The importance of extracellular antigens in *Pseudomonas cepacia* infections

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Summary. A clinical isolate of *Pseudomonas cepacia* from a cystic fibrosis patient was examined for its ability to produce extracellular toxic material. The organism was grown to stationary phase in a defined medium and toxic material was isolated by ultrafiltration, ion-exchange chromatography on DEAE-Sephacel and gel-filtration chromatography on Sepharose 4B. It consisted of a surface carbohydrate antigen, lipopolysaccharide and protein, and had an LD50 (when injected intraperitoneally into mice) of 395 ± 20 µg. The toxicity appeared to be associated with the lipopolysaccharide portion of the complex, because boiling for 15 min and exposure to proteolytic enzymes had no effect on toxicity. However, saponification destroyed the toxicity of the compound. Studies employing radial immunodiffusion with the sera of mice infected with this organism demonstrated production of the complex in vivo at levels approaching those sufficient to produce death. When sublethal amounts of this complex were placed in the lungs of specific-pathogen-free rats, the lung pathology observed after 12, 24, 36 and 48 h was extensive. However, antibody generated in rabbits against this material could protect mice against the complex, as well as against challenge by the homologous organism. These data indicate that extracellular toxic material produced by *P. cepacia* may be responsible for the lethality and lung tissue destruction normally associated with an active pneumonia caused by this organism.

Introduction

*Pseudomonas cepacia*, originally characterised as the cause of soft rot of onion bulbs, is now considered an important opportunist pathogen in compromised individuals, particularly in hospitals (Burkholder, 1950). While many patients colonised by *P. cepacia* apparently suffer no ill effects, other individuals experience severe life-threatening manifestations such as pneumonia, bacteraemia and meningitis (Goldman and Klinger, 1986). Cystic fibrosis (CF) centres across the USA demonstrate that CF patients are at especially high risk of colonisation and infection (Rosenstein and Hall, 1980; Corey et al., 1984; Isles et al., 1984; Tablan et al., 1985; Thomassen et al., 1985, 1986) and are currently the largest group suffering from the consequences of *P. cepacia* exposure (Goldman and Klinger, 1986). This finding is of considerable importance because of cases of fulminating pneumonia and bacteraemia in this population (Isles et al., 1984; Thomassen et al., 1985). Treatment of *P. cepacia* infections in CF patients is particularly difficult, because the organism is resistant to a wide range of antimicrobial compounds (Goldman and Klinger, 1986).

*P. cepacia*-associated pulmonary disease in CF patients was first described after examining aminoglycoside resistance in pseudomonads isolated from sputa (Blessing et al., 1979). This study reported a 4% *P. cepacia* colonisation rate in CF patients between 1974 and 1978. By 1981, rates were reported to be as high as 18% in a study in Toronto (Isles et al., 1984). An increase in *P. cepacia* colonisation from 5-1% in 1979 to 20% in 1983 was also reported (Thomassen et al., 1986). However, these workers reported a decrease in isolation and mortality due to *P. cepacia* once patients carrying *P. cepacia* were isolated from those not colonised by the organism.

The emergence of *P. cepacia* in CF patients is particularly alarming because colonisation with
this organism is often associated with rapid clinical deterioration. For example, Isles et al. (1984) reported that patients infected with *P. cepacia* had greater impairment of pulmonary function than those infected with *P. aeruginosa*. This syndrome was characterised by high fever, severe progressive respiratory failure, leucocytosis, an elevated erythrocyte sedimentation rate and a 62% fatality rate. Upon autopsy of these patients, histological examination of the lung showed severe necrotising pneumonia. Thomassen et al. (1985) also reported that CF patients colonised with *P. aeruginosa* followed by *P. cepacia* appeared to have more serious lung involvement and a poorer prognosis than those patients colonised with *P. aeruginosa* alone. Furthermore, there was a recent report of four female patients who showed leucocytosis, high fever and mortality following *P. cepacia* colonisation (Boxerbaum and Klinger, 1984). Interestingly, this organism was isolated from the blood of all four patients. This is of considerable importance because *P. aeruginosa*, which colonises the vast majority of CF patients, generally does not cause bacteraemia in this population (McCarthy et al., 1980). Recently, Tablan et al. (1987) showed that *P. cepacia* most often affected patients with moderate or advanced CF and this was usually associated with an adverse clinical outcome.

Such cases of death and severe necrotising pneumonia led us to speculate that there is an extracellular toxic substance produced by *P. cepacia* that is responsible for lethality and extensive lung pathology. Relatively little is known about the virulence properties of *P. cepacia*. McKevitt and Woods (1984) examined 48 *P. cepacia CF isolates and found that 88% were protease-positive; 69% of these strains produced lipase, but no exotoxin A or exoenzyme S activity was detected. Lonon et al. (1987) have recently demonstrated the production of lipolytic activity against a variety of substrates by 10 CF clinical isolates of *P. cepacia*.

In this study we describe an extracellular toxic complex (ETC) produced by *P. cepacia* that appears to be responsible for the lethality and extensive pulmonary necrosis associated with lung infections caused by this organism.

**Materials and methods**

**Bacteria, bacterial virulence and media**

The strains employed in these studies were *P. cepacia* 61 g and K30-6. *P. cepacia* 61 g, from the sputum of a patient with CF, was kindly supplied by J.D. Klinger, Rainbow Babies and Children's Hospital, Cleveland, OH, USA. *P. cepacia* K30-6, also isolated from the sputum of a CF patient, was kindly supplied by C.L. Prober, The Hospital for Sick Children, Toronto, Ontario, Canada. Both strains were serotyped by a typing scheme based on whole-cell antigen agglutination (McKevitt et al., 1987). *P. cepacia* 61 g is a serotype D organism and *P. cepacia* K30-6 is serotype A. The bacteria were maintained frozen in Trypticase Soy Broth and glycerol 20% v/v at -70°C. Liquid cultures were grown in Anwar's chemically defined medium consisting of 3 mM NaCl, 12 mM (NH₄)₂SO₄, 3.2 mM MgSO₄, 1.2 mM KH₂PO₄, 0.02 mM FeSO₄, and 20 mM glucose in 50 mM 3-(N-morpholino) propanesulphonic acid (MOPS, Sigma Chemical Company, St Louis, MO, USA), pH 7.4 (Anwar et al., 1983b). Lethality was determined by injecting graded numbers of the organism intraperitoneally (i.p.) into Swiss-Webster white mice and calculating the LD₅₀ value by the method of Reed and Muench (1938).

**Purification of the ETC**

Cultures were grown in 10-L quantities of Anwar’s medium for 96 h at 37°C in a shaker adjusted to 200 rpm. The bacteria were pelleted at 17 000 g at 4°C for 60 min, and the supernatant fluids were concentrated by ultrafiltration to a volume of 100 ml on an Amicon DC-2 concentrator (Amicon Corp., Danvers, MA, USA) with a H1P30-43 hollow-fiber filter that retains substances with a mol. wt > 30 000. Remaining supernatant fluids were lyophilised. This material was dissolved in 50 ml of 0.01 M Tris-hydrochloride buffer (pH 8.0) and dialysed overnight against 16 L of the same buffer at 4°C. The dialysate was then placed on a DEAE-Sephadex (Pharmacia Fine Chemicals, Inc., Piscataway, NJ, USA) column (2.5 × 40 cm) which had been equilibrated overnight at 4°C with 0.01 M Tris-hydrochloride buffer, pH 8.0. The column was washed with two bed volumes of the starting buffer, and the absorbing material was eluted with a linear NaCl (0-1.0 M) gradient in the starting buffer. The eluant was monitored for protein at 280 nm with a flow-through 2138 Uvicord S monitor (LKB Instruments). Fractions of 10 drops each (approximately 4.2 ml) were collected. In addition, the column eluate was monitored for total hexose, uronic acids and ketodeoxyoctonate (KDO) as follows. Every fifth fraction was concentrated to dryness by lyophilisation and 1.0 ml of deionised distilled water was added to each tube. A 0.2-ml sample was assayed for uronic acid by the procedure of Blumenkrantz and Asboe-Hansen (1973), with glucoronic acid (sodium salt; Sigma) as the standard.

A 0.5-ml sample was assayed for total hexose by the anthrone procedure (Loewus, 1952). Finally, a 0.2-ml sample was assayed for KDO by the method of Osborn (1963) employing *Escherichia coli* O55:B5 LPS (Sigma) or KDO (Sigma) as the standard. The various peaks were pooled, lyophilised and dialysed against 0.01 M Tris-hydrochloride buffer (pH 8.0). Pools that were lethal for mice after i.p. injection were further purified on Sepharose.
4B in the following way. Less than 100 mg of the ETC in 8 ml of 0.01 M Tris-hydrochloride buffer (pH 8.0) was applied to an ascending-flow column (2.5 × 90 cm) of Sepharose 4B at 4°C. Fractions (100 drops) were collected (approximately 4.5 ml) and protein peaks were monitored at 280 nm and recorded. Every fifth fraction was additionally monitored for uronic acid (Blumenkrantz and Asboe-Hansen, 1973), KDO (Osborn, 1963), and total hexose (Loewus, 1952). The appropriate fractions were pooled, lyophilised, and then dialysed against the 0.01 M Tris-hydrochloride buffer (pH 8.0). The pool that was toxic for mice represented the purified ETC and was used for chemical characterisation as well as LD50 determination and pulmonary necrosis studies.

Chemical characterisation of ETC

The purified ETC was assayed for protein by the procedure of Lowry et al. (1951) with bovine serum albumin (fraction V; Sigma) as the standard, for KDO by the procedure of Osborn (1963), and for total hexose by the anthrone procedure (Loewus, 1952). In KDO assays, E. coli O55:B5 LPS (Sigma) or KDO (Sigma) was the standard. In anthrone assays, glucose and mannose (Sigma) were standards.

To determine which portion of the ETC was responsible for its toxicity, 10 mg of the material was boiled for 15 min or exposed to bovine pancreas trypsin type I (Sigma; crystallised twice, ethanol-precipitated, and substantially salt-free) 1 g/L for 30 min at 25°C. To saponify the ETC, c. 1 g/L in deionised distilled water was brought to 1 N with 10 N NaOH in a screw-capped tube and heated at 66°C for 18 h. This material was then neutralised with concentrated HCl and examined for toxicity in mice. The fatty-acid analysis of ETC was performed on fatty acid methyl esters, prepared with boron trifluoride in methanol, after acid or base hydrolysis as described by Rietschel et al. (1972). A Varian Model 3300 gas chromatography equipped with a 12-m capillary column coated with SE-30 and a flame ionisation detector was used to detect the fatty acids. The analysis was performed with temperature programming from 150°C to 250°C, increasing at a rate of 4°C/min after an initial isothermal period of 4 min. Fatty-acid standards were obtained from Supelco, Inc. Identification of each fatty acid was made by comparison of retention times with those of standards from plots of the log of the retention times versus chain length and from comparison of retention times with standards obtained with a 30-m capillary column coated with SP-2330 held at 180°C. Hydroxy fatty acids were not analysed on this column.

Virulence studies in mice

The LD50 values of the ETC were determined by injecting 1-ml amounts in 0.01 M Tris-hydrochloride buffer, pH 8.0, i.p. into Swiss-Webster mice. After observation for 72 h, dead mice were counted and LD50 values were calculated by the method of Reed and Muench (1938). Before injection of the ETC preparation into mice, it was sterilised by exposure to UV irradiation (100 μW/cm² at room temperature for 30 min). The final ETC preparation was streaked on Trypticase Soy Agar (BBL Microbiology Systems, Cockeysville, MD, USA) plates for confirmation of sterility. For the virulence studies in mice, 0.1 ml of the ETC in 0.01 M Tris buffer, pH 8.0, were injected i.p., simultaneously with the bacteria into Swiss-Webster mice weighing 20–25 g. Control animals received the bacterial dilutions along with 0.1 ml of 0.01 M Tris-HCl buffer, pH 8.0.

Quantitation of in-vivo production of the ETC and Ouchterlony analysis of DEAE-Sephadel pools I and II

Radial immunodiffusion studies were performed to quantify the ETC found in the serum of infected mice that had received 10 LD50 of P. cepacia 61 g. Fifteen mice were inoculated i.p. with P. cepacia 61 g and observed for 24–48 h. When the animals appeared to be near death, they were exsanguinated by cardiac puncture. The amount of ETC in their sera was measured by radial immunodiffusion as follows. A 0.85% agarose solution in borate buffer was heated to boiling and allowed to cool to 50°C; 750 μl of homologous antiserum against the ETC was then added to 15 ml of the agarose suspension and poured on to a 3 × 4 in. (c. 8.1 × 10 cm) glass plate and allowed to solidify. The antiserum was prepared in New Zealand white rabbits by one subcutaneous injection of 500 μg of ETC in complete Freund's adjuvant and three subcutaneous injections of 500 μg of ETC in incomplete Freund's adjuvant, each spaced a week apart. The plates were incubated for 24 h at room temperature, and the zone diameters of the precipitin reactions within the gel were measured and compared with known quantities of ETC tested in the same manner. Five mice were also inoculated i.p. with 10 LD50 values of heat-killed (boiled for 15 min) P. cepacia 61 g as a control. The rabbit serum generated against purified ETC was employed to examine the antigenic relationship between DEAE-Sephacel pool II and purified ETC by double immunodiffusion in agar gels. These gels were prepared with Iogar No. 2 (Oxoid) 1% in borate-buffered saline (0.02 M, pH 8.0).

Pseudomonas cepacia inoculation

Twenty male Sprague-Dawley rats (specific pathogen-free, 200–220 g in weight, Charles River, Canada, Inc., St Constant, Quebec) were tracheostomised under ether anaesthesia and 0.05 ml of a suspension of P. cepacia 61 g, embedded in agar beads as previously described (Cash et al., 1979), was placed in a distal bronchus in the left lobe with a bead-tipped curved needle. After inoculation, animals were housed in wire-bottom cages, and food and water were given ad libitum. Ten and 20 days after inoculation, 10 animals were exsanguinated by cardiac puncture under anaesthesia. The lungs of the remaining five animals were processed for bacterial quantitation as previously described (Cash et al., 1979).
ETC inoculation

Eight male Sprague-Dawley rats were tracheostomised under ether anaesthesia, and 0·05 ml of a sterile suspension of ETC (50 μg) was placed in a distal bronchus in the left lobe with a bead-tipped curved needle. Eight control rats were inoculated in the same manner with 0·05 ml of sterile PBS. After inoculation, animals were treated as for P. cepacia inoculation. At 12, 24, 36, 48 h after inoculation, two animals from each group were exsanguinated by cardiac puncture under anaesthesia. The lungs were prepared for histological examination as previously described (Cash et al., 1979).

Animal protection studies

Antiserum to the purified ETC was prepared in New Zealand white rabbits by one subcutaneous injection of 500 μg of ETC in complete Freund's adjuvant and three subcutaneous injections of 500 μg of ETC in incomplete Freund's adjuvant, each spaced a week apart. To test the protective ability of this antiserum against purified ETC and homologous live P. cepacia, the following protocols were followed. Five mg of purified ETC in 5 ml of 0·01 M Tris buffer, pH 8·0, was placed in a beaker with 0·5 ml of rabbit anti-ETC (IRS). Five mg of purified ETC in 5 ml of 0·01 M Tris buffer, pH 8·0, was placed in a beaker with 0·5 ml of normal rabbit serum (NRS). Both beakers were stirred at room temperature for 1 h. After this time, 500 μg of ETC + NRS and 500 μg of ETC + IRS were injected i.p. into Swiss-Webster mice.

To determine if anti-ETC antibody would protect experimental animals against live homologous P. cepacia, the following protocol was followed. P. cepacia 61 g was grown to the stationary phase (OD540 = 0·45 M Tris-hydrochloride buffer, pH 8·0) in Anwar's defined media. For every 5 ml of washed P. cepacia 61 g added to a beaker, 1·0 ml of IRS was also added. The same amount of washed bacteria was added to a beaker containing 1·0 ml of NRS for every 5 ml of bacteria. The beakers were incubated at room temperature for 1 h. Then c. 10 LD50 of P. cepacia 61 g from each beaker was injected i.p. into Swiss-Webster mice. Deaths were recorded in both groups for 5 days.

To test the ability of the carbohydrate surface antigen (pool II from the DEAE-Sepharose column) prepared from P. cepacia 61 g to protect mice against challenge with live P. cepacia, the following protocol was followed. On days 1 and 8, each of 20 Swiss-Webster mice was immunised i.p. with 100 μg of pool II from the DEAE-Sepharose column in 1 ml of phosphate buffered saline (PBS), and each of another 20 Swiss-Webster mice received 1 ml of PBS i.p. On day 15, 10 of the pool II-immunised mice and 10 of the PBS control mice were inoculated i.p. with 1 × 108 cfu of the homologous P. cepacia 61 g strain. On day 15, 10 of the pool II-immunised mice and 10 of the PBS control mice were inoculated i.p. with 1 × 108 cfu of a heterologous P. cepacia strain K30-6. Deaths were recorded in all groups for 3 days.

Results

Purification of the ETC and the non-toxic surface antigen

When 1 ml (c. 1 mg dry weight) of the ultrafiltration concentrated supernate from 10 L was injected i.p. into 10 Swiss-Webster mice, all 10 animals were dead within 48 h. Control animals (10) that received uninoculated media treated in the same manner suffered no ill effects, indicating that some form of toxic material was elaborated by P. cepacia 61 g.

The elution profile of the extracellular material produced by P. cepacia 61 g on DEAE-Sepharose can be seen in fig. 1. This material could be separated into two distinct uronic acid-containing pools (OD540) by DEAE-Sepharose chromatography. Ouchterlony analysis with rabbit antiserum produced against purified ETC (DEAE-Sepharose pool I) gave a line of identity between DEAE-Sepharose pool I and II, with pool I showing a spur that was not present in pool II (data not shown). Pool I represented the material that bound to the ion-exchange resin and eluted between 0·10 and 0·45 M NaCl. Only pool I contained KDO-positive material (OD548) and was toxic for Swiss-Webster mice. Because DEAE-Sepharose pool II was not toxic for mice and appeared to be mostly carbohydrate, it was referred to as the carbohydrate surface antigen.

When animals were inoculated i.p. with 1 ml of pool I (c. 0·7 mg, dry weight), all mice died in 48 h. Pool I was lyophilised, dialysed to remove salt, dissolved in 8 ml of 0·01 M Tris-hydrochloride buffer, pH 8·0, and applied to an upward flow Sepharose 4B column (2·5 × 90 cm). The eluate was monitored as was done for the DEAE-Sepharose column. The elution profile of DEAE-Sepharose pool I on Sepharose 4B is shown in fig. 2. This material could be fractionated by Sepharose 4B into two distinct pools on the basis of absorption at 280 nm. When material from these two pools was injected i.p. into Swiss-Webster mice, only pool-I material (void volume material) was toxic. The purified ETC was found to be approximately 3% protein. The LD50 value for this material injected i.p. into 20–25 g Swiss-Webster mice was calculated to be 395 ± 20 μg. The average time to death for ETC injected animals was 36 h.

Presence of KDO in biologically active LPS preparations from P. cepacia

We have used the method of Osborn (1963) to assay culture supernate from P. cepacia 61 g and
EXTRACELLULAR ANTIGENS IN P. CEPACIA

Fig. 1. Elution profile of the ultrafiltration-concentrated extracellular material from P. cepacia 61 g (10 L stationary phase culture) on DEAE-Sephacel. The peaks were eluted with an increasing NaCl gradient (---) from 0 to 1.0 M in Tris-hydrochloride buffer (pH 8.0). The eluate was continuously monitored for protein by absorbance (OD) at 280 nm (○) and every fifth tube was assayed for uronic acid (○), KDO (△), and total hexose (△).

nine other P. cepacia strains grown to stationary phase in Anwar's defined medium for the presence of KDO. Eight of the ten strains showed no evidence of KDO (data not shown). However, the material from P. cepacia 61 g and one other strain gave a distinctly positive thiobarbituric acid reaction (Osborn, 1963). Spectral analysis indicated that this material absorbed strongly at 549 nm and was spectrally identical to the purified KDO used as a standard.

Ability of antibodies to ETC to neutralise ETC toxicity and passively protect against P. cepacia challenge

Antibody to the ETC can neutralise its toxicity in vivo since antibody to the ETC protected all the experimental animals from death, while all the animals that received ETC+NRS died (table I). In addition, we sought to determine whether antibodies to the ETC would protect experimental
Table I. Ability of antibodies to ETC to either neutralise the toxicity of the ETC or protect against live *P. cepacia* challenge or both

<table>
<thead>
<tr>
<th>Source of serum</th>
<th>Serum incubation</th>
<th>Source of ETC (500 µg/mouse)*</th>
<th>No. of mice dead/no. of mice inoculated (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rabbit anti-<em>P. cepacia</em> 61 g ETC (IRS)</td>
<td>5 mg ETC incubated with 1-0 ml of IRS for 1 h at 25 °C</td>
<td><em>P. cepacia</em> 61 g</td>
<td>0/10 (0%)†</td>
</tr>
<tr>
<td>Normal rabbit serum (NRS)</td>
<td>5 mg ETC incubated with 1-0 ml of NRS for 1 h at 25 °C</td>
<td><em>P. cepacia</em> 61 g</td>
<td>10/10 (100%)</td>
</tr>
<tr>
<td>Rabbit anti-<em>P. cepacia</em> 61 g ETC (IRS)</td>
<td><em>P. cepacia</em> 61 g (10^10 cfu) incubated with 1-0 ml of IRS for 1 h at 25°C‡</td>
<td>...</td>
<td>1/10 (10%)†</td>
</tr>
<tr>
<td>Normal rabbit serum (NRS)</td>
<td><em>P. cepacia</em> 61 g (10^10 cfu) incubated with 1-0 ml of NRS for 1 h at 25°C‡</td>
<td>...</td>
<td>10/10 (100%)</td>
</tr>
</tbody>
</table>

*Purified as described in Materials and methods.
†Significantly different from control values by Fisher's Exact test (Fisher, 1973) (p < 0.005).
‡Immediately after serum incubation, these mice received 10^9 cfu of *P. cepacia*.

Animals against infection with homologous *P. cepacia*. The results of these studies can also be seen in table I. This table shows that antibody to the ETC can passively protect mice against homologous challenge when the challenge organisms are delivered in combination with antibody; 90% of the animals that received 10^9 *P. cepacia* plus anti-ETC antibody survived, whereas all the animals that received 10^9 *P. cepacia* + NRS died.

**Ability of immunisation by DEAE-Sephacel pool II to protect experimental animals against homologous and heterologous *P. cepacia* challenge**

Table II shows that mice immunised against the DEAE-Sephacel pool II of *P. cepacia* 61 g were protected (90% survival) against 1 x 10^9 cfu of *P. cepacia* 61 g. Also, mice immunised against the DEAE-Sephacel pool II of *P. cepacia* 61 g were protected (80% survival) against 1 x 10^8 cfu of *P. cepacia* K30-6, a heterologous strain. Control mice that were not immunised all succumbed to 1 x 10^9 cfu of *P. cepacia* 61 g or *P. cepacia* K30-6.

**Bacteriology and pathology of lungs of rats inoculated with *P. cepacia***

In rats inoculated transtracheally with agar beads containing c. 10^4 cfu of *P. cepacia* 61 g, the number of recoverable organisms rose from the initial inoculum size to 3.7 x 10^5 (SD 2.1 x 10^4) cfu/lung at day 20. The histopathological picture seen in lungs infected with agar beads containing *P. cepacia* 61 g at day 10 was essentially identical to that seen at day 20. Microscopic examination demonstrated both bronchial and parenchymal changes (fig. 3). Purulent exudate and fibrin were present in bronchioles and terminal airways. Dense inflammatory cell infiltrate accumulated in the alveolar spaces and areas of frank necrosis were present.

**Pathology of ETC-inoculated lungs**

Microscopic examination of lungs inoculated with ETC demonstrated a progressively increasing accumulation of polymorphonuclear leucocytes in bronchial and alveolar lumina as well as in alveolar septae from 12 to 36 h (fig. 4). Injury and necrosis of bronchial epithelium, associated haemorrhage and fibrinous exudate was also noted.

**Fatty-acid analysis and determination of the component of ETC responsible for toxicity**

As the next step in the characterisation of the ETC, we performed experiments to determine which of the three components (LPS, carbohydrate surface antigen, or protein) present in the ETC was responsible for the observed animal toxicity. Boiling the ETC for 15 min in 0.01 M Tris-hydrochloride (pH 8.0), or exposure to trypsin (1 g/L) did not alter the toxicity of the ETC. After saponification, however, the ETC was no longer toxic as indicated by i.p. injection into experimental animals. Thus, it appeared that the LPS portion of the ETC was required for the toxicity to be expressed in mice.
Table II. Ability of immunisation by DEAE-Sephaec pool II (carbohydrate surface antigen) to protect experimental animals against homologous and heterologous P. cepacia challenge

<table>
<thead>
<tr>
<th>Source of carbohydrate surface antigen*</th>
<th>Challenge organisms‡</th>
<th>No. of mice dead/no. inoculated (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P. cepacia 61 g</td>
<td>P. cepacia 61 g</td>
<td>1/10 (10%)§</td>
</tr>
<tr>
<td>None (control)†</td>
<td>P. cepacia 61 g</td>
<td>10/10 (100%)</td>
</tr>
<tr>
<td>P. cepacia 61 g</td>
<td>P. cepacia K30-6</td>
<td>2/10 (20%)§</td>
</tr>
<tr>
<td>None (control)†</td>
<td>P. cepacia K30-6</td>
<td>10/10 (100%)</td>
</tr>
</tbody>
</table>

*Purified as described in Materials and methods. All immunised mice received 100 µg of P. cepacia 61 g DEAE-S pool II delivered i.p. in 1 ml of sterile PBS on days 1 and 8.
†Control mice received 1 ml of sterile PBS delivered i.p. on days 1 and 8.
‡On day 15, a 1 x 10⁸ cfu challenge of the appropriate organism was injected i.p. into these mice in 1 ml of PBS.
§Significantly different from control values by Fisher's Exact Test (Fisher, 1973) (p<0.005).

Because the lipid portion of the LPS had been shown to be responsible for its toxicity, a fatty-acid analysis of the ETC of P. cepacia 61 g was attempted by gas chromatography of methyl esters released by acid or base hydrolysis. As can be seen in table III, the major species were hexadecanoic acid and 3-hydroxytetradecanoic acid which, along with Δ²-tetradecanoic acid, accounted for 60% of the fatty acids by weight. The Δ²-tetradecanoic acid was probably derived from 3-hydroxytetradecanoic acid by dehydration during base hydrolysis.

LD50 value of P. cepacia 61 g and quantitation of in-vivo production of the ETC

The LD50 value of P. cepacia 61 g when performed as described in the Materials and methods in 20–25 g Swiss-Webster mice was 2.89 x 10⁸ cfu. Radial immunodiffusion was performed to quantify the ETC found in the sera of mice infected with 10 LD50 doses of P. cepacia 61 g (table IV). All mice had detectable ETC in their blood when they were killed. Mice had an average of 180 µg of ETC per ml of serum 25 h after inoculation. ETC values were in the range 50–325 µg/ml of serum. Mice that received 10 LD50 values of heat-killed P. cepacia 61 g i.p. did not die of their injection. After 48 h, 

Table IV. Quantitation of P. cepacia 61 g ETC in mouse sera by radial immunodiffusion*

<table>
<thead>
<tr>
<th>Mouse† no.</th>
<th>Time after inoculation (h)</th>
<th>ETC (µg/ml)</th>
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<tr>
<td>1</td>
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*Fatty acids released by base hydrolysis accounted for 98% of the total fatty acids present. No additional fatty acid was present in quantities >1% of the total.
†Data are reported as mean (SD), n = 3.

*Samples (10 µl) of serum were placed in each well, and plates were incubated for 24 h at room temperature.
†Mice received 10 LD50 of exponential phase P. cepacia 61 g cells i.p.
they were killed and we were not able to detect any circulating ETC in their blood.

**Virulence enhancement in mice**

Experiments were performed to determine if the ETC from *P. cepacia* 61 g could enhance the virulence of the homologous organism. The presence of 200, 100, or 50 μg of ETC in the peritoneal cavities of mice in concert with viable *P. cepacia* 61 g lowered the LD50 166-10-, and 4.6-fold, respectively.

**Discussion**

ETC produced by *P. cepacia* 61 g appeared to be composed of LPS, a carbohydrate surface antigen and protein and is probably released into the external environment by actively growing cells. Its component parts were not separable by ultrafiltration, ion-exchange (fig. 1), or gel filtration (fig. 2) chromatography. These results are reminiscent of those obtained with a similar compound produced by *Klebsiella pneumoniae* (Straus et al., 1985; Straus, 1987). The ETC of *P. cepacia* also appeared to contain a uronic acid which is also found in the carbohydrate capsule of *K. pneumoniae* (Straus et al., 1985; Straus, 1987). The significance of this is not clear.

The toxicity is thought to be associated with the LPS moiety, because boiling and protease treatment had no deleterious effect, and only saponification rendered the ETC non-toxic. However, the possibility that the toxicity may be associated with a protein has not been ruled out. It is not clear how the toxicity of the LPS molecule is expressed. When endotoxin enters the blood stream of an experimental animal, the resultant prompt and transient hypertensive state is followed by a progressive severe hypotension (Berry et al., 1982). The availability of blood to important tissues decreases, and death is thought to occur as a result of circulatory collapse. The action of LPS that produces these circulatory changes has not been fully elucidated.

Manniello et al. (1979) reported that there was no detectable KDO in biologically active LPS extracts from two strains of *P. cepacia* subjected to assay by the method of Osborn (1963). This finding was supported by Anwar et al. (1983a) in the isolation and characterisation of the outer membrane of *P. cepacia*. We have used the Osborn method to assay the culture medium of *P. cepacia* 61 g grown to stationary phase in Anwar’s defined medium and have detected material which we believe to be KDO. This material which resulted in a positive thiobarbituric acid reaction (Osborn, 1963), absorbed strongly at 549 nm. It appeared to be spectrally identical to the purified KDO used as a standard. This evidence coupled with the biological activity indicative of endotoxin led us to believe that the LPS of *P. cepacia* 61 g does in fact contain KDO. However, it should be noted that we examined the culture supernates from a total of ten *P. cepacia* strains for KDO (data not shown) and were able to detect KDO from only two strains. Therefore, it appears that while some strains of *P. cepacia* are capable of producing KDO, the majority of them probably are not.

Analysis of the lipid portion of ETC, the membrane anchoring site of the complex showed the predominance of 3-hydroxytetradecanoic acid and hexadecanoic acid with Δ⁹-hexadecanoic acid and cis-9, 10-methylene-hexadecanoic acid also present in significant amounts (table III). The unknown fatty acid chromatographed at a position corresponding to a methyleneheptadecanoic acid although it might also be a member of the octadecanoic acid series. The protein portion of the ETC probably plays no role in its toxicity, since treating ETC with trypsin did not effect its ability to kill mice. The role that the carbohydrate surface antigen plays in the toxicity of the ETC is not clear. We have not obtained the LPS moiety free of the carbohydrate surface antigen (fig. 1, pool I). The carbohydrate surface antigen free of the LPS (fig. 1, pool II) is not toxic in mice. The presence of the carbohydrate surface antigen in the ETC may enhance the toxicity of the LPS moiety by making it more difficult for the animal to excrete it from the body, for example in urine. Compounds such as individual LPS molecules of mol. wt <20 000 are secreted by the glomerular filtration system with relative ease. The aggregate size of LPS molecules would be hard to predict, especially *in vivo*. However, a compound with a molecular mass in excess of 2×10⁶ (fig. 2) would tend to lodge in the glomerular filtration system, thus setting up a gradient in the blood stream, and maintain the amount of the LPS-containing ETC at very high levels.

The ETC was shown to be lethal when injected i.p. into Swiss-Webster mice in sufficient quantities (395 μg or greater) and sublethal amounts of the ETC enhanced the virulence of the homologous organism. The ETC described here also produced extensive pulmonary pathology in the lungs of specific-pathogen free rats when placed there in the absence of live organisms (fig. 4). The damage produced by sublethal doses of the ETC placed transtracheally in the lungs of rats resulted in a
Fig. 3. Photomicrographs of hematoxylin and eosin-stained sections of rat lungs infected with *P. cepacia* 61 g (bar = 100 μm). (A) Agar bead containing *P. cepacia* lying in bronchial epithelium; suppurative inflammation obscures surrounding area. (B) Higher magnification of (A) demonstrating agar bead rimmed by polymorphonuclear leucocytes. (C) Purulent exudation and fibrin associated with airways and alveolar spaces 20 days after inoculation with *P. cepacia*. 
Fig. 4. Photomicrographs of hematoxylin and eosin-stained section of rat lungs inoculated with ETC (bar = 100 μm). (A) Injury and necrosis of bronchial epithelium and associated suppuration surrounding airway seen 12 h after inoculation. (B) Progressive parenchymal involvement associated with increasing accumulation of polymorphonuclear leucocytes in bronchial and alveolar lumina 24 h after inoculation. (C) Frank necrosis seen 36 h after inoculation, with airway and alveolar consolidation.
clinical picture even more devastating than the pathology produced by an active *P. cepacia* lung infection (fig. 3). This is in agreement with the observation that *P. cepacia* is relatively avirulent (Lonon *et al*., 1987) in normal animals. Immunisation with the ETC of *P. cepacia* 61 g elicited the production of mouse protective antibodies in rabbits. These antibodies neutralised the toxicity of the ETC and protected mice (probably by opsonisation) against challenge with the homologous organism (table I). Immunisation with another extracellular antigen (pool II from DEAE-Sephacel) protected mice against the homologous as well as a heterologous organism. This finding may have great significance if a non-toxic *P. cepacia* antigen is found to induce protection against various heterologous strains. We are currently conducting experiments to determine the extent of this protection.

All 15 mice infected with *P. cepacia* 61 g possessed circulating ETC. Although the values reported here are below the 395 μg LD50 for i.p. injection in mice, it should be noted that there are approximately 2 ml of blood in the average mouse and we did not attempt to quantify the ETC in the tissues. Therefore, had they not been killed, it seems likely that there would eventually have been sufficient ETC in the bodies of these animals to result in their deaths. None of the mice receiving 10 LD50 values of heat-killed *P. cepacia* 61 g had detectable levels of ETC in their circulatory system. It would appear then that the ETC detected in the blood of mice that received live *P. cepacia* 61 g is synthesised *in vivo* and is not only present in the bacteria themselves. We therefore propose that the ETC produced by *P. cepacia* is responsible for the lethality and extensive pulmonary tissue necrosis associated with pneumonia produced by this organism.

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