A SIMPLIFIED PLATE METHOD FOR DETECTING GELATINE-LIQUEFYING BACTERIA

BY

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Frazier (1926) described a plate method of testing for gelatine liquefaction. The gelatine was incorporated in a low-peptone agar medium. After growing the organism on the medium, the occurrence of digestion was shown by treating the plates with acid mercuric chloride or tannic acid. Frazier claimed that his method was more sensitive and gave quicker results than the usual gelatine “stab” method.

Barer (1946) applied Frazier’s method to freshly isolated bacteria from human material (chiefly intestinal Gram-negative bacilli) and found it satisfactory. Smith, Gordon, and Clark (1946) mention a simplification of Frazier’s method which they used in examining aerobic sporing bacilli. Gelatine was incorporated in a concentration of 0.4% in nutrient agar instead of in a specially prepared basal medium. Evans and Wardlaw (1952) tried a similar method, incorporating 0.5% gelatine in non-nutrient agar, for measuring the gelatinase activity of B. subtilis culture filtrates, but found it unsuitable for accurate measurement.

This paper reports an investigation of the simplified method (with a higher concentration of gelatine) which has been used in this laboratory for testing some bacteria of medical importance. The results have been compared with those of stab tests done at the same time.

Method

The following solutions are prepared: (1) 10% leaf gelatine (w/v) in distilled water is sterilized by steaming for 30 minutes on three successive days and stored at 4-6°C. (2) HgCl₂, 15 g., is dissolved in 20 ml. concentrated hydrochloric acid, and water added to make 100 ml.

For use, the gelatine is melted in the 37°C. water-bath. One volume is mixed with 9 volumes of infusion base nutrient agar and plates poured and allowed to set and dry. The organism is streaked or spotted on the surface. Six cultures may be tested on one 3½-in. plate. The plate is incubated at 37°C. for two or three days; two days is usually sufficient for intestinal Gram-negative bacilli. If desired, two plates may be inoculated and incubated for different times. The growth is wiped off with cotton-wool moistened in the HgCl₂-HCl solution with which the plate is then flooded. Digestion is shown by a clear zone around the colonies on an otherwise densely opaque plate. The high concentration of peptone in nutrient agar does not affect the results because proteolytic organisms digest it as well as the gelatine. No change in the opacity of the medium around the colony, or a slight change but without distinct clearing, is read as negative. For swarming organisms the agar content of the medium is increased to 6%.

The gelatine-nutrient agar plate and stab methods were compared at 37°C., using a variety of bacilli (70 strains in all), most of which were freshly isolated. For this purpose each strain was inoculated into one stab and streaked on three plates. The plates were treated with precipitating fluid after one, three, and five days’ incubation respectively. The nutrient gelatine stabs were made from the same batch of gelatine.
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as the plates by a standard method (Mackie and McCartney, 1948). Inoculated stabs were tested for liquefaction by cooling in the refrigerator after one, three, five, seven, 10, and 14 days' incubation at 37°C.

The plate results were found to be the same after five days' incubation as after three days with one exception, a recently isolated V.P.-negative, indol-

and M.R.-positive coliform bacillus which was positive at five days only. This strain did not liquefy a gelatine stab incubated for 28 days.

The results of the comparison are given in the Table. Digestion usually appeared faster on plates than in stabs, but with the exception mentioned above there was complete agreement between the two methods.

Attempts were also made to compare the plate and stab methods using coagulase-positive staphylococci. With these organisms, however, the positive results were often indistinct on the plates and very slow to develop in the stabs. Moreover, the strength of the reaction given by particular strains often varied greatly when re-tested by either method. The plate method was generally the more sensitive. Barer, using Frazier's original plate method, found that results with staphylococci were unsatisfactory.

Comparison with Frazier's Method

A further series of Gram-negative bacilli was tested in parallel on gelatine-nutrient agar plates and Frazier plates. The latter were prepared as follows:

Solution (1).—NaCl, 5 g., KH₂PO₄, 0.5 g., and K₂HPO₄, 1.5 g., were dissolved in 100 ml. distilled water.

Solution (2).—Bacto-gelatine (Difco), 4 g., was dissolved in 400 ml. distilled water, and 0.05 g. dextrose, 0.1 g. bacto-peptone, and 5 ml. of beef infusion (as used for infusion broth) were then added.

Solutions (1) and (2) were mixed and heated in the steamer. Then 500 ml. of 3% (w/v) agar powder in distilled water was added, the pH adjusted to 7.0, and the whole autoclaved at 5 lb. pressure for 20 minutes.

The following organisms (with numbers of strains in parentheses) were used to compare the two plate methods: Bact. coli (14), Bact. aerogenes (8), Alkali-genes faecalis (8), paracolon strains (4), Salmonellae

<table>
<thead>
<tr>
<th>Organism</th>
<th>No. Tested</th>
<th>Plate Method</th>
<th>Stab Method</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Positive</td>
<td>Negative at</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 Day* 3 Days</td>
<td>3 Days</td>
</tr>
<tr>
<td>Bact. coli</td>
<td>15</td>
<td>15</td>
<td>5</td>
</tr>
<tr>
<td>B. aerogenes</td>
<td>5</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>B. cloacae†</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Paracolons</td>
<td>2</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Proteus vulgaris</td>
<td>10</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Proteus mirabilis</td>
<td>3</td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td>Proteus morgani</td>
<td>8</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td>Ps. pyocyanea</td>
<td>9</td>
<td>9</td>
<td>3</td>
</tr>
<tr>
<td>Shigella sonnet</td>
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<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Salm. typhi and enteritidis</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Salm. para A, B, and C†</td>
<td>3</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Salm. schlechtein†</td>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Salm. abortus bovis</td>
<td>1</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Clostridium</td>
<td>4</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Genus bacillus</td>
<td>4</td>
<td>4</td>
<td>1</td>
</tr>
</tbody>
</table>

* The figures under the day columns show the numbers of cultures which became positive for the first time on that day. Strains marked † were obtained from the National Collection of Type Cultures.
Summary

An easily prepared plate method for detecting gelatine liquefaction is described. It was compared with the standard stab method, using 70 different organisms, mostly Gram-negative bacilli, from pathological material. With one exception there was complete agreement in results between the two methods, the plate method giving the results more quickly.

The plate method was also compared with Frazier’s plate method, using 82 strains of Gram-negative bacillus. The only discrepancy in results occurred with spreading Proteus strains, which failed to hydrolyse gelatine in the Frazier plates.

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A Simplified Plate Method for Detecting Gelatine-liquefying Bacteria
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