An evaluation of 12 methods for the demonstration of penicillinase

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SUMMARY Twelve methods for the demonstration of bacterial penicillinase production by strains of Haemophilus influenzae and Staphylococcus aureus are compared, and their suitability for routine clinical laboratory use is evaluated. The acidimetric agor agar plate method is recommended.

After the discovery of penicillinase, one of the β-lactamase group of enzymes, in 1940 (Abraham and Chain, 1940) and the demonstration of its role in the penicillin resistance of Staphylococcus aureus (Kirby, 1944) many methods began to be developed for the detection and quantitation of this enzyme. In 1962, Wolff and Hamburger published an evaluation of seven methods of determining staphylococcal penicillinase which were then current. Some of the principles being employed at that time have been adopted in more recent methods but none of the procedures described remains popular today.

Interest in demonstrating staphylococcal penicillinase in the routine clinical laboratory waned as the isolation of non-penicillinase producing staphylococci decreased. In recent years, however, the emergence of penicillinase producing strains of Haemophilus influenzae and Neisseria gonorrhoeae has given a new impetus to interest in these enzymes. A large number of new, simpler, or more rapid methods have been described in recent publications, and the present paper is an attempt to evaluate some of these methods and to compare their suitability for routine use in a clinical laboratory.

METHODS

Twelve methods were used to demonstrate penicillinase production or non-production by 20 penicillin resistant and five penicillin sensitive strains of Staph. aureus and by three ampicillin resistant and six ampicillin sensitive strains of H. influenzae. The methods employed fall into four categories, viz, iodometric, pH, cephalosporin, and inhibition.

IODOMETRIC METHODS
Penicillinase hydrolyses penicillin to penicilloic acid which, in turn, reacts with iodine. The removal of available iodine from the system is demonstrated by the use of starch. The presence of penicillinase in a test system is shown by the decolorisation of a starch-iodine complex (Alicino, 1946; Perret, 1954). Many methods have been developed using this principle, three of which have been evaluated in this survey.

1 Filter paper method (Perret, 1954)
In this method the working reagent consists of a solution of penicillin G containing soluble starch to which is added an iodine solution. This reagent is poured on to filter paper, and an organism is tested for penicillinase production by smearing a portion of a culture on to the paper. Penicillinase producing organisms give a zone of clearing of the starch-iodine surrounding the smear of culture.

2 Iodine vapour method (Fleming and Markowsky, 1975)
This is similar to the previous method. Penicillin-soaked paper is blued with iodine vapour.

3 Slide method (Rosenblatt and Neumann, 1978)
A test organism is emulsified on a slide in a penicillin-iodine solution to which a starch solution is then added. The original authors found this method effective in demonstrating penicillinase production by Staph. aureus, H. influenzae, Bacteroides melanogenicus, and N. gonorrhoeae.

pH METHODS
The production of penicilloic acid from penicillin lowers the pH of the system, and this change may be demonstrated by the use of conventional pH indicators.

4 Agar overlay method (Wong and Soo-Hoo, 1976)
A test organism is subcultured on to an agar plate, to give a confluent lawn growth, and to this is
Park et al., was modified by increasing the final concentration of phenol red from 0.001% to 0.01% and by adding penicillin G in solution rather than in powder form, thus simplifying the preparation of the plates. Organisms to be tested are smeared on to the surface of the agar and incubated at 37°C or at room temperature for 15-60 minutes. Penicillinase producing strains produce a brilliant yellow zone around the bacteria.

8 Commercial strip (Intralactam, Mast Laboratories Ltd, Bootle, Merseyside, UK) (Slack et al., 1977) Commercially prepared paper strips containing penicillin and bromocresol purple are moistened and smeared with test organism. A colour change from purple to yellow indicates β-lactamase production. This method is regarded as suitable for testing only Gram-negative organisms.

**CEPHALOSPORIN METHODS**

The chromogenic cephalosporin developed by Glaxo Research Ltd, code named 87/312, (3-(2,4 dinitrostyryl)-(6R, 7R)-7-(2-thienylacetamido)-ceph-3-em-4-carboxylic acid, E isomer), has the property of showing a distinct colour change from yellow to red when hydrolysed by a wide range of β-lactamases. This characteristic has been employed in a number of methods suitable for diagnostic use. The methods evaluated share a common simple buffered solution of 87/312 as the sole reagent (O'Callaghan et al., 1972; Glaxo Research Ltd, Package Insert).

9 Plate method

The reagent is dropped onto colonies growing on the original isolation plate. Colour develops in and around the colonies of penicillinase producing organisms.

10 Broth method

The reagent is added to a broth culture.

11 Cell suspension method

A saline suspension of the organism is tested by the addition of drops of the colour reagent.

A fourth cephalosporin method which was not evaluated is the broken cell method, in which, it is claimed, low concentrations of enzyme can be detected by testing a cell lysate.

**INHIBITION**

Some of the earliest methods for the demonstration of penicillinase, the Gots Test (Gots, 1945) and the Haight and Finland modification of it (Haight and Finland, 1952), use penicillin to suppress the growth of an indicator organism which is enabled to grow if penicillinase is produced by the test organism.
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12 Inhibition
A recent modification (Hodge et al., 1978) of this penicillin inhibition principle uses a plate seeded with a penicillin sensitive indicator organism on which is placed a penicillin disc and a radial streak of the test organism. Penicillinase production by the test organism distorts the sensitivity zone.

Results
Each of the 12 methods was tested against 20 penicillinase producing and five non-producing strains of *Staph. aureus* and against three penicillinase producing and six non-producing strains of *H. influenzae*. These were compared on the basis of reliability, versatility, sensitivity, reproducibility, readability, ease and speed of performance of test, ease of preparation of reagents, relative cost, and overall suitability for routine clinical use.

Reliability
In all methods tested, penicillinase producing strains of *H. influenzae* gave stronger and/or more rapid reactions than *Staph. aureus* strains.

Method 3 (iodometric slide) gave reactions that I found to be pale and difficult to interpret. None of the other methods gave any false-positive reactions. False-negative reactions were observed in four methods with some strains of *Staph. aureus*. Methods 4 (agar overlay), 11 (cephalosporin cell suspension), and 12 (inhibition) each showed one of the 20 penicillinase producing strains to be negative, and method 5 (rapid paper strip), two of 20 strains. Method 12 also gave a weak reaction with a further nine positive strains.

Method 8 (commercial strip) is recommended by the manufacturer for Gram-negative β-lactamase testing only and was shown to be unsatisfactory for testing staphylococci.

Method 9 (cephalosporin plate) also gave a poor reaction with penicillinase producing staphylococci.

Versatility
With the exception of methods 3 (iodometric slide), 8 (commercial strip), and 9 (cephalosporin plate), all methods were found to be suitable for the demonstration of penicillinase production in both the species tested.

Sensitivity
Prolonged incubation was found to be necessary with some strains of *Staph. aureus* when tested by four of the methods. Method 6 (modified one-minute tube) required overnight incubation with eight strains tested, method 8 (acidometric agar plate) with six strains, and method 10 (cephalo-

sporin cell suspension) required 72 hours to react with two strains. All positive reactions produced by strains of *H. influenzae* were rapid, in many cases being clearly readable in less than 1 minute.

Reproducibility
A high degree of reproducibility was seen in all methods except method 3 (iodometric slide).

Readability
Methods 1 (filter paper) and 2 (iodine vapour) often gave zones that were difficult to interpret with strains of *Staph. aureus*. Since all cultures, penicillinase producing and non-producing, tend to give a zone of clearing after a period of several minutes, careful comparison with positive and negative controls is essential.

The readability of the pH methods was very good with a most distinctive colour change, in most cases, from red to yellow. The methods requiring subculture of organisms were particularly good in this respect. Methods 10 (broth) and 11 (cell suspension), using the cephalosporin, were also very distinctive but slow in changing with staphylococci. Method 12 (inhibition) gave a readily readable result in most cases. In contrast, the cephalosporin plate method (9) was good only with haemophilus cultures, the red colour being quickly and clearly visible even on chocolate (heated blood) agar.

Ease and Speed of Performance
Methods 4 (agar overlay), 5 (rapid paper strip), 10 (cephalosporin broth), and 12 (inhibition) all require subculture of the organism isolated, with consequent delay in the determination of penicillinase production.

The iodometric methods (1, 2, and 3), cephalosporin methods (9, 10, and 11), and the pH tube method (6) all require reagents to be prepared immediately before use, although the actual testing is quick. Methods 7 (acidometric agar plate) and 8 (commercial strip), however, are entirely prepared in advance and require only the addition of the test organism to the prepared plate or moistened strip.

The use of iodine vapour in method 2 made the actual testing rather cumbersome in comparison with other methods.

Ease of Preparation and Relative Cost of Reagents
All methods (except 8, commercial strip) require some prior preparation of reagents, none of which is at all complex or time consuming although demanding some technical expertise (as in weighing 5 mg of cephalosporin). Many of the stock reagents, once prepared, can be stored for some time. The most
unstable reagent in my hands was the iodometric slide method working solution which decolorised in only a few minutes with the consequent need for a fresh solution to be prepared part way through the testing of a batch of cultures.

With the exception of the cephalosporin methods all the materials needed are inexpensive and readily available. The chromogenic cephalosporin is not commercially available.

The most expensive method must be the commercially prepared strip.

Discussion

In a clinical microbiology laboratory, apart from the epidemiological importance, the main significance of penicillinase production is directly related to the degree of urgency for the institution of specific antimicrobial therapy in a given patient. Standard antimicrobial sensitivity tests by diffusion methods require overnight incubation while automated procedures take several hours. A rapid reliable penicillinase test may make possible the early implementation of appropriate therapy.

Taking into consideration the criteria discussed above, one method stands out as meeting this need, the acidometric agar plate method (Figure). The preparation of plates is extremely simple; agar base may be prepared in quantity and stored for at least six months. When penicillin has been added to the base the poured plates are also relatively stable and have proved reliable after more than four weeks of storage at 4°C. Since the medium is non-nutrient there is no problem of bacterial contamination or deterioration to interfere with the results. Should spontaneous hydrolysis of penicillin occur during the storage of prepared plates, this will be shown by a colour change of the indicator. In use, a prepared plate may be taken from the refrigerator and directly inoculated with test organisms. As many as 14 cultures could be tested on a single 90 mm petri dish. Positive results are normally apparent within 15-30 minutes, but, unlike in some other methods, there is neither fading of positive results nor false-positive reactions on prolonged incubation; the colour zone tends to enlarge with time but, provided that inocula are not too close together, both positive and negative results remain clearly readable after 48 hours. After the completion of this study the acidometric agar plate method was adopted in this laboratory for the routine testing of all H. influenzae isolates. At the time of writing, a further 141 strains have been examined, 136 of which were ampicillin sensitive and negative for penicillinase production by this method, and five which were ampicillin resistant and penicillinase producing.

In every way the acidometric agar plate method appears ideal for the demonstration of bacterial penicillinase in the clinical microbiology laboratory.

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References


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