Modified Spot CAMP Test: A rapid, inexpensive and accurate method for identification of group B streptococci

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SUMMARY
A rapid modified spot CAMP test using 183 clinical isolates of β haemolytic streptococci was compared with the standard CAMP test described by Christie et al. The scheme of biochemical identification and serological confirmation was taken as reference method. The sensitivity of both tests was 100%, and the specificity of the rapid and standard tests was 96.8% and 88.9% respectively. The modified spot CAMP test is a rapid, inexpensive and accurate method for the identification of group B streptococci, and is more specific than the standard CAMP test.

Key words: spot CAMP test, Streptococcus agalactiae

INTRODUCTION
Infections in adults and neonates caused by group B streptococci are associated with high morbidity and mortality, especially in cases of neonatal septicemia and meningitis (6). Prompt and accurate identification of group B isolates will hopefully lead to more specific therapy and improved prognoses.

Although serological grouping is the test of choice for identifying group B streptococci, presumptive tests are often used by clinical laboratories due to their relative simplicity and cost effectiveness.

The CAMP test, originally described by Christie et al., is often selected from several presumptive tests because it requires minimal reagents and a simple inexpensive methodology (1).

In the CAMP test group B streptococci produce a factor (CAMP factor) that acts synergistically with staphylococcal beta-hemolysin (β-lysin) on sheep erythrocytes. Briefly, the traditional test consists of a single straight streak of the studied Streptococcus agalactiae and a beta-lysin producing Staphylococcus aureus strain. They must be perpendicular, with some millimetres in between. This test must be done on a sheep blood agar plate and requires overnight incubation. An attempt to standardize the CAMP test was made by Darling, but in an effort to make the standard CAMP test more rapid or simple or both, several modifications have been proposed (9,11). Kaplan et al (5) described a 20-minute spot CAMP test whose reagent is a crude beta-lysin-containing filtrate derived from a broth culture of S. aureus. The spot CAMP test can be performed directly on the blood sheep plate used like primary isolation medium, even when the colony(s) has been removed from the agar surface for other studies.

In order to standardize the spot CAMP test, Ratner et al (10) compared this test with the standard overnight one, using 350 clinical isolates of beta haemolytic streptococci.
They found 99% and 100% of sensitivity and specificity respectively.

Consequently, the Clinical Laboratory Standards Institute (ex National Committee Clinical Laboratory Standards) recommended the spot CAMP test for rapid identification of group B streptococci (7). However, the Kaplan test could not be used in a primary isolation medium different from the blood sheep agar, or in polymicrobial samples.

In this study, a new modification of the spot CAMP test was analysed, which consisted in performing the spot and develop the rapid test on the same day of the isolation.

We assessed the sensitivity and specificity of the modified spot and the standard overnight CAMP test for identification of *S. agalactiae*. Biochemical identification and serological confirmation were considered as reference method.

**MATERIALS AND METHODS**

Based on the prevalence of isolation at our laboratory (2), 183 beta haemolytic streptococci isolates (45 B group, 6 C group, 6 G group and 126 A group) were identified by conventional method, as described by Murray *et al* (6) including the serologic test confirmed by Lancefield precipitin test.

The standard CAMP test was performed as described by Darling (3). A streak of *S. aureus* ATCC 25923 (a known strain that produces a high level of beta toxin) was done in agar sheep blood plate. The strains of streptococci to be tested were streaked in a straight line 2 to 3 cm in length and at a right angle to these inoculum, with care not to touch the staphylococcal streak. The plate was incubated at 37 °C for 18 hours, and a haemolytic zone observed in “arrowhead” was considered as positive result.

**Spot CAMP reagent preparation**

The spot CAMP reagent was prepared inoculating two 5-ml tubes of brain-heart infusion broth with a small amount of growth from a fresh subculture of *S. aureus* (ATCC 25923), and incubated overnight at 35 °C in ambient atmosphere.

The two broth suspensions were mixed and sterilized by filtration in a laminar-flow biological safety cabinet using 0.45-μm cellulose–acetate filter.

Portions of 1 ml of the filtered-sterilized broth (spot CAMP reactive) were aliquoted into small tubes, with six-month expiration date when stored at –20 °C or lower temperature (7).

**Modified spot CAMP test**

Spot of microorganisms was done in agar sheep blood plate, and was incubated at 35 °C at 5% CO2-enriched atmosphere for at least 4 hours or until growth and haemolysis were evidenced.

One drop or 10-μl loopful of spot CAMP reagent was placed next to a suspected group B streptococcal spot. The plate was incubated at 35 °C in ambient air (right side up to prevent the spot CAMP reagent from running over the plate surface) for 20 minutes.

Transmitted light was used to detect a zone of enhanced haemolysis next to the spot. Initially negative reactions were reincubated for up to 30 minutes.

The presence of a clear zone of enhanced haemolysis was considered a positive reaction.

A negative reaction was the lack of haemolysis near to the spot of microorganisms in the presence of the spot CAMP reagent (staphylococcal haemolysis).

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**RESULTS**

All the *S. agalactiae* isolates were positive by spot CAMP test and standard CAMP test (sensitivity 100%). Only 4 Streptococcus pyogenes were positive by spot CAMP test (specificity 96.8%), and 14 by the standard method (specificity 88.9%) (Table 1) (p<0.01).

The reading time of the spot CAMP test need not be prolonged at any time for identification of *S. agalactiae*.

The stability of sheep agar plates used for the test was 2 months at 4 °C.

**DISCUSSION**

Since certain conditions should be met for the satisfactory performance of the CAMP test, many authors have evaluated it using tube or disk (9,12). However, the original CAMP test described by Christie *et al* (1) is commonly used in clinical laboratories because of its simplicity.

The NCCLS (7) suggests the use of the spot CAMP test described by Kaplan *et al* (5) for rapid identification of *Streptococcus agalactiae*, since this test has been previously validated by Ratner *et al* (10). Moreover, it would be very useful to make the primary isolation on a sheep blood agar plate.

In this study, we suggest that the modified spot CAMP may be done on the same day of primary isolation.

The stability of the sheep blood agar plate is another advantage of this test, which is an inexpensive option for routine diagnosis of *S. agalactiae* infections, and could be repeated on the same board during a month.

Some studies have demonstrated that *S. pyogenes* have a positive CAMP reaction under certain conditions,

<table>
<thead>
<tr>
<th>Microorganisms</th>
<th>$N$</th>
<th>CAMP test positive</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>“Spot” n/N (%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Standard n/N (%)</td>
</tr>
<tr>
<td><em>S. agalactiae</em></td>
<td>45</td>
<td>45/45 (100)</td>
</tr>
<tr>
<td><em>S. pyogenes</em></td>
<td>126</td>
<td>4/126 (3,2)</td>
</tr>
<tr>
<td>Streptococcus group C</td>
<td>6</td>
<td>0/6 (0)</td>
</tr>
<tr>
<td>Streptococcus group G</td>
<td>6</td>
<td>0/6 (0)</td>
</tr>
</tbody>
</table>

(1) n/N, number of positive reactions / total analysed.
particularly in anaerobic incubation (4,11). In our study, 14 *S. pyogenes* isolates were positive with the standard CAMP test, but not with the modified spot CAMP test.

However, the synergy seen in the system of positive CAMP in *S. pyogenes* reaction has been long considered a false positive reaction caused by small parts of streptolysin O unoxidized (11).

Notwithstanding that, Gase et al (4) demonstrated that *Streptococcus pyogenes* possess the gene that encoded an extracellular CAMP factor capable of participating in the positive reaction. In most cases, the haemolysis produced by these strains of *S. pyogenes* was less than that produced by *S. agalactiae*.

The time around in the modified spot CAMP test could have been insufficient evidence to this test of the haemolytic effect of low level CAMP factor of *S. pyogenes*.

Therefore, the modified CAMP test proposed has proved to be more specific than the standard; it could be done on the same day of microorganism isolation, and allowed to test several isolates on a single sheep blood agar plate. Thus, we consider this test a practical, accurate and inexpensive method for the identification of beta-haemolytic streptococci at any clinical laboratory.

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REFERENCES


