BACTERIAL PATHGENICITY

Adhesion of enteroaggregative Escherichia coli to formalin-fixed intestinal and ureteric epithelia from children

SUSAN HICKS, D. C. A. CANDY and A. D. PHILLIPS*

Department of Child Health, King's College School of Medicine and Dentistry, London SE5 8RX and *Academic Department of Paediatric Gastroenterology, Queen Elizabeth Hospital for Children, London E2 8PS

The adhesion characteristics of enteroaggregative Escherichia coli (EAggEC) to the mucosal surfaces of formalin-fixed paediatric intestinal and ureteral tissue were studied. The technique offers a means of overcoming the problem of limited tissue access in childhood and a way of examining the initial steps of bacterial adhesion. Five EAggEC strains isolated from children with diarrhoea in the UK and a well characterised, prototype EAggEC strain (221) were examined. Five of the six EAggEC strains showed preferential adhesion to jejunal mucosa with limited adhesion to ileum and colon. Five of the six also adhered to ureteric tissue. EAggEC can adhere to proximal, as well as distal, regions of the gastrointestinal tract in children, a previously unrecognised characteristic.

Introduction

Enteroaggregative Escherichia coli (EAggEC) is a recently described category of E. coli that adheres to HEp-2 cells in an aggregative or 'stacked-brick' pattern [1]. This pattern is distinct from localised and diffuse patterns of adhesion which are also detected in the HEp-2 cell assay; localised adhesion is characteristic of enteropathogenic E. coli [2] cells which attach to, and efface, the brush border of the intestine [3], whereas the role of diffuse adhering E. coli in diarrhoea is unclear [1, 4, 5].

EAggEC strains have been implicated epidemiologically in acute and chronic diarrhoea in children from developing countries [6–8] and more recently in the UK [9]. Ethical considerations prohibit the use of large quantities of ex-vivo tissue from children and investigations into the pathogenesis of EAggEC-associated diarrhoea have included the use of animal models [10, 11], isolated enterocytes [12], brief incubations with ex-vivo human tissue [13], and in-vitro organ culture [14]. The use of formalin-fixed surgical specimens offers a means of overcoming limited tissue access and has been used to study bacterial adhesion in man [13, 15]. The one EAggEC strain studied previously [13] showed preferential adhesion to colonic mucosa. Generally, it is considered that EAggEC strains bind preferentially to the distal rather than the proximal bowel [13, 14], although Raj et al. have demonstrated adhesion to isolated duodenal enterocytes from adults [12].

The use of formalin-fixed tissue is practicable because, although fixation is achieved by the cross-linking of neighbouring protein groups, surface carbohydrate receptors remain intact [16]. Furthermore, the use of this fixed material, which otherwise would be destroyed after definitive histological examination, enables studies on many regions of the gastrointestinal tract and other mucosal surfaces.

The ability of EAggEC to adhere to ureteral mucosa was also examined because of the association of diarrhoea with urinary tract infection (UTI) [17], and because some E. coli of colonic origin share phenotypic characteristics with those that cause UTI [18]. E. coli of UTI origin have been shown to adhere to human ureteral cells in vitro [19]. However, only
one strain (TL100) of EAggEC of intestinal origin has been examined with fresh and formalin-fixed adult ureteral tissue [13] – strong adhesion to fixed ureteral tissue was found.

The aim of this work was to examine the adhesion characteristics of EAggEC strains in studies with formalin-fixed surgical tissue from paediatric patients.

Materials and methods

Bacterial strains

Five enteroaggregative E. coli strains (3862 [O85:H1], WJ19/10 [O126:H27], AN11/13 [O55:H4], HR15/6 [O126:H27] and LC9/6 [O111:ab:H21]) that had been isolated from children with acute and chronic diarrhoea at Queen Elizabeth Hospital for Children (QEHC) were examined. The strains had been detected by their aggregative pattern in the HEp-2 cell assay and were aggregative adherence (AA) gene probe positive [9]. The sensitive and specific gene probe used was that described by Baudry et al. [20] which recognises a 1-kb region on the 60-MDa plasmid present in EAggEC. Strain 221, another well characterised EAggEC isolate from an American student who had returned from Mexico with travellers' diarrhoea [21], was also examined. This strain was used subsequently to challenge adult volunteers and was found to cause diarrhoea [22]. An enteropathogenic strain of E. coli (EPEC KH1/8) O114:H2, also isolated at QEHC, was used as a positive control. This strain showed an attaching-effacing lesion in vivo in a biopsy of the proximal small intestine of a patient with chronic diarrhoea (A. D. Phillips, personal communication). In the HEp-2 cell assay, EPEC KH1/8 showed a localised pattern of adhesion in vitro in human small intestinal organ culture (S. Knutton, personal communication). In the HEp-2 cell assay, EPEC KH1/8 showed a localised pattern of adhesion in vitro in human small intestinal organ culture (S. Knutton, personal communication). In the HEp-2 cell assay, EPEC KH1/8 showed a localised pattern of adhesion in vitro in human small intestinal organ culture (S. Knutton, personal communication). In the HEp-2 cell assay, EPEC KH1/8 showed a localised pattern of adhesion in vitro in human small intestinal organ culture (S. Knutton, personal communication).

All bacterial strains were streaked on to MacConkey agar. Five pure colonies were removed after overnight incubation and stored at -70°C in Microbank cryovials (Pro-Lab Diagnostics) as the stock bacteria for adhesion assays. Before incubation with fixed tissue, all EAggEC strains were examined in a HEp-2 assay to ensure that the aggregative phenotype had been maintained during storage.

Tissue samples

Macroscopically normal, formalin-fixed, surgical tissue available after histological examination was used for the adhesion experiments.

Four jejunal specimens (from patients aged 12 days, 9, 36 and 135 months), five ileal specimens (from patients aged 12 days, 10, 13, 51 and 65 months) and five colonic specimens (from patients aged 6 days, 4, 24, 53 and 138 months) were used. Ureteral specimens from four children (aged 12, 19, 49 and 65 months) were examined; tissue from children under 12 months of age was not available.

Adhesion assay

The adhesion assay was performed as described by Yamamoto et al. [13]. Bacterial strains were incubated aerobically in brain heart infusion (BHI) broth for 18 h at 37°C without shaking. Formalin-fixed tissue was cut into 3-mm² pieces and washed twice in 100 ml of Krebs-Ringer buffer for 3 h at room temperature to remove residual fixative. Preliminary experiments showed that any trace of fixative remaining after washing did not affect E. coli viability. The tissue was then added to a mixture of 1 ml of bacterial culture in BHI (c. 10⁸ cfu as measured by serial dilution and plating on to MacConkey agar) and 1 ml of sterile phosphate-buffered saline (PBS) containing D-mannose 0.5% w/v and incubated statically for 45 min at 37°C.

After incubation, the tissue was washed three times with sterile PBS to remove non-adhering bacteria, fixed in phosphate-buffered glutaraldehyde 3% and post-fixed with aqueous osmium tetroxide 1%. After washing with distilled water, the tissue was dehydrated through a graded series of ethanol before critical point drying with liquid CO₂. The samples were sputter-coated with gold-palladium and coded before being examined in a JEOL JSM-5300 scanning electron microscope (SEM) so that the operator was unaware of the identity of the sample. Ten random fields were examined from each specimen at a magnification of 3500 and the number of coliform bacteria adhering to the mucosa in each SEM field was recorded. Results from each of the specimens were pooled according to mucosal region and experimental E. coli strain, to give bacterial counts from a total of 40 fields of view for jejunal and ureteral tissue, and 50 fields for ileal and colonic mucosa.

Counts from control and experimental samples were compared by non-parametric analysis by the Mann-Whitney test, with a p value of < 0.05 recorded as significant.

Pieces of jejunum, ileum, colon and ureter with PBS alone were used to establish the normal range (95% confidence limits) for the presence of bacteria without the addition of the experimental strains. For these experiments, the numbers of coliform bacteria from 40 SEM fields per mucosal specimen, at the same magnification, were counted.
Results

Controls

The median number of bacteria adhering to jejunum gave 95% confidence limits of 0–3 bacteria/SEM field. This value was 0–2 for ileal and 0–1 for colonic tissue. Generally, fewer bacteria were found adhering to ureteric specimens and the median value gave 95% confidence limits of 0–0.5 bacteria/field.

Adhesion to jejunum

Five of six EAggEC strains adhered significantly to paediatric jejunal tissue (Table 1 and Fig. 1a). Strains 3862, HR15/6, LC9/6 and WJ19/10 all demonstrated highly significant adhesion to jejunum ($p < 0.001$). Strain AN11/13 adhered in lower, but still significant ($p < 0.04$), numbers than the previous strains. All strains demonstrated evidence of aggregative adherence to the mucosal surface (Fig. 2b). The median number of adhering bacteria varied from strain to strain and between specimens. However, strain 3862 adhered to jejunum in significantly greater numbers than any other strain ($p < 0.05$). An example of strain 3862 adhering to jejunal mucosa in comparison to control tissue is shown in Fig. 2a and 2b. EPEC strain KH1/8 (positive control) showed significant adhesion ($p < 0.02$). Neither of the negative control E. coli

<table>
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<td>HB101</td>
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+, represents significant adhesion ($p < 0.05$); –, represents insignificant adhesion ($p > 0.05$).

Table 1. Adhesion of E. coli to formalin-fixed mucosae

Fig. 1. Box and whisker plots showing the level of E. coli adhesion to fixed jejunal (a), ileal (b), colonic (c) and ureteral (d) mucosae. The boxes represent upper and lower quartiles about the median. The vertical lines represent the range unless outliers are present (represented as dots), in which case the lines represent one and a half times the interquartile distance. Total numbers of fields studied were: (a) control 160, E. coli strains 40, except HB101 30; (b) control 200, E. coli strains 50, except HB101 30; (c) control 200, E. coli strains 50 except HB101 30; (d) control 160, E. coli strains 40. §, $p \leq 0.05$
strains SC13 and HB101 showed significant adhesion (p = 0.71 and p = 0.11, respectively) to any of the four jejunal specimens.

Adhesion to ileum

Five specimens from this gut region were examined. Five of six EA
ggEC showed significant adhesion (p < 0.03) (Fig. 1b). Although fewer bacteria were found adhering to the ileum than to the jejunum, in all cases, strain 3862 again showed significantly greater adhesion than any other strain (p < 0.001) (Fig. 2d). As with jejunum, EPEC KH1/8 showed highly significant adhesion (p < 0.001) to ileum but neither of the negative control strains SC13 nor HB101 showed significant adhesion to it (p = 0.15 and p = 0.13, respectively).

Adhesion to colon

As with ileal mucosa, fewer bacteria were found adhering to the five colonic specimens than to jejunal tissue (Fig. 1c). Four of the six EA
ggEC strains showed significant adhesion to colonic mucosa – 221 (p < 0.03), 3862, AN11/13 and WJ19/10 (p < 0.001). Both the EPEC and normal flora control E. coli strains also showed significant adhesion (p < 0.001) but the negative control strain HB101 did not (p = 0.06).

Adhesion to ureter

Five of six EA
ggEC showed significant adhesion (p < 0.0001) to ureteral epithelial cells (Fig. 1d). Strain 221 did not show significant adhesion (p = 0.28). Bacteria adhered in a more diffuse manner than to gut mucosae with the bacterial cells appearing as scattered individuals rather than in small groups (Fig. 2h). Both the EPEC and normal flora control E. coli strains showed significant adhesion (p < 0.001). Strain HB101 did not adhere in significant numbers (p = 0.98).

No correlation was found between the degree of bacterial adhesion and the age of the child providing the sample.

Discussion

The results demonstrate that each of the six EA
ggEC strains showed significant adhesion to formalin-fixed cells of the intestinal samples from children although there were regional differences. The use of formalin-fixed tissue to examine EA
ggEC adhesion to the intestine was first reported by Yamamoto and colleagues [13,15]. Similar results were found between unfixed and fixed tissue and the technique was applied to other bacterial pathogens, including Vibrio cholerae [24] and enterotoxigenic E. coli [25]. However, only one strain of EA
ggEC was examined and the tissue samples used came from a narrow age range of children [13]. The present study included more strains of EA
ggEC, and this may, in part, explain the variable pattern of the results. Yamamoto et al. [13] make no mention of finding bacteria adhering to the mucosal surface of control specimens and, therefore, any adhesion by the EA
ggEC strain was considered significant. However, a low level of bacterial adhesion was found in controls in the present study. Evans et al. [26] showed that viable bacteria could be isolated from jejunal mucosal homogenates from children without diarrhoea, suggesting that, in man, bacteria may be present in vivo at the mucosal surface without disease. This could account for the presence of bacteria found on control specimens.

Another reason for variation in results may be the requirement, in this study, that definitive histological examination be complete before the release of the tissue. Therefore, tissue remained in phosphate-buffered formaldehyde 4% for up to 90 days before use. It is possible that such a lengthy fixation, with exposure to the atmosphere on several occasions, could have altered the binding capacity of the tissue and could account for the adhesion to colonic mucosa observed with the negative control. Yamamoto et al. [13] used tissue after an unspecified period of fixation. However, because adhesion under these experimental conditions involves carbohydrate receptors, which are unaffected by formaldehyde, we discount this as an important factor.

Data presented here indicate that adhesion to jejunal mucosa may be a particularly important virulence factor of EA
ggEC. In the normal fasting state, the proximal small intestine is not sterile and contains 10^3–10^4 bacteria/ml of luminal fluid [27], yet this normal flora does not cause diarrhoea. The ability to adhere in vivo to jejunal mucosa in significantly large numbers with or without the expression of other known virulence factors, such as haemolysin-like toxins [28] or enterotoxins, could lead to the diarrhoea associated with EA
ggEC.

Previous reports of the distribution of EA
ggEC adhesion to intestinal mucosae are at variance with the results presented here. Knutton et al. found that EA
ggEC strains (including some isolated from children with diarrhoea from QEHC) adhered preferentially to adult colonic mucosa in organ culture [14] and one isolate examined with fixed intestinal tissue from children was found to adhere to ileal mucosa [13]. These reports suggest that EA
ggEC strains colonise distal ileum or colon. However, in the present experiments with fixed tissues of a wide range of specimens of jejunum, ileum and colon from paediatric age groups, EA
ggEC strains adhered to them all but particularly to jejunal mucosa.
Fig. 2. Scanning electronmicrographs showing enteroaggregative *E. coli* strain 3862 adhering to formalin-fixed mucosae: *a*, jejunum control; *b*, jejunum with EAggEC strain 3862 (note the aggregative groups of bacteria); *c*, ileum control; *d*, ileum with EAggEC 3862; *e*, colon control; *f*, colon with EAggEC strain 3862; *g*, ureter control; *h*, ureter with EAggEC 3862.
Fig. 2. Scanning electronmicrographs showing enteroaggregative *E. coli* strain 3862 adhering to formalin-fixed mucosae: a, jejunum control; b, jejunum with EAggEC strain 3862 (note the aggregative groups of bacteria); c, ileum control; d, ileum with EAggEC 3862; e, colon control; f, colon with EAggEc strain 3862; g, ureter control; h, ureter with EAggEC 3862.
The finding of Raj et al. [12] that 35% of EAggEC strains adhered to isolated duodenal enterocytes from adults agrees with our observations that adhesion to the proximal small intestine is an important characteristic of this group of bacteria. Adhesion to jejunal mucosa in vivo could have a significant adverse effect on the absorptive and secretory capacity of the tissue, i.e., on the major functions of this gut region.

Most of the EAggEC strains also adhered to the epithelium of the ureter. This suggests that the urinary tract also expresses receptors for gastrointestinal pathogens that survive formalin fixation. This observation deserves further investigation with more E. coli strains, both experimental and control, of gastrointestinal origin. In light of evidence suggesting that E. coli strains of urinary origin can adhere to human ureteral cells in vitro [19], it would be of interest to examine this adhesive characteristic in EAggEC with a view to a possible role of ureteral colonisation leading to urinary tract infection.

The normal flora E. coli strain SC13 adhered significantly to colonic and ureteric tissue. This is a surprising result and the ability of normal flora bacteria to adhere to intestinal mucosae should be studied further.

Heterogeneity appears to be an important feature of the EAggEC group, demonstrated here by the variation between strains in adhesion to fixed tissue. Unlike EPEC, in which certain serogroups are described as ‘classic’, many serogroups have been associated with EAggEC [6]. Five different serogroups were present in the six strains examined. Another feature of some EAggEC strains is that their DNA hybridises with a DNA probe for the gene region encoding the EAST 1 toxin. Although this characteristic is thought to be common amongst EAggEC [29], of the six strains studied here only two gave positive results with the probe.

Several animal models have been proposed to demonstrate the putative lesion caused by EAggEC strains [10] and the diarrhoea that they induce [11]. Again there appears to be disagreement. Vial et al. [10], infected rabbit and rat ileal loops with EAggEC strains and found marked histological changes, including villus shortening and necrosis of villus tips. However, in the gnotobiotic piglet whole-animal model [11] only minor changes were found, such as oedema of villi and hyperaemia. Histological changes could not be examined in the work presented in this paper as the use of fixed tissue allowed only the initial stages of bacterial adhesion to be studied. In vitro organ culture of paediatric intestinal tissue is now being used to address some of these problems, and to further evaluate proximal intestinal adhesion of EAggEC.

In summary, the use of formalin-fixed tissue is an effective technique for studying the initial stages of bacterial adhesion in the paediatric age group and makes use of an untapped source of material that would otherwise be destroyed. By means of this technique, strains of a recently described group of bacteria, EAggEC, have been shown to adhere to jejunal mucosa, suggesting that EAggEC may be important pathogens of the proximal intestine in childhood.

We thank Dr S. Knutton for supplying the normal flora E. coli strain SC13 and the Histopathology Department, Great Ormond Street Hospital, London and Mr D. Drake for supplying the mucosal specimens. This work was funded by the King’s College Hospital Joint Research Council, grant number IRC 159.

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