Comparative study of two systems for detecting bacteraemia and septicaemia

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SYNOPSIS Two blood culturing systems were compared in parallel—a semi-automated radiometric method, BACTEC, and the conventional method used in our laboratory. BACTEC contains radioactive 14C labelled substrates and monitors the level of 14CO₂ evolved by actively metabolizing bacteria.

BACTEC was as efficient as the conventional method in detecting positive cultures and indicated that it was superior in detecting positive cultures earlier than the conventional system.

Rapid detection of bacteraemia is of utmost importance for patient care and prognosis. Commercial systems are available which attempt to decrease the time of detection and isolation; BACTEC is based on the detection of radioactive 14CO₂ produced by the decomposition of 14C labelled glucose and other 14C labelled substrates by microbial action. The semi-automated model BACTEC 301 was used in this comparative study. One hundred and thirty-nine paired aerobic and anaerobic blood cultures were examined by both systems from patients at Green Lane Hospital, Auckland.

Material and Methods

Each system employs two culture bottles, one for cultivating aerobes and one for anaerobes. All bottles were inoculated with 3 ml of blood drawn at the bedside by the clinical staff in the wards selected for this trial (2 cardiological and 2 surgical) and sent to the laboratory. Where requested, penicillinase or ß lactamase were added to the blood cultures.

CONVENTIONAL SYSTEM

The conventional system consisted of diphasic glucose broth containing 50 ml of brain-heart infusion broth with 1% glucose and 0.05% sodium polyanethol-sulphonate for the aerobes, and 20 ml thiglycollate broth with indicator-135 (Baltimore Biological Laboratories) and sodium polyanethol-sulphonate for the anaerobes. An atmosphere containing 5% CO₂ and 95% O₂ was added to the glucose broth bottles. Cultures were incubated at 36°C and examined visually twice daily for 10 days. Films were made from second day glucose broth cultures and stained with May-Grünwald/Giemsa for the presence of organisms not visually observable. Any visually suspicious or positive cultures were Gram stained and subcultured accordingly. At day 10 the remaining negative glucose broth cultures had films made and stained with May-Grünwald/Giemsa and were subcultured onto blood and chocolate agar half plates. These were incubated in a candle jar at 36°C for 48 hours before being discarded.

Anaerobic cultures were examined similarly, May-Grünwald/Giemsa stains being made and examined on second and fifth day cultures. All day 10 cultures and any suspicious or positive cultures were subcultured onto prereduced brain-heart infusion agar (BBL) and incubated anaerobically, and also onto blood agar for aerobic culture. All subcultures were kept for 48 hours before being discarded.

BACTEC SYSTEM (semi-automated model 301)

The BACTEC system consisted of two vials—the aerobic vial (6A vial) containing tryptic soy broth under a CO₂ atmosphere and the anaerobic vial (7B vial) containing reduced tryptic soy broth under a N₂/CO₂ atmosphere. Each vial contained sodium polyanethol-sulphonate and 14C labelled substrates (1-5 µCi).

All day 1 and day 2 aerobic cultures were placed on a reciprocating shaker in a 36°C waterbath and tested three times on day 1 and twice on day 2. The remaining negative cultures were placed in an
incubator (36°C) and tested on each of the remaining five days.

The anaerobic vials were not agitated and were tested first on day 2; the negatives were reincubated and tested once on the following days. All cultures were examined visually before being read on the instrument.

The atmosphere contained in the culture vials was drawn into an ionization chamber within the BACTEC 301 instrument and the amount of radioactivity was measured and displayed on an arbitrary scale, the growth index (GI) ranging from 0 to 100. GI readings of 25 or more were considered to be positive and were immediately Gram stained and subcultured as for the conventional method.

The flushing gases used were: for the aerobic vial 5% CO₂/air, and for the anaerobic vial 10% CO₂/10% H₂/N₂ which had been passed through a DEOXO catalyst (Englehard Industries, E. Newark, NJ) to remove any traces of oxygen.

All BACTEC vials were incubated for seven days, negative cultures being stained and subcultured at day 7 as for the conventional method.

Staphylococcus epidermidis, Bacillus species, and aerobic and anaerobic diphtheroids were considered to be contaminants unless they were growing in a number of cultures from the same patient. In these cases the significance of the isolates was determined by the patient’s clinical condition.

### Results

Of 139 blood cultures from 58 patients, 28 (20.1%) cultures were positive by one or both methods. Contaminants were isolated from 12 (8.6%) of these, leaving 16 (11.5%) positive cultures from six patients with true bacteraemia/septicaemia. Neither system detected all positive patients, each detected 14 (88%) positive cultures (table I). The organisms isolated by each system are given in table II. Others (Brooks and Sodeman, 1974; Smith and Little, 1974; Thiemeke and Wichers, 1975) have found a lower frequency in isolating coliform organisms by BACTEC. The results tabulated in table II indicate that this could be so although the number of enteric isolates is lower.

All the Propionebacterium acnes were isolated from one patient undergoing dental surgery and were considered significant.

An interesting isolate of a yeast, Hansenula anomala, grew in two sets of cultures from one patient with endocarditis. In the first set the conventional method was positive on day 3 compared with the BACTEC positive on day 7. The second set was positive in both systems by day 3.

Of the 12 paired positive cultures, four (33.3%) were detected first by BACTEC, one (8.3%) by the conventional method, and seven (58.3%) were detected simultaneously by each system.

### Table I  Summary of BACTEC results: 139 blood cultures from 58 patients examined in parallel

<table>
<thead>
<tr>
<th>Method</th>
<th>Positive Cultures</th>
<th>Positive Patients¹</th>
<th>Organisms</th>
<th>Contaminants²</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No.</td>
<td>%</td>
<td>No.</td>
<td>%</td>
</tr>
<tr>
<td>BACTEC</td>
<td>21</td>
<td>75</td>
<td>14</td>
<td>88</td>
</tr>
<tr>
<td>Conventional</td>
<td>19</td>
<td>68</td>
<td>14</td>
<td>88</td>
</tr>
<tr>
<td>One or both methods</td>
<td>28</td>
<td>100</td>
<td>16</td>
<td>100</td>
</tr>
</tbody>
</table>

¹Numbers of growing cultures from positive patients.

²Some vials grew more than one contaminant.

### Table II  Organisms isolated from positive patients

<table>
<thead>
<tr>
<th>Organism</th>
<th>Method</th>
<th>BACTEC</th>
<th>Conventional</th>
<th>One or both methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staph aureus</td>
<td></td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>E. coli</td>
<td></td>
<td>1</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>P. acnes¹</td>
<td></td>
<td>4</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Thiol dependent streptococcus</td>
<td></td>
<td>6</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Klebsiella</td>
<td></td>
<td>0</td>
<td>1²</td>
<td>1</td>
</tr>
<tr>
<td>Hansenula anomala (yeast)</td>
<td></td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>14</td>
<td>15</td>
<td>17</td>
</tr>
</tbody>
</table>

¹All were isolated from the same patient; 5 out of 9 cultures grew P. acnes and this was considered significant.

²One culture isolated two organisms, E. coli and Klebsiella.

Contaminants included S. epidermidis, Staph. aureus (grown in 1 out of 18 culture bottles), aerobic sporing bacillus, and aerobic and anaerobic diphtheroids.
All organisms isolated from positive patients were detected by day 7 by each system.

Two BACTEC vials grew Propionibacterium acnes and Staphylococcus aureus on subculture at day 7. These were considered to be contaminants: the Staph. aureus grew in one out of 18 culture bottles taken from one patient.

A number of cultures produced GI readings > 25 with no organisms demonstrable in a MayGrünwald/Giemska film or on subculture. These were referred to as 'false positive' cultures and included 17 (12.2%) of the total 139 cultures. Most of these gave a GI reading of less than 30, seven (5%) giving GI readings of more than 30. An explanation of this phenomenon was that actively metabolizing leucocytes may have contributed to the false positive readings (Thiemke and Wichcr, 1975). Indeed, leucocyte counts determined at the time of culturing showed that each of these patients had increased leucocyte counts ranging from 10-8 to 29.3 × 10^8 cells/mm^3 with an average count of 24.0 × 10^8 cells/mm^3. Cultures taken from two patients produced high GI readings (in excess of 30 at day 7), no organisms being demonstrated on repeated subculturing and staining. These may constitute a small proportion of cultures where organisms were present but were not demonstrable. One patient was suffering from a meningococcal meningitis with a haematological blood film picture consistent with severe infection. The other patient was admitted to hospital with abdominal pain and lobar pneumonia.

An alarming 12 (75%) positive cultures were visibly growing in the anaerobic 7B vial with GI readings less than 25, only four (25%) positive cultures producing a GI of more than 25.

There were no 'false negative' cultures among the aerobic 6A vials.

Discussion

The data presented indicated that the BACTEC system was as efficient as our routine method in isolating organisms from bacteraemic/septicaemic patients. BACTEC also appeared to be superior in detecting positive cultures earlier than our conventional method. There is general agreement with the results of this study and results obtained by others (Brooks and Sodeman, 1974; Thiemke and Wichcr, 1975). However, it must be stressed that the total number of cultures examined and the number of positive cultures detected was small. We were supplied with a limited amount of media for this initial comparative study and were unable to obtain further supplies in order to assess a larger range of patients.

From previous experience in this laboratory, Giemska staining was found to be superior in detecting small numbers of Gram-negative organisms in blood cultures which otherwise may have been easily missed in a Gram stain. Any positive Giemska films were subsequently Gram stained and subcultured onto appropriate media.

A major concern was the occurrence of the false positive cultures. The number of false positive and false negative cultures varied, depending upon what GI would be taken as showing the presence of viable organisms. In this study a GI of 25 or more was taken as an indication of growth, giving an incidence of 12.2% false positive cultures, all occurring in the aerobic 6A vial. There were no false negatives among the aerobic 6A vials but 75% of the positive anaerobic culture vials (7B vials) failed to register a positive GI reading. This emphasizes the need for visually checking cultures for signs of growth before reading on the instrument. Several factors may have contributed to the high incidence of false negative anaerobic cultures:

1. Inability of the organism to produce 14CO2 from the substrates supplied. This would indicate an expansion of the 14C substrates provided in the media by the manufacturers, to accommodate these organisms.

2. Differing modes of metabolism under anaerobic conditions. The thiol dependent streptococcus failed to register a positive GI reading in the anaerobic vial although they were visually growing, yet in the aerobic vial they were positive on the BACTEC before there was visible sign of growth. Most of the P. acnes isolated and two contaminating aerobic sporing bacilli also failed to register positive GI readings.

3. The anaerobic vials had expired their suggested period of use just before the beginning of this study and a further supply could not be obtained.

Brooks and Sodeman (1974) used a GI reading of 20 or greater with an incidence of 43% false positive cultures. They suggested that if a GI reading of 30 had been used, the number of negative vials subcultured would have been 1.5% at the expense of larger detection times for 95% of the positive Candida cultures. These generally gave a GI reading of less than 30 at the time of subculture.

Others (Smith and Little, 1974; Thiemke and Wichcr, 1975) have used a GI reading of 30 as indicative of growth. Thiemke and Wichcr (1975) found 6.8% false positive cultures overall; however, 6.8% of the positive cultures failed to register a positive GI. Of 14 isolates in this category, nine were fungi. Smith and Little (1974) found 8.9% false positive cultures in their study, 6.3% positive cultures giving a GI of less than 30.

After examining these studies we decided to compromise and take 25 as a positive GI reading.
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It became apparent that several organisms, although present in the BACTEC or conventional vials, failed to grow on subculture. This emphasized the need to be aware of their inability to grow on ordinary subculturing media and for designing a suitable medium for their cultivation. These organisms included the thiol dependent streptococcus and many of the anaerobic diphtheroids. Solid media for subculturing were improved by incorporating L-cysteine, menadione, haemin, and yeast extract in brain-heart infusion agar. Dithiothreitol and palladium chloride were added to the supplemented BHI agar and pre-reduced for the isolation of fastidious anaerobes.

Subculturing techniques must be of a high quality so that organisms found on primary blood culture can be subcultured for further studies (ie, identification and susceptibility testing). A good knowledge and high degree of expertise is essential for blood culture work. The BACTEC 301 semi-automated instrument, although removing the need for multiple staining procedures, does not alleviate the tedium of testing large numbers of cultures. In our small trial one person was committed to sampling for most of the day, and if all blood cultures were tested the BACTEC 301 would be unsuitable. In considering an automated method for blood cultures it would be necessary to use the fully automated BACTEC 225, which would allow the technologist freedom to give full attention to subculturing and testing positive cultures.

A further trial of at least 500 patients throughout the whole hospital is considered necessary to do justice to this type of equipment (preferably the 225).

No system has yet been devised that will detect all bacteraemic/septicaemic patients. Although this study involved only a small number of cultures the results indicated that BACTEC is equal to our conventional method in detecting bacteraemia/septicaemia.

I should like to thank Mr B. Cornere for his advice and assistance in the presentation of this paper.

References


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