Polyagglutinating and non-typable strains of *Pseudomonas aeruginosa* in cystic fibrosis

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Summary. Serologically polyagglutinating (PA) and non-typable (NT) strains of *Pseudomonas aeruginosa* are frequently isolated from cystic fibrosis (CF) patients, but are uncommon in other patients. From serologically typical parent strains, we isolated two variants (one PA, the other NT) which differed from the parent in bacteriophage susceptibility or in sensitivity to the bactericidal action of normal human serum. The PA and NT variants (strains 7/1 and 18S respectively) reacted with antiserum to the parent strains 7 and 18R but did not absorb homologous specific O antibody from antiserum to the parent strains. In contrast the parent strains absorbed anti-PA and anti-NT antibodies from antisera to the variant strains. The yield of lipopolysaccharide (LPS) from acetone-dried cells of the parent strain 7 was similar to that of the PA derivative; but the NT strain 18S yielded only half the LPS of its parent strain. LPS of the variant 7/1 gave a banding profile by SDS-PAGE similar to that of the parent LPS 7, but lacked high-molecular-weight components. LPS of the variant 18S appeared to be grossly different in profile from LPS 18R. Of 533 isolates of *P. aeruginosa* that were tested with O antisera and with antisera to the two variants, 15% were O-typable and 22% were O-non-typable; 26% reacted with anti-PA serum alone, 10% with anti-NT serum alone, and 27% were agglutinated by both sera. There was a statistically significant correlation between serum sensitivity of CF isolates and their reaction with the PA or NT antisera.

Introduction

*Pseudomonas aeruginosa* is serologically heterogeneous and at least 17 lipopolysaccharide (LPS) type-specific O antigens have been described (Liu et al., 1983). The detection of these antigens by agglutination forms the basis of most O-serotyping procedures, and approximately 93% of clinical hospital isolates can be allocated to a specific O type (Pitt, 1981). However a recent study by Penketh et al. (1983) of 49 patients with cystic fibrosis (CF) who had respiratory infections with *P. aeruginosa*, showed that a specific O serotype was detected in only 32% of 109 isolates from these patients. Approximately half (55%) of the cultures were agglutinated by three or more O antisera and were termed polyagglutinable (PA) and 9% were nontypable (NT), i.e., they did not react with any of the O antisera. Furthermore, PA and NT cultures were often sensitive to the bactericidal action of fresh normal human serum. Hancock et al. (1983) also found that the majority of 26 isolates from CF sputum were serologically PA or NT and that LPS of these cultures was deficient in oligosaccharide O-specific side chains.

We set out to investigate whether the serological anomalies observed in the previous study were due to changes in the LPS of *P. aeruginosa*, and to characterise the serological determinants of PA and NT strains in CF. This involved: (1) the isolation of two serological variants (one PA, the other NT) from two O-typable strains; (2) a comparison of their serological properties with the parent strains; and (3) the use of antisera against the variants for the detection of serologically atypical cultures from CF sputum.

Materials and methods

Strains of *Pseudomonas aeruginosa*

The origin and properties of the four experimental strains are given in table I. The 17 O-serotype strains of the International Antigenic Typing Scheme (IATS) for the production of O-typing sera were described by Liu et al. (1983), and O-factor strains O2b and O5d (Véron, 1961) were used to subdivide IATS serogroups O2 and O5. From 115 CF patients attending an out-patients'
clinics, 553 cultures were isolated. Sputum was digested with pancreatin (Rawlins, 1953), and cultured on Pseudomonas Isolation Agar (Difco) and King's 'A' agar (King et al., 1954). Cultures were identified as P. aeruginosa by their production of oxidase enzyme and pyocyanin, and their colonial appearance on King's 'A' agar.

**Media**

Tryptone Soy Broth (Oxoid) and Tryptone Soy Agar (Oxoid) were used throughout for liquid and solid culture respectively.

**Bacteriophages**

The 20 phages of the routine phage-typing set were as described by Asheshov (1974), and the LPS specific phages E79 and PLS1 (Jarrell and Kropinski, 1976 and 1977) were obtained from Dr A. M. Kropinski, Queen's University, Kingston, Ontario, Canada. Phage-resistant variants of strains were selected by the procedure of Bradley and Pitt (1974).

**Pyocin typing**

Cultures were typed by the method of Govan (1978).

**Preparation of antiserum.**

The methods described by Pitt and Erdman (1978) were used for the preparation of O-antisera, for agglutination tests and for the absorption of antibody from serum.

**Serum sensitivity test**

Cultures were tested for sensitivity to the bactericidal action of normal human serum by the method of Penketh et al. (1983).

**Isolation of lipopolysaccharide (LPS)**

LPS was isolated from whole acetone-dried cells by the method of Darveau and Hancock (1983). Total carbohydrate was measured by the anthrone reaction (Trevelyan and Harrison, 1952), total protein by the method of Lowry et al. (1951), and 2-keto-3-deoxyoctonate (KDO) by the procedure of Osborn (1963).

**SDS polyacrylamide gel electrophoresis (SDS-PAGE)**

This was performed by the method of Palva and Mäkelä (1980) with a 12.5% acrylamide resolving gel. LPS (10 μl containing 5 mg/ml) was applied to each lane. Protein bands were stained with Coomassie brilliant blue 0.1% w/v in acetic acid 10% v/v and methanol 25% v/v. Gels were destained with the above solution without the stain. Carbohydrates were detected by the silver stain of Tsai and Frasch (1982). Gels were calibrated with an Electrophoresis Calibration Kit (Pharmacia) consisting of standard proteins with mol. wt in the range 14 400–94 000.

**Indirect haemaggulination**

Sera were inactivated at 56°C for 30 min and absorbed with thrice-washed sheep red blood cells (SRBC) by adding 100 μl of serum to 0.9 ml of SRBC suspension 10% v/v. LPS (50 μl containing 1 mg/ml) was added to 0.3 ml of saline and 0.15 ml of 10% SRBC and the mixture was incubated at 37°C for 1 h with shaking in a water bath. The sensitised cells were washed thrice in saline by centrifugation at 1000 g for 10 min and resuspended in 5 ml of phosphate buffered saline (PBS) 0.02 M, pH 7.5, containing bovine serum albumin (Armour Pharmaceutical Co. Ltd, Eastbourne, Sussex) 0.1% w/v. Control mixtures not containing antigen were included in each batch of tests. Sera were diluted serially in PBS-albumin by two-fold dilutions of 25 μl in 'U'-well microtitration trays (Dynatech), and 25-μl volumes of sensitised SRBC were added to each of the serum and control wells. The trays were shaken gently and incubated first for 2 h at 37°C and then overnight at 4°C.

**Results**

**Selection of variant strains**

The variant strain 7/1 was selected from the reference phage propagating strain 7 by its resistance to phage 7. Strain 7/1 was classified as PA because it was strongly agglutinated by seven O-typing antisera (01, 03, 05d, 06, 09, 010 and 013); the parent strain 7 was agglutinated by serum 06 alone. Strain 7/1 was sensitive to the bactericidal action of normal human serum, in contrast to the parent which was serum resistant. The pyocin type of both strains was identical; but the variant was not lysed by any of the typing phages and was insensitive to the LPS-specific phages E79 and PLS1 (table 1).

The other variant strain (18S) was isolated from the same sputum specimen of a CF patient as a serum-resistant culture (18R) which was of serotype 05d. Strain 18S was fully serum-sensitive but was not agglutinated by any of the O-typing sera. It was assumed that 18S was derived from 18R because, in serial subculture on agar, approximately 1 in 20 colonies reverted to serum resistance and expressed O antigen 5d. Also, the opposite conversion of 18R to 18S occurred at a similar frequency. The pyocin type of 18R and 18S was identical, and their phage-typing patterns were similar although only 18S was...
Table I. Characteristics of reference strains of *P. aeruginosa*

<table>
<thead>
<tr>
<th>Strain no.</th>
<th>Description</th>
<th>Serum sensitivity</th>
<th>O-serotype</th>
<th>Pyocin type</th>
<th>Phage-type pattern</th>
<th>Lysis by LPS-specific phage</th>
</tr>
</thead>
<tbody>
<tr>
<td>7</td>
<td>Propagating strain for phage 7</td>
<td>R</td>
<td>6</td>
<td>22/a</td>
<td>7 44 119×</td>
<td>Col.11</td>
</tr>
<tr>
<td>7/1</td>
<td>Variant of strain 7 (resistant to phage 7)</td>
<td>S</td>
<td>PA</td>
<td>22/a</td>
<td>NT</td>
<td>- -</td>
</tr>
<tr>
<td>18R</td>
<td>Isolate from sputum of patient with CF</td>
<td>R</td>
<td>5d</td>
<td>1/c</td>
<td>7 F8 109 352 1214</td>
<td>+ -</td>
</tr>
<tr>
<td>18S</td>
<td>Variant isolated from same sputum specimen as 18R</td>
<td>S</td>
<td>NT</td>
<td>1/c</td>
<td>7 44 F8 109 352 1214</td>
<td>+ -</td>
</tr>
</tbody>
</table>

R = resistant, S = sensitive, to bactericidal action of normal human serum; PA = polyagglutinable (with antisera O1, O3, O5d, O6, O9, O10 and O13); NT = non-typable; LPS = lipopolysaccharide.

Sensitive to phage 44. Both strains were lysed by the LPS-specific phage E79, but not by PLS1.

The four strains produced pyocyanin and fluorescein, and gave typical colonies on agar; no differences in biochemical activity were observed between the parent and variant strains. The variants grew less well in overnight broth culture than the parents (viable count 10-fold less), and broth cultures of the variants became granular after heating at 100°C.

**Agglutination reactions of variant strains**

Antisera were raised against boiled cells of the four strains, and were tested by slide agglutination with each of the strains. Table II shows that serum to each of the parent strains contained agglutinin only to the homologous vaccine strain and its variant. The heterologous titres of the four antisera were at least 4-fold lower than their homologous reactions in repeated tests. Anti-18S serum cross reacted with each of the heterologous strains to the same titre, but the agglutination reactions were incomplete—a proportion of cells were not agglutinated and remained in suspension even after mixing for 5 min on the slide. The titres of unabsorbed and absorbed sera were reproducible; values did not vary more than two-fold in tests performed on two separate occasions.

Absorption of anti-7 serum with the homologous vaccine strain. These results indicated that the variant antigens were present in the parent strains, but the latter did not induce agglutinating antibody to the variant antigens as efficiently as did the variant strains.

The O-antigen specificity of the four antisera, diluted to one-quarter of their homologous titres, was determined by agglutination tests with the reference O-type strains. Anti-7 serum reacted only with strain O6, which was the serotype of strain 7.

Table II. Agglutination of parent and variant strains of *P. aeruginosa* by homologous and heterologous O-antisera before and after absorption with heterologous strains

<table>
<thead>
<tr>
<th>Antiserum to strain</th>
<th>Absorbed with suspension of</th>
<th>Agglutination titre* with strain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7</td>
<td>7/1</td>
</tr>
<tr>
<td>7 (parent)</td>
<td>nil</td>
<td>160</td>
</tr>
<tr>
<td>7/1</td>
<td>70</td>
<td>-</td>
</tr>
<tr>
<td>7/1 (variant)</td>
<td>nil</td>
<td>20</td>
</tr>
<tr>
<td>7</td>
<td>7</td>
<td>-</td>
</tr>
<tr>
<td>18R (parent)</td>
<td>nil</td>
<td>-</td>
</tr>
<tr>
<td>18S</td>
<td>18S</td>
<td>...</td>
</tr>
<tr>
<td>18S (variant)</td>
<td>nil</td>
<td>20</td>
</tr>
<tr>
<td>18R</td>
<td>18R</td>
<td>...</td>
</tr>
</tbody>
</table>

* = <10; ... = not done.

* Tested by slide agglutination. In repeated tests, the titres did not vary more than 2-fold from those stated (see Results).
Table III. Percentage yield of lipopolysaccharide (LPS); and total carbohydrate, protein and KDO content of LPS

<table>
<thead>
<tr>
<th>Strain</th>
<th>Yield of LPS*</th>
<th>Analysis of LPS</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Protein†</td>
<td>Carbohydrate†</td>
<td>KDO‡</td>
</tr>
<tr>
<td>7</td>
<td>10.5</td>
<td>3.8</td>
<td>68</td>
<td>2.7</td>
</tr>
<tr>
<td>7/1</td>
<td>9.5</td>
<td>3.3</td>
<td>44</td>
<td>3.6</td>
</tr>
<tr>
<td>18R</td>
<td>8.5</td>
<td>1.4</td>
<td>44</td>
<td>2.4</td>
</tr>
<tr>
<td>18S</td>
<td>4.5</td>
<td>1.8</td>
<td>38</td>
<td>3.2</td>
</tr>
</tbody>
</table>

KDO = 2-keto-3-deoxyoctonate.
* Percentage of dry weight of acetone-dried whole cells.
† Percentage of purified LPS.
‡ Percentage of total carbohydrate.

Anti-7/1 serum gave weak reactions with type strains O15 and O17 but did not agglutinate any of the other type strains. Both anti-18R and anti-18S sera reacted with strain O5d (O serotype of 18R), and showed cross reactions with strains O2 and O16 which are known to be serologically related to type O5.

Lipoplysaccharide of variant strains

To identify the basis of the change in the expression of heat-stable surface antigens, the LPS of each strain was characterised. The yields of LPS expressed as a percentage of the dry weight of acetone-dried whole cells are given in table III. Strains 7 and 7/1 contained similar amounts of LPS, but strain 18S yielded 47% less LPS than the parent strain 18R. Protein contamination was minimal; values were in the range 1.4% (for 18R) to 3.8% (for strain 7). The total carbohydrate of the two variant strains was similar (38% and 44%) but there was a considerable difference in yield between the parent strains (44% and 68%). The carbohydrate content of LPS 7/1 was approximately two-thirds of LPS 7, but the difference between LPS 18R and LPS 18S was only 6%. The percentage KDO content of the carbohydrate of the variant 7/1 was higher than that of the parent strain, but 18S had more KDO than 18R.

A variety of bands was resolved by SDS-PAGE (see figure). A densely stained zone, apparently consisting of three bands, was given by LPS 7 (lane 2) in the region of the gel which corresponded to mol. wts c. 10 000–17 000 when compared with standard proteins. These bands were interpreted as possibly representing core components of LPS, because two of them were common to LPS 7/1 and

![SDS-PAGE](image)

Fig. SDS-PAGE of lipopolysaccharide (LPS) from four strains of *P. aeruginosa* on 12.5% acrylamide gels. Samples contained 10 µl of LPS (5 mg/ml) in buffer. Lane 1, standard proteins; lane 2, LPS 7; lane 3, LPS 7/1; lane 4, LPS 18R; lane 5, LPS 18S. Mol. wt (10^3) markers are shown.

LPS 18R (lanes 3 and 4 respectively); but only one was present in the track of LPS 18S (lane 5). Narrow bands forming a regular ladder throughout the middle part of the gel were distinguishable in the tracks of LPS 7 and 7/1; but the densely staining bands of high mol. wt (60 000–95 000), of LPS 7, were absent from the track of variant LPS 7/1. The LPS of strain 18R (lane 4) did not exhibit molecular size heterogeneity, as evidenced by the lack of multiple banding; but two major bands corresponding to mol. wts c. 35 000 and 50 000 were developed by the silver stain. These bands were not found in the track of the variant LPS 18S. No protein bands were detected in any of the LPS tracks on staining with Coomassie blue.

Serological reactions of lipopolysaccharides

The LPS of strain 7 adsorbed efficiently to sheep erythrocytes, and high homologous titres of haemagglutination with antisera were obtained (table IV) in tests repeated at least three times. The titres given by LPS 7/1 were variable and appeared to be
**Table IV.** Indirect haemagglutination of lipopolysaccharide (LPS) of parent and variant strains of _P. aeruginosa_ by homologous and heterologous O antisera

<table>
<thead>
<tr>
<th>Antiserum to strain</th>
<th>Haemagglutination titre with LPS of strain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7</td>
</tr>
<tr>
<td>7</td>
<td>40 960</td>
</tr>
<tr>
<td>7/1</td>
<td>640</td>
</tr>
<tr>
<td>18R</td>
<td>0</td>
</tr>
<tr>
<td>18 S</td>
<td>0</td>
</tr>
</tbody>
</table>

* Range of titres in repeated tests.
  
  0 = <20.

influenced by the batch of LPS used to sensitise the erythrocytes, because homologous titres ranged from 40 to 2560. A similar variation was observed in tests with anti-18S serum and LPS 7/1; but the homologous titres of the parent strains 7 and 18R were reproducible. The results obtained by haemagglutination were in general agreement with those given by agglutination of whole cells (see table II), and confirmed that LPS 7/1 was distinguishable from the parent LPS 7. Moreover, the partial serological relationship between the variant strains, as indicated by agglutination tests, was confirmed by the reaction of anti-18S serum and LPS 7/1 in haemagglutination tests. The LPS of strain 18R gave the same titre with anti-18S serum, although the LPS of 18S failed to react with antibody to 18R.

**Detection of antigenic variants in CF isolates**

It was evident on examination of primary cultures of sputa that most patients were colonised by two or more colonial forms of _P. aeruginosa_. In total, 553 cultures of _P. aeruginosa_ were isolated from specimens taken from 115 CF patients on at least two separate occasions. These were tested for serum sensitivity, and were serotyped by agglutination with O-antiserum and with the two antisera to the variant strains 7/1 and 18S. Table V shows that 414 (75%) of the isolates were fully sensitive to serum, and only 70 (13%) were resistant; 69 were partially sensitive to serum, giving a 10- to 1000-fold decrease in viable count compared with a control.

The association of serum sensitivity and loss of O-specific reaction observed previously by Penketh _et al._ (1983) was confirmed with this series of cultures. O-typable cultures were most often serum resistant or only partially sensitive to serum. Amongst the 17 cultures which were fully sensitive to serum but reacted with a single O serum, 6 were serotype O11 and were isolated from the same patient on repeated visits to the out-patients clinic; the other serotypes in this group were O3, O6, O9, O10 and O16.

Approximately one-fifth of the cultures did not react with any of the O-antisera, and 84 (69%) of 122 were serum sensitive. A number of these strains may have had specific O antigens, but were not agglutinated because of their extreme mucoid nature (35%) which made them non-emulsifiable in saline. Of the 553 isolates, 144 (26%) were agglutinated by anti-7/1 (PA) serum alone, in contrast with 56 isolates (10%) by anti-18S serum; but 147 cultures (27%) were agglutinated by both sera. Three O-typable serum-resistant cultures gave a reaction with the PA serum (anti-7/1), and seven of the serum-resistant group which were O-non-typable were agglutinated by this serum. Statistical analysis of the data confirmed that most serum-

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**Table V.** Sensitivity to normal human serum, and agglutination reactions, of 553 isolates of _P. aeruginosa_ from 115 CF patients

<table>
<thead>
<tr>
<th>Serum sensitivity</th>
<th>Number (percentage) of isolates in agglutination category</th>
<th>Total number (%) of isolates</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>O-typable</td>
<td>O-non-typable</td>
</tr>
<tr>
<td>Resistant</td>
<td>40</td>
<td>19</td>
</tr>
<tr>
<td>Partially sensitive</td>
<td>27</td>
<td>19</td>
</tr>
<tr>
<td>Sensitive</td>
<td>17</td>
<td>84</td>
</tr>
<tr>
<td>Total</td>
<td>84 (15)</td>
<td>122 (22)</td>
</tr>
</tbody>
</table>

PA = polyagglutinable; NT = non-typable.
resistant and partially resistant strains were O-typable or O-non-typable, whereas strains which were agglutinated by either variant anti-serum were most often sensitive to serum ($\chi^2 = 193.03$, d.f. = 8, $p < 0.0005$).

Discussion

Holloway and Cooper (1962) were the first to show that a somatic antigen of P. aeruginosa was altered by lysogenisation with phage. Subsequently, Liu (1969) and Bergan and Midtvedt (1975) demonstrated conversion of the O serogroup of reference strains by phage. We used a typing phage to isolate resistant variants of its homologous propagating strain, and chose a culture which had become resistant and partially resistant strains were O-typable or O-non-typable, whereas strains which were agglutinated by either variant anti-serum were most often sensitive to serum ($\chi^2 = 193.03$, d.f. = 8, $p < 0.0005$).

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Holloway B W, Cooper G N 1962 Lysogenic conversion in
Govan J R W 1978 Pyocin typing of
mittent antibiotic therapy the organisms persisted.
and yield, and were readily serogrouped by specific
Darveau R P, Hancock R E W 1983 Procedure for isolation of
Cryz S J, Pitt T L, Furer E, Germanier R 1984 Role of
infection into the blood stream, but despite inter-
progressively lost their O-specificity and became
bladder for several months and even years, but they
infecting strains of E. coli were serum resistant, had
a normal LPS as determined by gross composition
in CF, where a progressive antigenic change has
accompanied increase of sensitivity to serum during
the course of respiratory infections (Penketh et al.,
1983).

The loss of O-specific side chains by P. aeruginosa
in CF may result in a decrease of virulence; Cryz et al.
(1984) showed that for burned mice a LPS-
defective variant was 1000-fold less virulent than its
parent. The continued survival of these organisms
in the CF lung is therefore paradoxical in that they
may be avirulent, sensitive to serum and hyper-
sensitive to antibiotics (Irvin et al., 1981) and yet
contribute to mortality in CF patients (Penketh et al.,
1983). Antigenic variation of P. aeruginosa in
CF may occur as a result of a selective pressure of
the host’s immune response towards the typical
antigens of the initial infecting strain. The emergence
of variant strains with defective antigens may
not be recognised immunologically and the lack of
antibody response towards these antigens may
facilitate the survival of these strains in the already
diseased lung of the CF patient.

We thank Drs J. C. Batten and M. E. Hodson for access to the
patients and Sister F. Duncan for the collection and dispatch of
specimens. The work was funded in part by the Cystic Fibrosis
Trust.

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