Significance of the isolation of *Clostridium welchii* from routine blood cultures

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**SYNOPSIS** *Clostridium welchii* has been demonstrated in approximately 20% of contact plates taken from the antecubital fossa of 185 inpatients and outpatients and healthy staff. The highest incidence was in a group of 40 very ill patients. The isolation of the organism from blood cultures is not always of clinical significance. Skin preparation as at present practised is often inadequate to remove the spores when contamination is relatively heavy, for example, in bedridden patients.

In previous years a proportion of our blood cultures have yielded *Clostridium welchii* in one only of the three bottles, all of which have been shown to be capable of supporting its growth.

In 1974 the laboratory processed about 3000 blood cultures. In 30 of these cultures yielding growth the organism isolated was *Cl. welchii*. The majority of patients from whom these cultures were taken did not seem to have systemic infection and in no case did bacteraemia with *Cl. welchii* seem to have had pathological effects.

A possible explanation was that *Cl. welchii* spores might exist on the skin of the antecubital fossa in the way described by Parker (1969) and Ayliffe and Lowbury (1969) for the skin of the buttock and thigh. In an investigation of a small group of 25 individuals, Collee and Watt (1971) have demonstrated a few colonies of *Cl. welchii* on swabs taken from the hands and forearms in three by direct plating. This proportion rose to over 50% of 17 hand swabs after enrichment, as would of course occur in normal blood culture techniques.

**Material and Methods**

Nagler medium screened with 100 μg/ml neomycin, as described by Lowbury and Lilly (1955), was used in the form of Contact plates. Colonies growing on this medium and showing lecithinase activity after 18 hours' anaerobic incubation were accepted as presumptive *Cl. welchii*. Few other organisms grow on this medium.

Presumptive *Cl. welchii* isolates were subjected to the following confirmatory tests:

1. Subculture aerobically and anaerobically on blood agar to prove that there were strict anaerobes
2. A half antitoxin plate using *Cl. welchii* antitoxin and including a known *Cl. welchii* as a control to demonstrate specific neutralization of lecithinase activity
3. Inoculation of litmus milk to show the 'stormy clot' formation
4. Demonstration of acid production from lactose in a peptone water sugar tube.

The last two tests were done to distinguish *Cl. bifermontans*, which does not ferment lactose but shows cross neutralization with welchii antitoxin.

**Sampling Methods**

The skin of the antecubital fossa of both arms was sampled by the use of surface Contact plates. The skin was then disinfected with 70% isopropyl alcohol (Medi Swab) using 20 strokes for 30 seconds. The disinfectant was allowed to dry, and sampling was repeated in the same area. Both plates (before and after skin treatment) were incubated anaerobically overnight, examined for presumptive *Cl. welchii* colonies, and compared to demonstrate the effect of skin preparation.

One hundred inpatients were studied over an eight-week period. A control group of 85 ambulant patients attending the anaemia outpatient and staff members were investigated in a similar way.

**Results**

The results of these investigations are shown in tables I to III.
Table I  Numbers of Cl. welchii isolated from patients studied

<table>
<thead>
<tr>
<th>Subjects</th>
<th>Total</th>
<th>No. positive</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inpatients</td>
<td>100</td>
<td>21</td>
<td>21</td>
</tr>
<tr>
<td>Outpatients and staff</td>
<td>85</td>
<td>16</td>
<td>18</td>
</tr>
</tbody>
</table>

Table II  Cl. welchii isolated from inpatients according to severity of their illness

<table>
<thead>
<tr>
<th>Total Inpatients</th>
<th>Before Skin Preparation</th>
<th>After Skin Preparation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. Positive %</td>
<td>No. Positive %</td>
</tr>
<tr>
<td>100</td>
<td>21</td>
<td>16</td>
</tr>
</tbody>
</table>

Table III  Effect of skin preparation with isopropyl alcohol

Repeat sampling of nine patients giving positive results and of nine negative patients as a control, after a two-week interval, showed that of nine long-stay female orthopaedic patients mainly suffering from fractured neck of the femur, three demonstrated persisting carriage over this period (table IV).

Table IV  Persistence of carriage of Cl. welchii over a two-week period

<table>
<thead>
<tr>
<th>No. of Patients</th>
<th>No. carrying Cl. welchii after 2 Weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td>9 previously positive</td>
<td>3 still carrying Cl. welchii</td>
</tr>
<tr>
<td>9 negative</td>
<td>1 now carrying Cl. welchii</td>
</tr>
</tbody>
</table>

Discussion

In our study of 100 inpatients, which revealed 21% antecubital fossa carriage of Cl. welchii, most carriers were elderly women who had been in hospital for a long time, for example, geriatric orthopaedic patients. When lying in bed their arms were in such a position that the antecubital fossa was in contact with the buttock area or the sheaths in this area. These patients were heavily contaminated with Cl. welchii spores, ie, more than 100 colonies per plate, and attempts at disinfection had little effect in removing the spores.

Three out of nine carried the organism for a period of more than two weeks. One new carrier previously negative was demonstrated at the end of the period. This may have represented only a sampling error in the original test.

The effectiveness of isopropyl alcohol in removing light contamination (ie, one or two colonies) can be explained as mechanical removal since alcohol is not sporicidal. Compresses of the iodophor, povidone-iodine, applied for 15-30 minutes have been found to be useful by Lowbury et al (1964) and Ayliffe and Lowbury (1969). Washing with soap and water was found to be ineffective by Ayliffe and Lowbury (1969). We did not try these methods because they are not used in skin preparation before sampling blood for culture.

The method of preparing the skin for this purpose was that recommended by Washington (1975). The skin is cleansed with 70-90% ethyl alcohol followed by an alcoholic solution of iodine (1-20%). Using this method, 70% alcohol alone has been found to decrease the skin microflora by at least 90%. However, it will certainly not kill all the bacteria in the deep layers of the dermis, particularly in structures such as sweat glands and hair follicles (Updegraft, 1964). Alternatively, Washington recommended the use of povidone iodine swab sticks. The intended site of needle entry should not be touched unless the fingers used for palpation have been similarly disinfected or gloves are worn. Upon completion of the venepuncture residual iodine should be removed with an alcohol sponge or pad.

Our tests again illustrate the ineffectiveness of much skin preparation before venepuncture, showing that at least when carriage is heavy the perfungatory techniques often employed will not remove clostridial spores. The finding of Cl. welchii in blood cultures is not necessarily of clinical significance, and even its repeated isolation over a period of weeks may not represent infection in the patient but merely, in some cases, contamination from the site of venepuncture.

Our thanks are due to Mr M. Kreuzer, of Sterilin Limited, for making available the supply of Contact plates used in this study.

References

Significance of the isolation of Clostridium welchii from routine blood cultures.
F J Ahmad and J H Darrell

doi: 10.1136/jcp.29.3.185

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