An in vitro System to Study Interactions between Bacteria and Epithelial Cells at the Molecular Level

By JAAP M. MIDDELDORP*† AND BERNARD WITHOLT
Department of Biochemistry, University of Groningen, Nijenborgh 16, 9747 AG Groningen, The Netherlands

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This paper describes an experimental system to study interactions between porcine enterotoxigenic Escherichia coli (ETEC) and porcine intestinal epithelial cells in vitro at the molecular level. Radiolabelled bacteria or bacterial membrane fractions were incubated with brush borders prepared from purified epithelial cells, which were then washed repeatedly. The bacterial components removed by washing or retained by the brush borders were analysed to determine their composition and source. For this it was necessary to develop a minimal medium in which attachment factors of porcine ETEC could be radiolabelled. Furthermore, an improved method for the isolation of porcine intestinal epithelial cells was developed, since other procedures did not yield sufficiently pure preparations. The resulting method was rapid and yielded large quantities of viable epithelial cells, free from crypt cells and contaminating intestinal contents. Finally, we adapted existing procedures to isolate brush borders from these epithelial cells with special emphasis on the removal of nuclear and cytosolic material and on the isolation of morphologically intact brush borders. Using this system, mixtures of bacterial cytoplasmic and outer membranes were incubated with brush borders. Cytoplasmic membranes were easily removed by washing, while the outer membranes were not.

INTRODUCTION

Enterotoxigenic strains of Escherichia coli (ETEC) cause diarrhoea in calves, piglets and man by colonizing the small intestine and transferring enterotoxins to the epithelial cells (Richards & Douglas, 1978). Interactions between ETEC and host cells are usually investigated microscopically. The numbers of bacteria which bind per host cell have been determined by light microscopy (Evans et al., 1978; Sellwood et al., 1975; Wilson & Hohmann, 1974) and the morphology of the interaction has been investigated by electron microscopy (Hohmann & Wilson, 1975; Moon et al., 1977). However, these techniques do not provide much information on the various bacterial and host components involved at the molecular level. Such information can best be obtained by studying the interactions of radioactively labelled bacteria or bacterial components with unlabelled epithelial cells or epithelial cell brush borders; bacterial components involved in adhesion should bind specifically to host cells or brush borders and should be identifiable even in the presence of large amounts of host cell material (King & Swanson, 1978).

For these experiments, it is necessary to meet two requirements. First, ETEC have to be grown in a medium in which both incorporation of radioactive amino acids and expression of bacterial attachment factors are optimal. Since the rich media usually employed to grow ETEC (Hohmann & Wilson, 1975; Jones & Rutter, 1972; Nagy et al., 1976, 1977; Shipley et al., 1978; Stirm et al., 1967; Wilson & Hohmann, 1974) are not suitable for the incorporation of

† Present address: Department of Clinical Immunology, State University Hospital, Oostersingel 59, 9713 EZ Groningen, The Netherlands.
radioactive amino acids, a minimal medium was developed which allows labelling as well as the synthesis of attachment factor. Second, it is essential to work with purified epithelial cells and brush borders to avoid non-specific binding. Procedures for the preparation of intestinal epithelial cells and brush borders have been described for a number of animals including rat (Forstner et al., 1968; Webster & Harrison, 1969), guinea-pig (Evans et al., 1971), man (Maestracci et al., 1975) and pig (Holmgren et al., 1975; Sellwood et al., 1975; Wilson & Hohmann, 1974). Although adequate for morphological investigations, these procedures were not suitable for biochemical investigations due to the binding of radioactive bacterial material to non-epithelial cells or cell nuclei. We therefore modified existing methods to develop a procedure for the isolation of purified intact epithelial cells in high yields from porcine small intestine, and a procedure for the isolation of highly purified intact brush borders from these epithelial cells.

In this paper we describe the effect of growth in minimal or rich medium on the ability of bacteria to adhere to isolated epithelial cells. We also describe in detail the modified procedures for isolating epithelial cells and purifying brush borders from these cells. Such preparations are useful tools for investigating interactions between porcine ETEC and host cells at the molecular level, as illustrated by adhesion experiments involving bacterial membrane fractions and purified brush borders.

**METHODS**

**Buffers and media.** The following buffers were used: 10 mM-sodium phosphate (pH 7-4) buffered Ringer’s mammalian solution (PBR) (Altman & Dittmer Katz, 1976); PBR supplemented with 1 mg glucose ml−1 and 5 μg trypsin inhibitor ml−1 (Sigma) (IPBR); Eagles minimal essential medium (Altman & Dittmer Katz, 1976) supplemented with 9 mM-CaCl2, 10 000 U penicillin l−1, 0-5 g streptomycin l−1 and 2-5 mg fungizon l−1 (Eagles MEM); 0-2 M-sorbitol, 12-5 mM-NaCl, 10 mM-sodium phosphate buffer (pH 7-4) (final buffer); final buffer supplemented with 0-5 mM-EDTA (isolation buffer); final buffer with 10 mM-MgCl2 (Mg2+-buffer).

**Bacteria and growth conditions.** The E. coli strains used are shown in Table 1. Most were originally isolated from pigs with diarrhoeal disease (Guinée et al., 1977); they were the kind gift of Dr P. A. M. Guinée (Rijks Instituut voor de Volksgezondheid, Bilthoven, The Netherlands). To maximize expression of adhesion factors in rich media, strains were grown for 1 or 2 weeks at 37°C in stationary tubes containing 10 ml of one of the following media (P. A. M. Guinée, personal communication): 0-8% (w/v) Nutrient Broth (Difco), Nutrient Broth supplemented with 10% (v/v) horse serum or 3-7% (w/v) Brain Heart Infusion (Difco). Bacteria were also grown to the stationary phase (16 h) on a rotary shaker at 200 r.p.m., which gave a density of about 1-2 mg dry weight ml−1 (Witholt et al., 1976) in a minimal medium containing E-salts (Vogel & Bonner, 1956), 0-5% (w/v) dextrose and 1% (w/v) Basal Medium Eagle vitamins (Flow Laboratories).

**Isolation of attachment factor K88α.** Escherichia coli strain 2100 cells were harvested from 8-4 l minimal medium and suspended in 150 ml 0-1 M-phosphate buffer (pH 7-5). Attachment factor was sheared off the cells in a Waring blender (1 min at half maximum power) and isolated by acid precipitation according to Stirm et al. (1967). The final K88α pellet was dissolved in water, dialysed against water at 0–4°C for 1 week, lyophilized to a white powder, and stored at −20°C.

Samples of purified K88α, K88αs, K88αd and K88αs/d were kindly provided by Dr F. Mooi and Dr Gaastra (University of Amsterdam, The Netherlands). Antiserum against K88α was kindly provided by Dr P. A. M. Guinée.

**Radioactive labelling and isolation of bacterial membrane fractions.** Strain 2100 was labelled continuously with [35S]methionine (28-49 Td mmol−1, Amersham) in minimal medium at 37°C or at 18°C. Harvested cells were converted to sphaeroplasts (Witholt et al., 1976) and lysed by sonication (7 periods of 15 s at 50 W 0–4°C, Branson Sonifier B12, Branson Sonic Power Co., Danbury, Conn., U.S.A.) in the presence of DNAase and RNAase (De Leij & Witholt, 1977). Large fragments were removed from the lysate by centrifugation at 5000 g for 10 min and total membranes (TM) were obtained from the supernatant fraction by centrifugation at 190 000 g for 2 h. Outer membranes and large cell fragments were removed from the lysate by centrifugation at 30 000 g for 15 min and a crude cytoplasmic membrane (CM) fraction was obtained by centrifuging the supernatant at 190 000 g for 2 h. This CM fraction contained approximately 20% outer membrane material which was not sedimented by the first centrifugation step.

**Isolation of intact epithelial cells from porcine small intestine.** Intestines were removed from one year old slaughter pigs starved for 12–18 h prior to slaughter. Immediately after death the intestines were dissected out and a 1-5 m length from the middle of each small intestine was placed on ice. The luminal contents of these segments were
were stored at -80 °C was most useful when large numbers of cells were needed for the isolation of brush borders. Added to the cell suspension to a final concentration of 15% (v/v) and small volumes, frozen in liquid nitrogen, 400 ml beaker; a large surface-to-volume ratio appeared necessary to maximize oxygen diffusion. Such cells were still 70% viable after storage for 4 h. Prolonged storage at 0-4 °C in a 300-400 ml beaker, a large surface-to-volume ratio appeared necessary to maximize oxygen diffusion. Such cells were still 70% viable after storage for 4 h. Prolonged storage at 0-4 °C renders the cells more sensitive to agitation, so for most adhesion experiments the cells were used within 1 h of isolation. For long-term storage, glycerol was added to the cell suspension to a final concentration of 15% (v/v) and small volumes, frozen in liquid nitrogen, were stored at -80 °C. The viability of such stored cells, which was high directly after thawing, decreased rapidly. Storage at -80 °C was most useful when large numbers of cells were needed for the isolation of brush borders.

Isolation of brush borders. The isolation of brush borders from rat (Forstner et al., 1968), pig (Forsyth et al., 1978; Sellwood et al., 1975) and other animals (Mooseker, 1976) has been described. Some of these procedures (Forstner et al., 1968; Sellwood et al., 1975) were unsatisfactory, since they yielded fragmented brush borders or failed to separate brush borders from the underlying cellular material (Mooseker, 1976) or from cell nuclei (Forstner et al., 1968; Forsyth et al., 1978). An improved procedure (Fig. 1) using SDS-PAGE to follow the removal of non-brush border proteins and phase-contrast microscopy to follow the morphology of the brush borders at various stages of purification was developed. Epithelial cell suspensions (106 cells ml-1) were washed twice in PBR (0 °C) and centrifuged at 200 g for 10 min (Fig. 1a). The pellet was suspended in 5 mM-NaHCO3 (pH 8.2) by repeated passage through a syringe with a 0.7 mm bore needle. The cells were gently homogenized by about 40 up and down strokes in a Dounce tissue grinder with a loosely fitting pestle (clearance 0.005 cm) until all epithelial cells were disrupted and little cytoplasmic material was left on the brush border basal side as seen by phase-contrast microscopy (Fig. 1b). To remove terminal web filamentous material from brush borders, the cell homogenate was centrifuged at 450 g for 10 min, and the pellet resuspended in isolation buffer (Dounce, 10 strokes) and centrifuged again. The brush borders were separated from underlying material by one to two washes in isolation buffer. Homogenization in hypotonic medium and low speed centrifugation easily removed most cell fragments (Fig. 1c) except nuclei (Fig. 1d). To remove these, the final pellet was resuspended in Mg2+-buffer using a Dounce tissue grinder and stored at 0-4 °C in a glass column (Fig. 1e) for 30-60 min until a substantial sediment had developed (modified from Kessler et al., 1978). The supernatant fraction was carefully removed and filtered through glass wool (1 g to 10 ml suspension) as indicated in Fig. 1(f) (from Forstner et al., 1968). The resulting filtrate was centrifuged at 450 g for 10 min and the brush border pellet was suspended in final buffer (Fig. 1g). Purity was assessed by SDS-PAGE (Fig. 3, lane 9), phase-contrast microscopy and sucrose-density gradient centrifugation analysis (a typical pattern is shown in Fig. 1h).

For long-term storage, brush border suspensions in final buffer were frozen in liquid nitrogen and kept at -80 °C. The brush borders in such suspensions remained intact and showed unchanged protein profiles for several months. Before use in adhesion tests the brush borders were washed and suspended in PBR.

Adhesion between bacteria and epithelial cells or between bacterial membrane fragments and brush borders. Adhesion experiments with intact epithelial cells were performed by a modification of the method of Svanborg-Eden et al. (1977). Epithelial cells from fresh isolates were generally used. Bacteria were suspended in PBR after harvesting at 5000 g for 10 min. Bacteria (108 in 200 l) were added to epithelial cells (105 in 200 l) and the mixture was incubated for 45 min at 37 °C. PBR was used for both incubation and washes. There was increased adhesion to

<table>
<thead>
<tr>
<th>Strain</th>
<th>Serotype</th>
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<tr>
<td>1900 (B41)*</td>
<td>O101:K-:K99</td>
</tr>
<tr>
<td>2000</td>
<td>O149:K91:K88,</td>
</tr>
<tr>
<td>2100 (G7)*</td>
<td>O8:K87:K88,</td>
</tr>
<tr>
<td>2208†</td>
<td>C600(K88,bo)</td>
</tr>
<tr>
<td>2800</td>
<td>O8:K442</td>
</tr>
<tr>
<td>2900‡</td>
<td>O9:K103</td>
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<tr>
<td>3000‡</td>
<td>O9:K103</td>
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* Designations used by Orskov et al. (1977).
† K88,bo plasmid from strain 2100 (G7) introduced into E. coli C600 by P. A. M. Guinée.
‡ A single strain was stored on solid media to select for maximal (2900) and minimal (3000) P-987 production (Nagy et al., 1977).

minimal so that a two-step wash was sufficient to remove non-mucous material. Segments of approximately 20 cm were cut immediately after removal of the intestine and flushed with ice-cold IPBR, directly followed by gentle flushing with 30 ml ice-cold IPBR per segment. The segments were transported in IPBR on ice and everted. The everted segments were dipped in fresh IPBR, and transferred to 200 ml IPBR at room temperature. Epithelial cells were separated from the underlying tissue by shaking at 100 r.p.m. for 5-10 min. After removal of the gut segments, the epithelial cells were harvested by centrifugation at 200 g for 10 min at 0 °C, gently resuspended, washed carefully in PBR (0 °C), and centrifuged again. The washed cells were gently suspended in cold Eagles MEM to a density of about 106-107 cells ml-1. When needed immediately, the cells were stored at 0-4 °C in a 300-400 ml beaker, a large surface-to-volume ratio appeared necessary to maximize oxygen diffusion. Such cells were still 70% viable after storage for 4 h. Prolonged storage at 0-4 °C rendered the cells more sensitive to agitation, so for most adhesion experiments the cells were used within 1 h of isolation. For long-term storage, glycerol was added to the cell suspension to a final concentration of 15% (v/v) and small volumes, frozen in liquid nitrogen, were stored at -80 °C. The viability of such stored cells, which was high directly after thawing, decreased rapidly. Storage at -80 °C was most useful when large numbers of cells were needed for the isolation of brush borders.

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areas where the epithelial cell membranes had been damaged ('puffs'). Therefore, only intact epithelial cells were counted in the adhesion experiments. Brush borders were incubated with radiolabelled bacterial membranes by adding 200 μl brush border suspension to 100 μl labelled membranes and shaking them in polycarbonate tubes for 45 min at 37 °C (100 cycles min⁻¹). The incubation was stopped by adding 1 ml ice-cold PBR. Brush borders were sedimented by centrifugation at 800 g for 3–5 min. The supernatant fraction was stored and the brush borders were washed with 1 ml PBR (0 °C) followed by centrifugation as above. This treatment was repeated five times. Supernatant fractions (washes) and the final washed brush border pellet were analysed by scintillation counting and sucrose-density gradient centrifugation.

Analytical procedures. Phase-contrast microscopy was used to follow the appearance of epithelial cells during their isolation, and cells were counted in a Bürker–Türk counting chamber. Phase-contrast microscopy was also used to distinguish epithelial cells, with their characteristic columnar shape and brush border region (Evans et al., 1971), from other intestinal cell types. Viability was tested by trypan blue exclusion. The removal of epithelial cells from the intestinal villi was followed by microscopy. Gut samples taken during isolation were fixed in 2.5% (v/v) glutaraldehyde and prepared for thin-section microscopy. Fixed gut samples were dehydrated, embedded in paraffin and thin sections were stained with the periodic acid–Schiff reagent to show the mucus layer.

Sucrose-density gradient analysis was performed according to De Leij et al. (1977), and radioactivity was counted as described by De Leij et al. (1979). SDS–PAGE was done according to Laemmli (1970) using 12.5%
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(w/v) acrylamide gels. Gels were stained with Fast Green. Protein was determined by the Lowry method using bovine serum albumin as standard.

Slide agglutination tests were performed by adding one drop of bacterial suspension to one drop of antiserum at 1:20 dilution on a microscope slide and agglutination was observed after 1 min. Ouchterlony double diffusion was performed as described by Isaacson (1977).

RESULTS

Isolation of porcine intestinal epithelial cells and the corresponding brush borders

Early observations indicated that epithelial cells were very easily detached from the villi. Incubation times longer than 15 min decreased the viability of the epithelial cells and caused the release of other cell types, resulting in cell preparations which contained mostly crypt and lamina propria cells. The viability of epithelial cells was increased markedly by buffering to pH 7.4 (Evans et al., 1971) and by addition of glucose and trypsin inhibitor during isolation. EDTA, as used by Evans et al. (1971), destabilized the cell membranes and was therefore not used. Despite these precautions and the speed with which the first flushing steps were carried out, epithelial cells were already detached from the lamina propria when the intestinal segments were everted, as can be seen in Fig. 2(a, b). Gentle shaking of these everted segments released the epithelial cells into the medium in large sheets (Fig. 2c) which formed small clumps of cells on subsequent washing or on agitation of the suspension. Individual cells were more easily damaged than cells still attached to one another. It was important therefore to avoid unnecessary agitation during the isolation of epithelial cells.

Suspensions prepared as described above (see Methods), contained more than 95% epithelial cells, with a viability of 80–90% in fresh isolates. The entire isolation could be completed within 30–40 min of killing the pig, which minimized damage to the epithelial cells due to autolysis. These suspensions were used either to test the effect of different growth media on the adhesion of various ETEC strains to epithelial cells, or to prepare brush borders.

To purify brush borders and to separate them from nuclei we combined a series of purification steps as illustrated diagrammatically in Fig. 1. The resulting procedure yielded open brush borders which, when compared to nuclei, accounted for 95–99% of the cell fragments seen by phase-contrast microscopy. The enrichment of two typical brush border constituents (actin and myosin with mol. wt of 42000 and 200000, respectively; Mooseker, 1976) and the corresponding decrease of nuclear protein (histones with mol. wt of 10000, 12500, 13000 and 14000; Altman & Dittmer Katz, 1976) during the brush border isolation is shown in Fig. 3. Based on the relative histone content of brush borders before (Fig. 3, lane 7) and after purification (lane 9) the final brush border suspension contained, at most, 5% nuclear material, as routinely found for more than 25 separate isolations. The brush borders isolated by this procedure remained intact, with very little contamination on the cytosolic side, as seen by phase-contrast microscopy. They banded at a density of 1.185–1.210 g cm⁻³ on sucrose gradients, corresponding to 39.3–44% (w/w) sucrose at 20 °C (Fig. 1h).

Development of a medium suitable for radiolabelling and expression of functional attachment factors

In a previous paper we described a minimal medium suitable for labelling of ETEC with radioactive amino acids (Middeldorp & Witholt, 1981). The utility of this medium for our studies depended on the synthesis of functional attachment factors. Slide agglutination tests with anti-K88ab antiserum showed that K88ab strains produced K88ab antigen in minimal as well as in rich media.

To establish the identity of the K88 antigen produced by strain 2100 in minimal medium, it was isolated to near homogeneity (90–95%) and compared to K88 variants isolated by Mooi & De Graaf (1979). SDS–PAGE and immunodiffusion against anti-K88ab antiserum identified the K88 variant produced by strain 2100 as K88ab (data not shown). To determine whether the K88ab antigen synthesized in minimal medium was expressed as a functional attachment factor, we measured the adhesion of whole bacteria to isolated epithelial cells. Strains possessing K88ab
antigen showed pronounced adhesion in minimal medium (Fig. 4, upper two strains). Although the K88as-mediated adhesion occurred over the whole epithelial cell, the bacterial density was by far the greatest on the brush border side of the cell, indicating that the K88-receptor is concentrated at the brush border region, which is the only point of contact in vivo. This was especially the case for strain 2100, which showed a complete coverage of the brush border area. Strain 2100 was also cultured at 18 °C, which prevents K88 formation (Jones & Rutter, 1972; Middeldorp & Witholt, 1981). After one growth cycle at 18 °C some adhesion to epithelial cells

Fig. 3. SDS-PAGE of samples taken during the isolation of brush borders from porcine small intestinal epithelial cells. Lanes 1, 6 and 10: reference proteins (from top to bottom: bovine serum albumin, hen ovalbumin, aldolase, chymotrypsinogen, and cytochrome c with molecular weights of 68000, 43000, 40000, 25000 and 11700, respectively). Lane 2: whole epithelial cells. Lanes 3, 4 and 5: supernatant fractions from hypotonic homogenization (3) and washing (4, 5) steps. Lane 7: material before the Mg2+-sedimentation step. Lane 8: supernatant fractions of the Mg2+-sedimentation step. Lane 9: purified brush borders after glasswool filtration. Lane 11: material remaining in the glasswool after filtration and recovered by squeezing.

Fig. 2. Thin sections of fixed gut segments, taken after transportation and eversion as described in the text. Segments were stained with the periodic acid–Schiff reagent to show the presence of the mucus layer (dark lining) and individual slime cells (S). The time elapsed after killing the pig was about 15 min. At this point epithelial cells (E) had already become detached from the lamina propria (LP) [(a) and at higher magnification (b)]. The epithelial cells became detached as large sheets of cells (a) and were found as such in the medium (c). The bar markers in (a) and (c) represent 30 μm and the bar marker in (b) represents 10 μm.
Fig. 4. Binding of various *E. coli* strains to isolated porcine intestinal epithelial cells. Six strains of *E. coli* were grown in minimal medium or in one of three rich media for one (1) or two (2) weeks and added to epithelial cells as described in Methods. MM, minimal medium; BHI, Brain Heart Infusion; NB, Nutrient Broth; NB-HS, Nutrient Broth supplemented with horse serum. The bars of each histogram show four classes of epithelial cells: \[0\], 0-9 bacteria; \[\], 10-19 bacteria; \[\], 20-29 bacteria; \[\], >30 bacteria. The number of bacteria indicates the number of organisms attached per epithelial cell.

was still evident but repeated cultivation at 18 °C completely abolished adhesion and cell agglutination with anti-K88<sub>ab</sub> serum. A K12 strain harbouring a K88-coding plasmid (strain 2200) also produced functional K88 pili; total adhesion was slightly less than for strain 2100 (Fig. 4). Other piliated strains, such as O8 : K"442" : P+ (this strain was derived from strain 2800) and a K99+ strain [O101 : K- : K99 (B41)] also showed adhesion in this system although the total number of bound bacteria was less than that with K88 strains (data not shown).

Finally, differences among adhering and non-adhering strains were clear when they were grown in minimal medium but were reduced considerably or eliminated when these same strains were grown in rich media (Fig. 4). During repeated experiments using identical conditions for bacterial growth we always found similar results with each type of K88-receptor in different isolates as has been reported by Sellwood *et al.* (1975).

**Interaction of labelled bacterial fragments and isolated brush borders**

*Escherichia coli* strain 2100 was labelled with [35S]-methionine in minimal medium, and the various isolated membrane fractions were added to purified brush borders. After incubation for 45 min at 37 °C, the brush borders were washed five times to remove non-binding material. The results for two such experiments are shown in Table 2 and Fig. 5. The total membrane fraction (TM) consisted of unseparated outer and inner membranes as shown by sucrose density gradient analysis (Fig. 5). The first wash removed nearly three-quarters of the radioactivity originally added to the brush borders; this wash preferentially removed cytoplasmic membrane material (Fig. 5), implying that the brush borders preferentially bound the outer membrane material present in the total membrane preparation. The same was true for a partially purified cytoplasmic membrane preparation (Table 2) in which case it was contaminating outer
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Fig. 5. Sucrose-density gradient centrifugation of radiolabelled E. coli total membranes before and after interaction with brush borders. ○, Total membranes before adhesion; ⋄, membranes removed in the first wash; □, material which remained bound to the brush borders after five washes; △, density. Gradient profiles were normalized to fit within the same figure. The 100% value for ○, ⋄ and □ represents 10⁵ c.p.m., 10⁴ c.p.m. and 10³ c.p.m., respectively. Repeated experiments always gave similar results, however, before each experiment the membrane composition of the total membrane was always analysed separately on sucrose-gradients in order to check for complete (OM- and CM-) membrane separation. OM, outer membrane; CM, cytoplasmic membrane.

Table 2. Binding of radiolabelled bacterial fragments to isolated brush borders

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<thead>
<tr>
<th>Bacterial fraction</th>
<th>Washes</th>
<th>Washed brush borders</th>
<th>Total recovery (%)</th>
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<tr>
<td></td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>TM</td>
<td>71-6</td>
<td>4-5</td>
<td>1-6</td>
</tr>
<tr>
<td>CM</td>
<td>76-0</td>
<td>6-4</td>
<td>1-7</td>
</tr>
</tbody>
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membrane material which remained bound to the brush borders (data not shown). This outer membrane material was bound tightly to the brush borders since it was not removed by the third, fourth and fifth washes (Table 2) and on sucrose-density gradient analysis it ran at the same density as purified brush borders (compare Fig. 5 and Fig. 1h). The same was true for the partially purified cytoplasmic membrane preparation (data not shown). Analysis of the various fractions by SDS–PAGE and fluorography established that the material which remained bound to the brush borders consisted of outer membrane fragments enriched with respect to K88ab, as has recently been described elsewhere (Middeldorp & Witholt, 1981). The results shown in Table 2 and Fig. 5 are typical of this type of experiment and show that molecular details can be studied even in the presence of large amounts of non-binding material.

DISCUSSION

In this paper we describe improved methods for the preparation of purified epithelial cell brush borders and of radioactively labelled bacterial components recently used to study the interactions of bacteria and epithelial cells in vitro and at the molecular level (Middeldorp & Witholt, 1981).
Epithelial cells were easily lost in the initial washing steps when preparation procedures described by other workers were employed. By using a few gentle washes in the presence of trypsin inhibitor to block intestinal proteases, satisfactory epithelial cell suspensions were obtained. These suspensions could then be used to prepare nearly homogeneous brush border preparations as determined by phase-contrast microscopy and SDS-PAGE.

By growing bacteria in a minimal medium, which allowed radiolabelling of membrane proteins, full expression of K88\textsubscript{ab} pili and probably other attachment factors as well was achieved. Although a minimal medium is unlikely to mimic the in vivo intestinal environment, the same might be said of rich media such as nutrient broth or brain heart infusion. Synthesis of K99 pili is blocked in the presence of L-alanine, explaining why rich media, which contain L-alanine, inhibit the synthesis of K99 (De Graaf \textit{et al.}, 1980). Synthesis of K88 also seems to be blocked in rich media (Fig. 4). Other components of rich media which block the synthesis of K88 may include trace metals, since no synthesis of functional K88 occurred when trace metals were added to the minimal medium.

Gram-negative bacteria such as \textit{E. coli} are endowed with a cytoplasmic and an outer membrane. It might be expected that attachment factors such as pili, which presumably originate in the outer membrane, attach to receptors on host cells and thus link whole cells or outer membrane fragments to such cells. When mixtures of outer and cytoplasmic membranes interact with brush borders (Table 2 and Fig. 5) it is the cytoplasmic membrane fraction which is removed by washing, presumably leaving outer membrane fragments behind. More detailed experiments which utilize the adhesion system described in this paper showed that outer membranes do indeed bind to brush borders (Middeldorp & Witholt, 1981). This binding depended on the presence of K88\textsubscript{ab} attachment factor, and was independent of how the outer membranes were prepared (Middeldorp & Witholt, 1981). Thus, outer membrane fragments which are released into the medium (Gankema \textit{et al.}, 1980; Hoekstra \textit{et al.}, 1976; Mug-Opstelten & Witholt, 1978; Wensink & Witholt, 1981), periplasmic outer membrane fragments (Gankema \textit{et al.}, 1980; Wensink \textit{et al.}, 1978) and cellular outer membrane (De Leij & Witholt, 1977) all bound to purified brush borders, while the corresponding cytoplasmic membranes, or outer membranes of non-piliated cells, did not. Since the release of outer membrane fragments by growing cells is a normal phenomenon (Gankema \textit{et al.}, 1980; Hoekstra \textit{et al.}, 1976; Mug-Opstelten & Witholt, 1978; Wensink & Witholt, 1981), and since the heat-labile enterotoxin (LT) of \textit{E. coli} is associated with the outer membrane (Gankema \textit{et al.}, 1980; Wensink \textit{et al.}, 1978), the interaction of released outer membrane fragments and host epithelial cells may be physiologically significant. Specifically, released outer membrane fragments may function as toxin carriers as we have suggested in a recent model (Middeldorp & Witholt, 1981).

The experimental system described above provides the basic tools to study such interactions in detail, and should be useful in elucidating the molecular events which occur when bacterial and eukaryotic membranes happen to encounter one another under conditions of symbiosis or pathogenesis.

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