasite must serve the purposes of both. Earlier studies in our laboratory21 22 demonstrated the presence of antibodies (particularly IgA) against intestinal bacteria in rat bile and found, in particular, that the presence of coliforms was a major determinant in inducing normal levels of circulating and intestinal immunoglobulins. The present studies have shown that prevention of bile can cause subtle changes in the bacterial composition of caecal contents in most animals so far examined. Also, enhancement of the numbers of coliforms and their more obvious presence in the small intestine lead to the suggestion that factors present in bile contribute to the normal distribution of these bacteria. These observations may be relevant to the well known increased incidence of Gram negative sepsis in hepatobiliary disorders. Related to this is the recent finding that a seven day absence of normal bile flow in rats leads to translocation of bacteria (particularly coliforms) into mesenteric lymph nodes.23 The likely overgrowth of intestinal bacteria in this condition, coupled with a decreased level of natural antibody or other as yet undetermined factor(s) may be important features in facilitating invasion.

In summary, we propose that bile contributes to the control of homeostatic processes of both host and parasite in the gastrointestinal tract and that disturbance of its natural flow is likely to put the host at risk from invasion by its resident bacteria. Of current concern is how the changes observed in this study might influence specific phases of an infection by exogenous pathogens.

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