Semiautomated method for quantitative urine culture

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Examination of urine must be the most frequently performed task in any bacteriology laboratory with a large service commitment. Generally, an attempt is made to determine the numbers of viable bacteria present in the specimen, and a variety of methods have been used to this end. At the time of publication of his papers in the 1950s,1-3 from which the criteria for interpretation of bacterial counts in urine samples were defined, Kass used the agar pour plate method. Hoeprich, in 1960,4 introduced the use of the calibrated loop for making a surface count of organisms on agar media (this method had originally been described in 19285 as a means of enumerating bacteria in samples of milk). In 19646 Leigh and Williams introduced the blotting paper strip method. Finally, in 19657 Mackey and Sandys took the business of inoculation outside the laboratory when they published their paper outlining the use of dipspoons.

All of these methods remain in use. Some modifications have been made, most especially in the method of Mackey and Sandys with the introduction of commercial "kits," some bearing two types of indicator medium and others using material other than agar as a support.8 We do not wish to discuss the merits or demerits of the breakpoints commonly used in counting bacteria in urine. Discussions concerning such breakpoints, however, are valid only when they are founded on results which do not vary from operator to operator (and from laboratory to laboratory). In this paper we present a method which we believe approaches this ideal more closely than others currently in use.

Material and methods

A Hamilton MicroLab M programmable diluter (VA Howe and Co, London) (Fig. 1) was programmed to perform the sequence of functions shown in the Table. Urine specimens are mixed by inversion. Seventy microlitres is withdrawn and 60 μl is dispensed into one well of a microtitre tray for estimation of the number of cells using an inverted microscope.9 The remaining 10 μl of urine is dispensed with 990 μl of diluent (sterile peptone) into a 1:5 ml autoanalyser cup (Sarstedt Limited, Leicester); the vortex produced achieves mixing automatically. Then 120 μl of the diluted sample is taken up and dispensed as six 20 μl drops on to a cysteine lactose electrolyte deficient (CLED) agar plate in the pattern shown in Fig. 2. The sixth drop is spread using the tip of the dispenser. All the manipulations above are performed using the automatic diluter.

Program for the Hamilton Microlab M

<table>
<thead>
<tr>
<th>No.</th>
<th>Instruction</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Automatically pick up 1000 μl peptone</td>
</tr>
<tr>
<td>2</td>
<td>Automatically deliver 1000 μl peptone</td>
</tr>
<tr>
<td>3</td>
<td>Manually pick up 70 μl specimen</td>
</tr>
<tr>
<td>4</td>
<td>Manually deliver 60 μl specimen to microtitre tray</td>
</tr>
<tr>
<td>5</td>
<td>Automatically pick up 990 μl peptone</td>
</tr>
<tr>
<td>6</td>
<td>Manually deliver 1000 μl diluted specimen</td>
</tr>
<tr>
<td>7</td>
<td>Manually pick up 120 μl diluted specimen</td>
</tr>
<tr>
<td>8/9</td>
<td>Deliver six 20 μl drops separately; five to one side and the last to the opposite side of a CLED plate (spread the latter)</td>
</tr>
<tr>
<td>10</td>
<td>Automatically pick up 500 μl peptone</td>
</tr>
<tr>
<td>11</td>
<td>Manually pick up 500 μl chloros</td>
</tr>
<tr>
<td>12</td>
<td>Manually deliver 1000 μl of peptone/chloros</td>
</tr>
</tbody>
</table>

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The plates are incubated in an atmosphere containing 10% CO₂ overnight. The next morning the plates are examined. Using the spots numbered 1 to 5 a count is made using an illuminated electric colony counter (Scientifica and Cook Electronics Ltd) (Fig. 3). A total count of colonies in all five spots is made. If the count reaches a value over 100 then it is not taken any further and a value of >100 is recorded; otherwise the actual count is recorded. The total volume of diluted urine sampled in the five spots is 100 μl. The dilution of the specimen was 1/100 so that the five spots are equivalent to a 1 μl sample of the original specimen. Thus each colony present is equivalent to $1 \times 10^6$ colony forming units per litre (cfu/l)—counts of more than 100 colonies represent “significant growth” ($>10^6$ cfu/l) and counts of between 10 and 100 colonies represent “questionable growth” ($10^7$–$10^6$ cfu/l). Spot number
6, which has been spread, allows more certain detection of mixed cultures and allows sampling of single colonies. By examining "dummy" specimens (sterile peptone water) with this method immediately after positive specimens, carry of bacteria from one sample to the next has not been found to occur.

Where an automatic diluter is not available it is possible to employ an automatic pipette (for example, Eppendorf, Germany) using a separate disposable tip for each specimen. In this case, dilutions are more conveniently made in larger volumes (60 μl in 6 ml) in plastic or glass containers. The disinfection step using hypochlorite is omitted.

Discussion

Each method has its own disadvantages and in use there will be a balance between ease of use and accurate enumeration of bacteria. The pour plate method requires considerable skill and colonies may have to be retrieved from the depths of the agar. The loop method of Hoeprich appears to be very simple. Experience shows, however, that to obtain a really even spread requires considerable effort. Much variation may occur between operators, depending on the zeal with which they spread. The blotting paper strip method of Leigh and Williams is quick and is economical of media. Because of the small surface area inoculated, however, examination of specimens containing a mixed population of bacteria may not be easy. Some laboratories have moved towards overcoming some of these difficulties by combining the last two methods: the blotting paper strip method is used as a screen for specimens containing no leucocytes (or only small numbers) and the loop method is reserved for specimens containing leucocytes in numbers exceeding a locally chosen level.

The method that we have outlined for counting bacteria is now in routine use on the urine bench in the microbiology department at Cheltenham. No extra laboratory technician time has been required in order to incorporate this method—the time taken to set up each culture plate is about 30s. As far as cost is concerned, the major expense is the purchase of the programmable diluter (about £2400 excluding VAT). In addition, there will be an increase in the amount of agar used (the extent depending on the currently used method).

In principle, this method is the same as that described by Miles and Misra in 1938.10 It provides a system for obtaining counts of bacteria in urine which strikes a more satisfactory balance between accuracy and ease of operation. As mentioned above, we have not set out to discuss the validity of the "breakpoints" commonly used in enumeration of bacteria in urine specimens. If, however, there is to be useful discussion on this topic results should be subject to as little variation as possible. A system such as the one presented here should be of value in standardising the results obtained without adding greatly to the time taken to examine each specimen.

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References


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