PREVENTION OF SWARMING OF *CLOSTRIDIUM SEPTICUM*

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PLATES XXXII–XXXV

In earlier papers (Williams and Willis, 1970; Willis and Williams, 1970; Williams, 1971) we reported that the surface swarming of *Clostridium tetani* in plate cultures is prevented by incorporation into the medium of commercial tetanus antitoxic serum, and that this technique provides a superior alternative to firm agar for obtaining surface viable counts of *C. tetani*. Because all strains of *C. tetani* are known to share a common O antigen, it seemed likely that this inhibitory effect on swarming was due to the presence in the antiserum of agglutinating O antibody to the vegetative cell.

Strains of *C. septicum* are divisible into two groups on the basis of specific components of the O antigen, neither of which is shared with *C. chauvoei* (Moussa, 1959). Batty and Walker (1963) applied this difference in the O antigenic components of the two organisms to the preparation of fluorescent labelled antibodies which may be used as specific stains to differentiate between *C. chauvoei* and *C. septicum* in smears and in tissue sections.

In the present study we report upon the use of *C. septicum* O antiserum for the prevention of swarming growth of *C. septicum* in plate cultures.

**Materials and methods**

Organisms. Twenty-five strains of *C. septicum* were used. These were kindly provided by Dr S. P. Lapage from the National Collection of Type Cultures (NCTC), and Dr P. D. Walker from the Wellcome Research Laboratory Culture Collection (CN). The strains were respectively nos. NCTC281, 282, 283, 284, 285, 501, 504, 547, 549, 550, 551; and nos. CN424, 2779, 3232, 3608, 3847, 3848, 3941, 3942, 3957, 4212, 5115, 5421, 5545. In addition, two strains of *C. chauvoei* (NCTC8070 and 8590) and stock strains of *C. welchii* type A, *C. bifermentans*, *C. oedematios* type A and *C. histolyticum* were used.

Preparation of O antigen. This was as described by Batty and Walker.

Preparation of antiserum. The method used was that described by Batty and Walker. Antiserum was prepared against a mixture of four strains of *C. septicum*, each of the two serological groups of Moussa being represented by two strains. The strains used were nos. NCTC501, 551, 281 and 282. Courses of immunisation repeated over a period of months yielded sera with agglutinating titres of the order of 2560.

Media. Fresh horse blood agar was used as the plating medium, and was prepared in the usual way from an ox heart digest broth base (Cruickshank, 1965); layer plates containing 5 per cent. of defibrinated horse blood (Wellcome) were used. The ordinary agar medium contained 1-5 per cent. of New Zealand agar; firm agar plates contained 4 per cent. of New Zealand agar. For the plate culture of *C. chauvoei* the medium was enriched with...

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0.5 per cent. of glucose and 1 per cent. of liver extract. All media contained 0.05 per cent. of cysteine hydrochloride. *Clostridium septicum* antiserum blood agar was prepared by spreading the blood agar plates with 0.2-0.5 ml of the undiluted rabbit serum.

**Seeding of plates.** In order to assess inhibition of swarming growth, half-antiserum plates were seeded in a chordal area across the centre of the plate at right angles to the antiserum area. Surface viable counts were performed by the method of Miles, Misra and Irwin (1938) with 10-fold dilutions of the inoculum prepared in 0.9 per cent. NaCl in water containing 0.05 per cent. of cysteine hydrochloride (BDH). Cultures were incubated for 24 and 48 hr in an anaerobic atmosphere containing 10 per cent. CO₂. In all cases the inoculum was the supernatant from a fresh 18-hr cooked meat broth culture of the organism under test.

<table>
<thead>
<tr>
<th>Strain of <em>Cl. septicum</em></th>
<th>Surface viable count per ml on antiserum blood agar</th>
<th>Surface viable count per ml on firm blood agar</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NCTC281</td>
<td>10⁸ D*</td>
<td>10⁸ R⁺</td>
</tr>
<tr>
<td>NCTC501</td>
<td>2.5 × 10⁷ D D</td>
<td>5 × 10⁷ R</td>
</tr>
<tr>
<td>CN3847</td>
<td>4 × 10⁸ D</td>
<td>4 × 10⁸ R</td>
</tr>
<tr>
<td>CN3848</td>
<td>1.5 × 10⁶ D</td>
<td>RNCP‡</td>
</tr>
<tr>
<td>CN3941</td>
<td>10⁸ D</td>
<td>10⁸ R</td>
</tr>
<tr>
<td>CN3942</td>
<td>2 × 10⁸ D</td>
<td>3.5 × 10⁸ R</td>
</tr>
<tr>
<td>CN3957</td>
<td>2.5 × 10⁸ D</td>
<td>1.5 × 10⁸ R</td>
</tr>
<tr>
<td>CN4212</td>
<td>1.5 × 10⁹ D</td>
<td>RNCP</td>
</tr>
<tr>
<td>CN5115</td>
<td>10⁷ D</td>
<td>1.3 × 10⁷ R</td>
</tr>
<tr>
<td>CN5545</td>
<td>10⁷ D</td>
<td>10⁸ R</td>
</tr>
</tbody>
</table>

*D* = Discrete colonies; † *R* = rhizoidal colonies; ‡ RNCP = rhizoidal, no count possible.

**RESULTS**

Inhibition of swarming growth of *Clostridium septicum* by antiserum

All 25 strains of *Cl. septicum* showed complete inhibition of swarming growth on the antiserum half of the plate. Colonies and areas of confluent growth of *Cl. septicum* on the antiserum medium showed an irregular edge, but this rhizoidal structure was much more truncated than is usually seen on ordinary blood agar. Discrete colonies showed a particularly compact structure, commonly presenting an entire appearance to the naked eye; under the plate microscope these colonies showed a mildly lobulated edge. As expected, most strains showed haemolytic activity which was not inhibited by the antiserum (figs. 1 and 2).

Comparison of surface viable counts on antiserum blood agar and firm blood agar

The surface viable counts of ten strains of *Cl. septicum* (including two of the strains used in the preparation of the antiserum), performed on firm blood agar and on antiserum blood agar, are summarised in the table; they represent the mean of quadruplicate counts in each case. None of the strains produced overt swarming on either of these media.
**Fig. 1.**—*Cl. septicum* on half-antiserum horse blood agar. The swarming growth is inhibited by *Cl. septicum* antiserum on the left-hand half of the plate. Haemolytic activity is not affected. ×1-2.

**Fig. 2.**—Discrete colonies of *Cl. septicum* on antiserum-treated horse blood agar. ×72.

**Fig. 3.**—Discrete colonies of *Cl. septicum* in a surface viable count inoculation zone on *Cl. septicum* antiserum horse blood agar. ×14.
PREVENTION OF SWARMING OF \textit{Cl. septicum}

Fig. 4.—Rhizoidal colonies of \textit{Cl. septicum} in a surface viable count inoculation zone on firm (4 per cent.) agar. \( \times 14 \).

Fig. 5.—Higher-power view of two of the colonies shown in fig. 4. \( \times 72 \).
FIG. 6.—Discrete colonies of *Cl. septicum* and *Cl. chauvoei* on *Cl. septicum* antiserum horse blood agar. ×36.

FIG. 7.—Colonies of *Cl. chauvoei* overgrown by swarming growth of *Cl. septicum* on horse blood agar. ×36.

FIG. 8.—Colonies of *Cl. oedematiens* type A on either side of a discrete colony of *Cl. septicum* on *Cl. septicum* antiserum horse blood agar. ×72.
PREVENTION OF SWARMING OF *Cl. septicum*

**Fig. 9.**—Colony of *Cl. oedematiaiens* type A being engulfed by swarming growth of *Cl. septicum* on horse blood agar. ×18.

**Fig. 10.**—Discrete colonies of *Cl. welchii* and *Cl. septicum* on *Cl. septicum* antiserum horse blood agar. ×36.

**Fig. 11.**—Colonies of *Cl. welchii* being engulfed by swarming growth of *Cl. septicum* on horse blood agar. ×36.
All strains of _Cl. septicum_ gave countable colonies on antiserum blood agar (fig. 3). The colonies were small, circular and discrete, with a mildly lobulated edge and usually an absence of rhizoidal outgrowth. On firm blood agar, on the other hand, two of the strains produced growth that was so irregular in structure that it was not possible to identify separate colonies accurately and counts could not be obtained. Colonies of the other eight strains were markedly irregular but were easily countable (figs. 4 and 5).

Among the eight strains for which surface viable counts were obtained on both media, seven showed essentially comparable results, and the remaining one gave a count that was marginally higher on the firm agar medium.

Isolation of _Clostridium chauvoei_ from mixtures with _Cl. septicum_ on _Cl. septicum_ antiserum blood agar

The principle of inhibition of swarming growth of _Cl. septicum_ by antiserum was applied to the separation of two strains of _Cl. chauvoei_ (nos. NCTC8070 and 8590) from mixtures with _Cl. septicum_ in plate cultures. Mixtures of the two species were examined on half-antiserum horse blood agar plates enriched with glucose and liver extract. It was a simple matter to pick off discrete colonies of _Cl. chauvoei_ from the antiserum half of the plates; the untreated half of each plate showed a confluent growth of _Cl. septicum_ overlying colonies of _Cl. chauvoei_ (figs. 6 and 7).

The use of antiserum-treated plates was equally successful in the separation of other anaerobes from mixtures with _Cl. septicum_, including _Cl. welchii_ type A, _Cl. bifermens_, _Cl. oedematiens_ type A and _Cl. histolyticum_ (figs. 8–11). Unlike concentrated agar media, antiserum-treated plates had no adverse effect on the growth of other clostridia, but allowed the free development of their characteristic colonies. This was especially marked in the case of _Cl. chauvoei_, which may grow poorly or not at all on firm agar media.

**DISCUSSION**

The formation by _Cl. septicum_ of discrete surface colonies in the presence of _Cl. septicum_ O antiserum is probably the result of "agglutination" of the growing culture. Watt (1972) made use of this procedure for the enumeration of _Cl. septicum_ and obtained useful inhibition with a selected antitoxic serum. It is important to note that the definite results obtained in our present study with many strains of _Cl. septicum_ are presumably attributable to the use of a polyvalent serum intentionally prepared against the O antigenic components of the organism.

Although the use of firm agar is almost as effective as the use of plates bearing _Cl. septicum_ antiserum in preventing surface swarming growth of _Cl. septicum_, it is less satisfactory as a method for the isolation of other clostridia from mixtures with _Cl. septicum_. This applies particularly to exacting species such as _Cl. chauvoei_, the growth of which may be so retarded by high concentrations of agar that macroscopic colonies fail to develop.
It seems that the antiserum method of restricting surface growth of *Clostridium septicum* may find a useful application in veterinary anaerobic bacteriology. *Clostridium septicum* is a normal inhabitant of the intestinal tract of herbivorous animals, whence it rapidly invades the tissues post mortem. Consequently, the isolation of *Clostridium septicum* from these animals does not imply a causal relation to a pathological process. In the case of isolation of *Clostridium chauvoei* from blackleg lesions, difficulty is often encountered when *Clostridium chauvoei* is overgrown by the nutritionally less demanding *Clostridium septicum*, a state of affairs that is aggravated in primary plate cultures by the swarming habit of *Clostridium septicum*. Batty and Walker (1963) utilised the dissimilarity of the O antigenic components of *Clostridium septicum* and *Clostridium chauvoei* to develop a fluorescent antibody-staining technique that serves to differentiate between the two species in smears and tissue sections. We have extended this exploitation of the property of antigenic dissimilarity to the separation of *Clostridium chauvoei* from *Clostridium septicum* in artificial culture.

Surface viable counts of *Clostridium septicum* may be conveniently obtained by the method of Miles and Misra when horse blood agar treated with *Clostridium septicum* O antiserum is used as the plating medium. Antiserum-treated agar is marginally superior to firm agar for this purpose.

**SUMMARY**

A *Clostridium septicum* O antiserum, prepared in the rabbit against the two serological groups of Moussa, inhibits the swarming growth of *Clostridium septicum* in plate cultures. Advantage may be taken of this in the separation of other organisms from mixtures with *Clostridium septicum* and in performing surface viable counts with *Clostridium septicum*.

We are greatly indebted to Mr J. Harrison for production of the photographs.

**REFERENCES**


Moussa, R. S. 1959. Antigenic formulae for *Clostridium septicum* and *Clostridium chauvoei*. *J. Path. Bact.*, 77, 341.


