A Preliminary Investigation on the Chemical Composition of the Cell Surface of Five Enteropathogenic *Escherichia coli* Serotypes

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The cell surfaces of five enteropathogenic *Escherichia coli* serotypes (O111:H2; O111:H12; O125:H9; O119:H6; O26:H11) were assayed by chemical methods, lectin agglutination tests and spectroscopy associated to transmission electron microscopy. Results of lectin agglutination assays showed that all strains reacted with mannose-binding lectins. Strains belonging to serotype O125:H9 also agglutinated with lectins which recognize galactose and N-acetyl-galactosamine residues. The bacterial cells were treated with 0.01M phosphate buffered saline (pH 7.0) at 100°C for 2hr and the extracts were submitted to precipitation and fractionated by Cetavlon. Phosphate, total sugar and protein contents were determined. Gas liquid chromatography-mass spectrometry analysis of alditol acetates showed the presence of galactose, mannose, fucose, glucose and traces of ribose. Spectroscopic analysis of intact cells showed the presence of a capsule-like structure which was not totally preserved after extraction. Some cells were still surrounded by an amorphous capsular-like material after polysaccharide extraction.

Key words: enteropathogenic *Escherichia coli* - cell surface - lectins

Enteropathogenic *Escherichia coli* (EPEC) is a leading cause of diarrhea in children living in developing countries, especially among infants from 6 months to 1 year old (Gomes et al. 1989, Regua et al. 1990, Albert et al. 1995, Tamura et al. 1996). Although diarrhea is often severe and protracted, EPEC virulence factors are not completely well understood (Baldwin 1998).

Bacterial adhesion to intestinal epithelial cells has been considered an important step for the pathogenicity of EPEC serotypes leading to ultrastructural and functional changes (Donnenberg et al. 1998). It has been observed that EPEC strains present multifactorial virulence mechanisms and chemical and physical properties of their surfaces are assumed to be related to pathogenicity (Orskov et al. 1977). In nature, bacteria adhere to different substrata by a mass of tangled polysaccharides fibers, a component outside the cell wall (Costerton et al. 1978). This exopolysaccharide network mediates adhesion of bacteria to various substrata, including certain epithelial surfaces of the human host (Peterson & Quie 1981). On the other hand, extracellular capsular polysaccharides of different bacteria, including *E. coli*, have been frequently associated with diseases (Muhldorfer & Hacker 1994).

The aim of the present study was to investigate and compare the chemical nature of exopolysaccharides isolated from different *E. coli* serotypes. The cellular topology of these components was also assayed by transmission electron microscopy.

**MATERIALS AND METHODS**

**Strains** - EPEC strains belonging to five different serotypes (O111:H2; O111:H12; O125:H9; O119:H6 and O26:H11) were analyzed. They were selected on the basis of the frequency of serotypes isolated during a study previously performed in our laboratory (Regua Mangia et al. 1993). For all experiments, bacteria cells were grown in brain heart infusion broth (BHI; Difco Laboratories, Detroit, Michigan, USA) for 18 hr.

**Lectin agglutination** - Agglutination tests were performed in 96-well, round bottom microtiter plates (Takatsy microtiter plates; Cooke Engineering Co., Alexandria, Virginia, USA), using a pro-
cEDURE based on that described by Wong et al. (1986) and the 17 lectins listed in Table I. For that, equal volumes (0.025 ml) of bacterial cells suspensions (2 X 10^8 bacteria/ml) and lectin solutions (0.24 to 1000 µg/ml) were mixed and incubated at room temperature or at 4°C for 1 and 2 hr, respectively. Cell agglutination was scored by visual inspection.

*Extraction and purification of the capsular polysaccharide* - Capsular polysaccharide were extracted as previously described (Lloyd 1970). Briefly, strains were grown in 250 ml of BHI and, after centrifugation, 3 vol of ethanol were added to the supernatant to precipitate the polysaccharides. Precipitates were redissolved in water and reprecipitated three times with ethanol. The crude polymers were fractionated by stepwise precipitation with hexadecyltrimethylammonium bromide (Cetavlon) at pH 7.0 (Fraction A) and Cetavlon-borate at pH 8.8 (Fraction B). Polysaccharides in Fraction B were recovered from the Cetavlon-borate complex by treatment with glacial acetic acid and reprecipitation with 3 vol of ethanol. The resulting precipitate was removed by centrifugation and washed with 2% acetic acid in ethanol and then ethanol only. Fraction C was obtained by adjusting the pH of the supernatant to pH 9.5 with 2N NaOH and the product was isolated as for Fraction B. The fractions were acidified, dialyzed, evaporated to a small volume and the resulting polymers were precipitated with excess ethanol, and then isolated. The different fractions obtained were submitted to both analytical and chemical methods.

*Analytical methods* - Total carbohydrate contents was determined by the phenol sulfuric acid assay (Dubois et al. 1956). Total protein contents was determined according to Lowry et al. (1951), and phosphate contents by the procedure recommended by Ames (1966).

*Determination of polysaccharide composition* - Fractions A, B, and C were hydrolyzed with 0.5 M sulfuric acid for 18 hr at 100°C and the resulting monosaccharides were examined as their alditol acetates by gas liquid chromatography-mass spectrometry (GC-MS) (Sawadeker et al. 1965). The chromatographic separation was carried out by using a 30 m x 0.25 mm glass capillary column containing OV-225, programmed to 50-220°C at 50°C/min, and hold. Helium was used as the carrier gas at a linear speed of 40 cm/sec.

*Ultrastructural cytochemistry* - Bacterial cells were fixed or not with 2.5% glutaraldehyde, washed with 0.1 M phosphate buffer for 60 min at room temperature and at 4°C. The washed cells were then incubated in a solution containing 100 mg/ml of cationized ferritin (Sigma Chemical Co., St. Louis, Missouri, USA) for 60 min at room temperature and at 4°C. Subsequently, they were washed, post-fixed in 1% osmium tetroxide, dehydrated in acetone and embedded in Epon (Dannon et al. 1972). The resulting thin sections were examined by transmission electron microscopy in a CEM 902 Zeiss electron microscope.

*Electron spectroscopic imaging* - A drop of concentrated bacteria (before and after extraction) was placed on a formvar coated grid. The grid was air-dried. Electron spectroscopic imaging was carried out on a CEM 902 Zeiss electron microscope operated at 80 kv with a 15 µm objective aperture and an energy less acceptance window of 20 eV for the so called zero loss filtered or elastic bright field image. Contrast tuning was achieved by altering the energy losses of the electrons used to form the microscope image. Electrons with particular energy losses were used to form the image of thick specimens, obtaining variations on contrast which helped in the observation of some structures as proposed by Bauer (1988).

**RESULTS**

Results of lectin agglutination tests are shown in Table I. All strains agglutinated in the presence of most mannose binding lectins, although high concentrations were required. By contrast, concentrations as low as 0.24 µg/ml of Bauhinia purpurea lectin agglutinated serotype 0125:H9 E. coli. Microorganisms belonging to this serotype also agglutinated in the presence of Glicine max, Wisteria floribunda and Ricinus communis I lectins, and reacted with Lotus tetragonolobus and Ulex europaeus I (fucose-binding lectins) as well. No agglutination was detected with the other lectins tested as indicated in Table I.

The chemical composition of fractions obtained from bacterial cells belonging to five different EPEC serotypes was determined by GC-MS and colorimetric methods. Analysis of the fractions showed variation in their total carbohydrate, protein and phosphate values; however, Fractions A derived from all EPEC serotypes yielded higher contents than that of the other fractions. Colorimetric analysis of Fractions A (Table II) revealed that values of total carbohydrate range from 29% to 61%, protein from 8.5% to 25% and phosphate from 5% to 8.5%. Carbohydrates were also found in Fraction B obtained from serotype 0125:H9 (63%), and in Fractions C from EPEC serotypes O125:H9 (19%) and O119:H6 (51%). Proteins were detected in Fraction B from serotype O125:H9 (28%) and in Fractions C obtained from serotypes O125:H9 and O119:H6 with contents of 61% and 57%, respectively.

GC-MS analysis showed that galactose and
mannose were the predominant monosaccharide components present in most of the serotypes studied. The crude polysaccharide preparation from serotype O125:H9 was fractionated into two components. Fraction B contained 53% of galactose, 45% of mannose and 28.5% of peptide. Mannose and galactose in a ratio of 1:1 were found in Fraction C. This fraction had a peptide content (61%) higher than that of the fraction B. Mannose, galactose and glucose in a ratio of 1:1:1 and traces of ribose and fucose were found in the capsular polysaccharide from serotype O119:H6 strain. Glucose and traces of ribose were present in serotype O26:H11. Neutral monosaccharide were not found in Fractions B and C obtained from serotypes O111:H2 and O111:H12.

Data from ultrastructural cytochemistry studies showed no labeling with cationized ferritin in bacteria collected after incubation at 37°C or 4°C either in glutaraldehyde-fixed or in nonfixed cells (Fig. 1).

Analysis of intact bacteria on coated grids showed the presence of a continuous and thick surface material outside the cell wall observed by conventional transmission electron microscopy (Fig. 2) and clearly demonstrated by using inelastic images of whole bacteria (Fig. 3). This material resembling a capsule-like structure was observed surrounding individual or colony cells. After extraction, this structure was partially lost and an amorphous capsular material remained surrounding some of the bacteria cells (Fig. 4).

DISCUSSION

Electron microscopy images of the EPEC strains studied showed a continuous structure around the bacterial cells closely resembling a capsule. Furthermore, spectroscopic images of bacteria collected after hot extraction, additionality suggested the capsular nature of the such material. E. coli capsular polysaccharides (K antigens) are arranged into several distinct serotypes and many structures of the repeating units have been reported.

### TABLE I

Agglutination of enteropathogenic *Escherichia coli* (EPEC) serotypes by lectins of various specificities

<table>
<thead>
<tr>
<th>Lectin</th>
<th>Specificity</th>
<th>EPEC serotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wheat germ agglutinin</td>
<td>β-GlcNAc; α-NeuAc</td>
<td>O111:H2</td>
</tr>
<tr>
<td>Bauhinia purpurea</td>
<td>α-GalNAc; β-GalNAc</td>
<td>O111:H12</td>
</tr>
<tr>
<td>Glycine max</td>
<td>α-GalNAc</td>
<td>O125:H9</td>
</tr>
<tr>
<td>Wisteria floribunda</td>
<td>α-GalNAc</td>
<td>O119:H6</td>
</tr>
<tr>
<td>Ricinus communis</td>
<td>β-GalNAc; D-Gal</td>
<td>O26:H11</td>
</tr>
<tr>
<td>Dolichos biflorus</td>
<td>α-GalNAc</td>
<td></td>
</tr>
<tr>
<td>Vicia villosa</td>
<td>β-GalNAc; D-Mannose</td>
<td></td>
</tr>
<tr>
<td>Jacaline</td>
<td>D-Gal</td>
<td></td>
</tr>
<tr>
<td>Concanavalin A</td>
<td>D-Mannose</td>
<td>125</td>
</tr>
<tr>
<td><em>Lens culinaris</em></td>
<td>α-D-Mannose; α-Gli; α-GluNAc</td>
<td>125</td>
</tr>
<tr>
<td><em>Canavalina bonariensis</em></td>
<td>D-Mannose</td>
<td>31</td>
</tr>
<tr>
<td><em>Lotus tetragonolobus</em></td>
<td>α-Fucose</td>
<td>250</td>
</tr>
<tr>
<td><em>Ulex europaeus I</em></td>
<td>L-Fucose</td>
<td>31</td>
</tr>
<tr>
<td><em>Phaseolus lunatus</em></td>
<td>α-GalNAc</td>
<td>250</td>
</tr>
<tr>
<td><em>Ulex europaeus II</em></td>
<td>β-GluNAc</td>
<td>31</td>
</tr>
<tr>
<td><em>Solanum tuberosum</em></td>
<td>β-GluNAc</td>
<td>250</td>
</tr>
<tr>
<td><em>Limulus polyphemus</em></td>
<td>α-NeuAc</td>
<td>31</td>
</tr>
</tbody>
</table>

a: values (in µg/ml) are relative to the minimal concentration required for agglutination.

### TABLE II

Colorimetric analysis of Fractions A obtained by Cetavlon fractionation of different enteropathogenic *Escherichia coli* (EPEC) serotypes

<table>
<thead>
<tr>
<th>Content (%)</th>
<th>EPEC serotypes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbohydrate</td>
<td>50.0</td>
</tr>
<tr>
<td>Protein</td>
<td>13.0</td>
</tr>
<tr>
<td>Phosphate</td>
<td>8.5</td>
</tr>
</tbody>
</table>
Fig. 1: thin section of enteropathogenic *Escherichia coli* (EPEC) serotype O111:H2 showing no labeling of cationized ferritin on cell surface. Fig. 2: zero loss filtered image of EPEC serotype O111:H2 before extraction showing a continuous and thick material surrounding individual and colony cells. Fig. 3: inelastic imaging (E=20 \( \Delta eV \)) of EPEC serotype O111:H2 before extraction, showing similar results with a capsule-like structure surrounding the cells. Fig. 4: inelastic imaging (E=20 \( \Delta eV \)) showing the same cells after extraction. The structure outside the bacterial cell wall was only partially preserved.
Our findings related to the detection of anionic residues and neutral sugars such as phosphate, glucose, mannose, rhamnose, galactose, ribose and fucose are in accordance with previous reports which described such residues as components of repeating units devoid or not of aminosugar, acidic sugar and labile substituents (Jann et al. 1978, Hackland et al. 1991, Hahne et al. 1991). The simultaneously detection by colorimetric and GC-MS methods of carbohydrate, protein and phosphate residues in the same fraction suggest be components of a unique structure. Bacterial glycoproteins have been demonstrated, and acceptor, only recently, as cell envelope components such as membrane-associated glycoproteins, surface-associated glycoproteins and crystalline surface layers (S-layers) as well as secreted glycoproteins and exoenzymes. Although bacterial envelope components are usually constituted of polianionic species, no labeling with cationized ferritin was observed. However, it is known that, during the dehydration steps of routine preparations for electron microscopy, the very heavily hydrated bacterial glycocalyx may collapse and the polymeric matrix may not be preserved (Chan et al. 1982). This could explain the non-observation of such cationized molecules labeling the bacterial cells surfaces. The electron spectroscopic imaging technique permits to obtain images of thick specimens without the conventional chemical routine procedures. Analysis of these images, before extraction, allowed us to observe the presence of a continuous cell surface-associated structure which was disrupted after extraction suggesting the presence of a structure similar to a bacterial envelope.

The results of the determination of monosaccharide components obtained from the extracted material are in accordance with the data obtained in the lectin agglutination tests using intact bacterial cells. These findings in association with results obtained by transmission electron microscopy, suggest that the materials analyzed are superficial components of the cell surface. Agglutination tests using lectins as probes for the identification of the terminal carbohydrate on the surface of E. coli strains suggested the presence of the following sugar units: D-galactose, D-mannose, or mannose-like structures, L-fucose and N-acetyl-D-galactosamine. It should be pointed out that high concentrations of lectins were required to induce agglutination, which may indicate that the bacterial cells investigated are not as rich in surface neutral carbohydrate as most of the eucaryotes already analyzed (Winzler 1970).

Previous results reveal that these strains carry a net negative surface charge due to acidic groups which are susceptible to trypsin and phospholipase C treatments (Regua Mangia et al. 1995). Such ionogenic surface groups are residues mainly associated with glycoproteins and glycolipid and, in bacteria, these components have been found to be associated to capsular antigens or LPS from outer membranes (Messner 1997). Electronegative charges were also detected in the present study since the major product obtained after fractionation was Fraction A, a fraction that contained acidic compounds obtained by Cetavlon precipitation, a cationic complex.

To our knowledge the presence of capsular structures have only been described in extraintestinal E. coli strains (Jann & Jann 1992). However, transmission electron micrographs images associated to physico-chemical analysis performed in the present study suggest the existence of an envelope structure surrounding individual EPEC cells.

The interaction between bacteria and host cells include many specific and non-specific effects (James 1979). Although EPEC has long been known as an important category of diarrheagenic E. coli specific virulence mechanisms, such as bundle-forming pilus and intimin, a 94 Kda outer membrane protein which play an important role in the adherence step to enterocytes, were only recently characterized (Baldwin 1998). Non-specific factors include physico-chemical properties such as surface charge due to specific chemical groups localized on the cell surface.

In summary, this study presents preliminary information on the chemical composition of the cell surface of five EPEC serotypes and on the cellular topology of the components. Additional investigation is needed in order to establish the chemical composition of EPEC capsule-like structures as well as the role played by such components during the interaction of the bacteria with host cells.

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