Comparison of Sentinel and Bactec blood culture systems


Abstract

Aims: To evaluate the Sentinel automated blood culture system and to compare its performance with that of Bactec.

Methods: The Sentinel blood culture system was evaluated in three centres. The performance of the system was assessed in comparison with the routine blood culture method used in these centres, the Bactec system.

Results: Blood culture sets (n = 2180) consisting of Sentinel aerobic and anaerobic, and Bactec aerobic and anaerobic bottles yielded 218 (10%) clinically important isolates. One hundred and fifty five (71%) of the isolates were detected by both systems; 35 (16%) were detected by Sentinel only; and 28 (13%) by Bactec only. For the duration of the evaluation, the Sentinel system was deliberately configured so that it was impossible to detect positive results during the first 12 hours. The times to positivity after the first 12 hours were similar. Data gathered during and subsequent to the evaluation have been used by the manufacturer to refine the algorithm so that positive results can be detected at a minimum of 2-25 hours.

Conclusions: After a period of familiarisation the Sentinel system was considered easy to use. Sentinel is a useful addition to the methods available for the detection of bacteria in blood cultures.

Blood cultures represent one of the most important specimens submitted to clinical microbiology laboratories. They are the mainstay of the microbial diagnosis of septicaemia, infective endocarditis, and conditions associated with a clinical presentation of fever of unknown origin. Many methods have been described and reviewed, but there remains a need for systems which monitor blood cultures continuously and automatically. Sentinel (Difco Laboratories Ltd, East Molesey, Surrey) is an automated blood culture system that detects bacterial growth by monitoring voltages produced within the bottles containing a blood culture medium mixture.

An evaluation of the system was commissioned by the Medical Devices Directorate of the Department of Health, and carried out at three centres: the Westminster Hospital, London, and the Public Health Laboratories at Luton and Leicester. The system's performance was assessed in comparison with the routine blood culture method used in these centres, the Bactec system. A full report of the evaluation has been produced.

Methods

SENTINEL SYSTEM

The Sentinel system consists of a base plinth on which are placed between one and five incubation drawers, each holding 80 bottles containing blood and broth culture media (fig 1). A five drawer (400 bottle) system measures 112 cm high, 75 cm wide, and 75 cm deep. The bottles contain electrodes and the system detects microbial growth by measuring relative changes in voltage produced in the bottle. The Sentinel bottles contain one gold-plated and one aluminium alloy electrode in the base (fig 2). The electrodes are concealed, but on insertion into a Sentinel drawer they pierce a thin membrane and enter the culture medium. This results in the bottle becoming a simple battery, with two dissimilar metals in an electrically conducting fluid. The voltage is generated by the aluminium electrode (anode) slowly dissolving, liberating aluminium ions; the resulting electrons transfer to the gold electrode (cathode) via the drawer circuitry. At the gold electrode, the electrons are transferred to reducible chemical species (electron acceptors) in the medium. In the aerobic system dissolved oxygen is the major electron acceptor, but in the oxygen free anaerobic system alternative compounds fulfil this function. When micro-organisms grow, oxygen (in the aerobic bottle) and the alternative electron acceptors (in the aerobic bottle) are reduced.

Figure 1 Complete Sentinel three drawer system, with computer.
and the voltage between the two electrodes falls.

The drawer electronics detect these voltage changes and an algorithm function within the computer software determines when the voltage has dropped sufficiently to flag the bottle as positive. The Sentinel drawers monitor the voltage between the two electrodes in each bottle at about 14 second intervals. The average value of these readings for each bottle is transmitted to the computer every 15 minutes. Voltage readings less than 40 mv default to zero; that particular 15 minute reading is then excluded from the algorithm. This ensures that removal of bottles does not cause flagging of false positive results. The algorithm has the same format for both aerobic and anaerobic bottles, but there are different coefficients used for the two types of bottle. The system checks the absolute voltage (a voltage level test) and any drop in voltage from the previous reading (the fall off test).

The algorithm used during the evaluation allowed a minimum time of 12-5 hours for the detection of a positive result in the aerobic bottle and 3-75 hours in the anaerobic bottle. In the case of the aerobic bottle the longer period for detection was due to incomplete determination of the algorithm coefficients for the test before evaluation began.

EVALUATION METHODS

The evaluation was carried out in three separate laboratories by an agreed protocol. Different Bactec models were used in each of the three participating laboratories. Westminster Hospital used a Bactec 460 (radiometric detection system) using 6B and 7D media, the Public Health Laboratory at Luton used a model NR 730, and the Public Health Laboratory at Leicester used a model NR 660; the latter two systems used infra-red spectroscopy and 6A and 7A media. The detection thresholds were the same at all centres: a growth value of ≥ 30, or, if there was a rise of ≥ 15 in between readings, the δ value. Although two different detection systems were used, results are reported as being comparable.

Sets of the Sentinel and Bactec bottles (aerobic and anaerobic in each case) were distributed to hospital wards with instructions that 20 ml of blood should be taken and divided equally between each of the four bottles in the set. In the event that less than 20 ml was taken the user was asked to distribute the volume equally between all four bottles. To determine blood volumes introduced into each type of bottle, 10% of the total number of Sentinel and Bactec bottles were weighed before despatch to the hospital wards. These bottles were individually coded so that they could be identified and weighed again on return to the laboratory. The weights were converted to volumes by multiplying them by the value of the mean density of human blood (1.0595 g/ml). The time at which bottles were entered into each system and became positive was recorded; this was automatic in the Sentinel system.

All bottles were incubated for a minimum of seven days, longer if there was a clinical requirement, such as in the case of suspected bacterial endocarditis. After completion of the incubation period all bottles were subcultured on to a blood agar plate, for anaerobic incubation, a heated blood agar plate for incubation in an atmosphere of CO₂, and a MacConkey agar plate for aerobic incubation. All plates were incubated at 37°C for 48 hours.

RESULTS

A total of 2180 complete sets of blood cultures was received during the evaluation—aerobic and anaerobic bottles for both Bactec and Sentinel. The totals for each of the three centres were Westminster 663, Leicester 922, and Luton 595 specimens. In addition, 304 sets, consisting only of Sentinel and Bactec aerobic bottles were received from paediatric patients.

The 2180 complete sets yielded 218 (10%) clinically important isolates (table 1). Of these, 35 (16%) were isolated in the Sentinel system only and 28 (13%) in the Bactec system only: 155 (71%) were detected by both systems.

Among the 304 paediatric samples, two were positive; one was <i>Streptococcus pneumoniae</i>, which was detected only in the Bactec aerobic bottle, the other was <i>Haemophilus influenzae</i>, which was detected only in the Sentinel aerobic bottle.

STRAIN ISOLATED ONLY ON TERMINAL SUBCULTURE

Ten clinically important strains were isolated only on terminal subculture. Four (<i>Streptococcus pneumoniae</i>, β haemolytic streptococcus group C, <i>Staphylococcus aureus</i> and <i>Candida glabrata</i>) were isolated from the Sentinel system. These were not indicated positive by Sentinel but were positive in the Bactec system. Two clinically important isolates, <i>Pseudomonas aeruginosa</i> and <i>S. sanguis</i>, were isolated on terminal subculture from the Bactec system; these were not indicated positive by Bactec but were positive in the Sentinel system.

In addition, three clinically important isolates (<i>Pseudomonas testosteroni</i>, <i>Gardnerella vaginalis</i>, and <i>Proteus mirabilis</i>) were isolated on terminal subculture from Sentinel bottles, which were not indicated positive by either Sentinel or Bactec and were not isolated from the Bactec bottles. One organism, <i>Pseudomonas aeruginosa</i>, was isolated on terminal subculture from the Bactec system, although it had not been indicated positive by either Bactec or Sentinel; it was not isolated from the Sentinel bottles.

FALSE POSITIVE RESULTS

During the trial, 67 (1·5%) Bactec bottles and 130 (3%) Sentinel bottles indicated positive, but were negative by cultural and microscopic methods.

TIME TO POSITIVITY

The comparison of detection times is shown in table 2. Bactec detected 20% of the positive...
samples before six hours compared with 0·5% for Sentinel. This was due to the fact that timings started when samples were entered into each system. In the case of the Bactec this was when they were first tested, often after prior incubation.

The average times to positivity for each type of bottle were: Bactec aerobic 22·4 hours; Bactec anaerobic 28·7 hours; Sentinel aerobic 22·7 hours; Sentinel anaerobic 31·4 hours.

**AMOUNTS OF BLOOD**

The average amounts of blood introduced into each type of bottle were: Bactec aerobic 3·3 ml; Bactec anaerobic 3·2 ml; Sentinel aerobic 3·9 ml; Sentinel anaerobic 3·9 ml. The recommended amount is between 3 ml and 5 ml for Bactec bottles and 5 ml into each Sentinel bottle.

A total of 174 (8%) of all sample sets grew organisms that were regarded as not clinically important. Some sets grew such isolates from more than one bottle, so that a total of 278 bottles (3·2% of the total of 8720 bottles) grew organisms. The numbers of each type of bottle which grew non-significant isolates were: Bactec aerobic 88 (4·0%), Bactec anaerobic 61 (2·8%), Sentinel aerobic 78 (3·6%), Sentinel anaerobic 51 (2·3%).

**Discussion**

The results indicated that the Sentinel system was at least as good as the Bactec systems in the rate of detection of organisms.

Sentinel detected more isolates than Bactec but the numbers were small and the difference not significant. The types of organisms detected were diverse, doubtless reflecting the different patient populations tested in a multicentre evaluation. The range of organisms detected was comparable in both systems. The number of complete blood culture sample sets examined (2180) and the significant positive rate (218) over the period of evaluation, exceeded the minimum numbers (2000 and 200 respectively) recommended by Illstrup. Of particular note was the fact that Sentinel detected seven isolates of *Escherichia coli* which were not detected by Bactec while Bactec in turn detected two strains of *E coli* which Sentinel did not; *E coli* was detected in 27 samples by both systems. Bactec detected six isolates of *Streptococcus pneumoniae* which Sentinel did not; Sentinel detected three isolates which Bactec did not; 16 isolates of *S pneumoniae* were detected by both systems. In the case of one patient with *S pneumoniae* infection a Sentinel bottle did not flag a positive, although the Bactec bottles did. However, when the growth curve of that Sentinel bottle was examined, it showed evidence of growth in the period before 12·5 hours. It is presumed that the organisms grew before and during this period and had become non-viable before the 12·5 hour minimum time for a positive result had elapsed.

There was a slightly higher false positive rate with Sentinel than with Bactec, probably due to operator error in incorrectly venting the Sentinel aerobic bottle.

It was difficult to determine accurately the time to positivity as this depended on the time of collection of the blood sample being recorded. Moreover, because the algorithm was deliberately configured so that it was not possible for the Sentinel to detect positive results during the first 12·5 hours, the Bactec had an inherent advantage during this period. The Bactec was indeed faster in detecting positive results up to the first 12 hours of the detection period. Since completion of the evaluation, the manufacturers of the Sentinel system have refined the algorithm to allow positive results to be detected at a minimum of 2·5 hours.

The Sentinel system had slightly fewer con-

### Table 2  Time to detection of clinically important isolates for each system

<table>
<thead>
<tr>
<th>Time</th>
<th>Bactec</th>
<th>Sentinel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Days</td>
<td>0-6</td>
<td>6-12</td>
</tr>
<tr>
<td>Hours</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bactec:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of strains isolated</td>
<td>44</td>
<td>7</td>
</tr>
<tr>
<td>Percentage cumulative positive</td>
<td>20</td>
<td>23</td>
</tr>
<tr>
<td>Sentinel:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of strains isolated</td>
<td>1</td>
<td>28</td>
</tr>
<tr>
<td>Percentage cumulative positive</td>
<td>0·5</td>
<td>13</td>
</tr>
</tbody>
</table>

*Bactec and Sentinel detected 84% and 87% of the total number of clinically important isolates.
taminants than Bactec, perhaps representing the non-invasive monitoring technique of Sentinel.

When two blood culture systems are evaluated it is usual that a greater volume is taken than for one system and subsequently more isolations are made. The average volume of blood inoculated into both systems was 14.3 ml, less than the 20 ml recommended. On average, a slightly higher volume of blood was introduced into the Sentinel bottles. Because the yield of organisms increases with the volume of blood introduced, it seems likely that both systems would have performed better if the recommended amount of blood had been introduced.

The continuous monitoring feature of Sentinel offers advantages in that the samples flagged as positive overnight are known immediately at the start of the day, enabling speedier reporting.

Continuous monitoring should also increase the speed of detection, although laboratory logistics may need to be changed to make full use of this technology. For example, the best scenario would be for blood culture bottles to be entered into the system immediately on receipt in the laboratory, including during the night, rather than batching samples, as is commonly practised at present with other methods.

The only potential safety hazard was the use of a venting needle in the Sentinel aerobic bottle. Electricity was supplied to the instrument drawers at a maximum of 24 volts which was considered a good safety feature.

The Sentinel software is menu-based and controls the data collection and analysis function of the system; it also incorporates a data search facility for current and stored samples. The system was user friendly and was liked by staff involved with the evaluation.

The Sentinel system is a useful addition to the methods available for the detection of bacteria in blood.

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