The rapid recognition of Lancefield group B haemolytic streptococci

R. J. FALLON

From the Department of Laboratory Medicine, Ruchill Hospital, Glasgow

SYNOPSIS When grown overnight on Columbia agar in an atmosphere of hydrogen and carbon dioxide, haemolytic strains of Lancefield group B streptococci (Streptococcus agalactiae) produce orange or brown pigmented colonies. This production of pigmented colonies can be used for the rapid presumptive identification of these organisms as belonging to group B without the need for grouping by serological methods.

The production of pigmented colonies by streptococci has received little attention since first described by Orla Jensen (1919). He observed that red colouration was produced by several strains of pathogenic streptococci in casein peptone agar and noted that 'Streptococcus mastidis' (now known as Strep. agalactiae) formed an orange colour in casein peptone broth with added soluble starch. Durand and Giraud (1923) published a study of chromogenic streptococci in which they noted the importance of starch as a medium constituent and the maintenance of anaerobic conditions as factors necessary for the production of pigment. Eleven of the 125 strains they examined produced pigmented colonies including strains which were clearly pathogenic as well as some which they regarded as saprophytic. With one exception all strains came from human sources and appeared to be serologically related.

The isolation of a pigmented strain of streptococci from a case of pyelonephritis was reported by Olivieri (1929). Lancefield (1934) described studies on a strain of group B haemolytic streptococci which formed a yellow-brown pigment. In passing she commented that other strains produced pigment including nine of 29 strains classified in groups B and D and that pigment was only apparent under conditions of partial or complete anaerobiosis.

Plummer (1941) in a study of 522 cultures of haemolytic streptococci found that 128 of 187 strains of group B produced pigment. Pigment production was also seen in one of group D strains examined and noted in her table I (but not in the text of the paper) and in one of 28 strains which were not grouped. Apart from a passing reference by Esseveld, Daniëls-Bosman, and Leijnse (1958) there appear to have been no other reports on this subject.

Materials and Methods

VAGINAL SWABS These were taken with charcoal-impregnated cotton wool swabs and were transferred to the laboratory in Stuart’s (1959) transport medium.

ORGANISMS In addition to strains isolated in this laboratory, cultures of the four serotypes of group B streptococci were kindly provided by Dr M. T. Parker; in addition non-haemolytic strains of group B were provided by Dr H. W. Wilkinson. These latter strains were described by Wilkinson, Thacker, and Facklam (1973).

MEDIA The media used were Oxoid Columbia agar; blood agar made by adding 5% horse blood to Columbia agar base; Oxoid nutrient agar; and Oxoid nutrient broth.

Results During the routine examination of blood agar plates inoculated with vaginal swabs from patients seen either for gynaecological or antenatal examination, it was noted that some plates contained haemolytic colonies which were pink in appearance. On Gram film examination of these colonies it was found that they were streptococci. They did not appear to belong to serogroups A, B, C, D, F, or G and hence were forwarded to Dr M. T. Parker at the Central
Public Health Laboratory, Colindale, for grouping. He reported that all the strains forwarded were of group B and included serotypes Ia, Ib, II, and III. Our failure to recognize them was due to using extraction with 0-2N HCl at 100°C instead of at 55°C in the Lancefield grouping technique. These apparently pink colonies were only seen on plates incubated anaerobically.

**Proof of Pigmentation**

An experiment was performed to confirm that the colonies were pigmented. A sterile Millipore membrane was placed on a blood agar plate and the streptococci were streaked across it. After overnight anaerobic incubation, orange pigmentation of the colonies was seen clearly against the white background of the membrane. No pigment was seen on aerobic incubation. The reason pigmentation had not been noted previously was not clear until it was realized that the recent introduction of the GasPak system of anaerobiasis into the laboratory may provide the explanation.

**Cultural Conditions**

Cultures were set up and incubated at 37°C under the following conditions: (1) aerobically, (2) aerobically with 5-10% CO₂, (3) anaerobically in H₂ with CO₂ (GasPak system), (4) anaerobically in H₂ alone, and (5) in 90% N₂ and 10% CO₂. The results are shown in table I. Pigment was only produced overnight in cultures incubated in an atmosphere containing CO₂ and was only consistently seen in cultures incubated in the jars containing H₂ and CO₂. This was probably because the anaerobiasis produced using N₂ and CO₂ was less complete than with H₂ and CO₂ as even using the latter system, if free gas exchange between the petri dish and the atmosphere in the jar was prevented by a film of moisture between the lid and the rim of the plate, pigment production was poor or absent.

**Influence of Culture Medium**

Blood was omitted from the medium and it was found that not only was pigment still produced after overnight incubation in H₂ and CO₂, but that it was much easier to see than on blood agar. Strains varied in the depth of colonial pigmentation from very pale orange to a deep orange-red colour. The impression was gained that some strains seemed able to produce much more pigment than others. Even the poor pigment-producing colonies were distinguishable from the white colonies produced by the same strains under aerobic conditions. The experiment was repeated using nutrient agar instead of Columbia agar but no pigment was formed after overnight incubation. As Columbia agar contains starch, 0-1% soluble starch was added to nutrient agar and it was found that pigment was produced on this medium, although not as well as on Columbia agar. Hence, starch appeared to be a significant component of the medium. The observation of Lancefield (1934) that pigment was produced on plain agar containing less than 1% dextrose was not confirmed using Oxoid nutrient agar (table II). Unlike the observation of Plummer (1941) no pigment was seen in broth culture even in the presence of added starch.

<table>
<thead>
<tr>
<th>Medium</th>
<th>Pigment Production</th>
</tr>
</thead>
<tbody>
<tr>
<td>Columbia agar</td>
<td>++</td>
</tr>
<tr>
<td>Nutrient agar</td>
<td>-</td>
</tr>
<tr>
<td>Nutrient agar plus 1% dextrose</td>
<td>-</td>
</tr>
<tr>
<td>Nutrient agar plus 0.1% dextrose</td>
<td>-</td>
</tr>
<tr>
<td>Nutrient agar plus 0.1% starch</td>
<td>+</td>
</tr>
</tbody>
</table>

Table II Pigment production by group B haemolytic streptococci when grown in an atmosphere of H₂ and CO₂ on various media.

**Effect of Medium pH on Pigment Production**

Occasionally batches of Columbia agar plates were encountered where pigment production by group B streptococci was poor. In order to investigate this, plates were prepared at different levels of pH. Pigment production by colonies was poor on medium at pH 7 but was easily seen at pH 7-3 or above. It is important, therefore, to ensure that the medium pH is not below 7-3.

**Examination of Known Group B Strains**

Nineteen strains supplied by Dr M. T. Parker were examined. Seventeen of these were haemolytic and included serotypes Ia, Ib, II, and III. The results of examination for pigment production are shown in table III. Other non-haemolytic strains (Wilkinson et al, 1973) were also examined and the results are also shown in table III. It can be seen that non-haemolytic strains consistently failed to produce pigment.

<table>
<thead>
<tr>
<th>Atmosphere</th>
<th>Pigment Production</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>-</td>
</tr>
<tr>
<td>Air plus 5-10% CO₂</td>
<td>-</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>-</td>
</tr>
<tr>
<td>Hydrogen plus CO₂ (GasPak)</td>
<td>+</td>
</tr>
<tr>
<td>Nitrogen plus 10% CO₂</td>
<td>±</td>
</tr>
</tbody>
</table>

Table I Effect of atmosphere on pigment production by group B haemolytic streptococci grown on Columbia agar.
Use in the Diagnostic Laboratory

In view of these observations, all vaginal swabs were inoculated onto Columbia agar which was incubated in H₂ and CO₂ as well as onto the media in routine use in this laboratory. All β haemolytic streptococci seen on the blood agar plates were grouped and the Columbia agar plates were examined for pigmented colonies. Where pigmented colonies occurred, they were present in virtually the same numbers as the haemolytic colonies on the blood agar plate inoculated from the same specimen except on the rare occasion when two different types of haemolytic colony were seen due to the presence of more than one group of streptococci in the specimen. When pigmented colonies were seen, they were always of group B. Over 120 strains of group B haemolytic streptococci so far examined produce pigmented colonies, whereas strains of haemolytic streptococci of groups A, C, G, and F isolated from vaginal swabs have consistently failed to produce pigment. None of 18 haemolytic strains of Strep. faecalis produced pigment.

A prospective study to determine what proportion of strains of group B haemolytic streptococci produce pigment gave the results shown in table IV.

<table>
<thead>
<tr>
<th>No. of Strains</th>
<th>Pigment Production on Primary Isolation</th>
</tr>
</thead>
<tbody>
<tr>
<td>54 (85.7%)</td>
<td>+</td>
</tr>
<tr>
<td>9 (14.3%)</td>
<td>-</td>
</tr>
<tr>
<td>Total 63</td>
<td></td>
</tr>
</tbody>
</table>

Table IV Pigment production by group B haemolytic streptococci on primary isolation

It will be seen that some group B strains did not produce pigment on first isolation. This may have been due to the fact that they were unable to produce pigment, that the medium pH was unsatisfactory or that, where unvented plastic petri dishes were used, proper anaerobic conditions were not obtained due to moisture forming a seal between the plate lid and rim.

Discussion

The work described confirms earlier observations that group B streptococci frequently are able to produce pigmented colonies under anaerobic conditions in a medium containing starch. Pigment may be produced on other media without added starch either to a lesser extent, eg, Hartley digest agar as made by the Streptococcal Reference Laboratory, Colindale, or after a longer period of incubation so that the requirement of starch for pigment production is probably relative rather than absolute. The requirement for the presence of carbon dioxide has not previously been noted nor has the influence of pH on pigment production. The failure of non-haemolytic strains of group B streptococci to produce pigment agrees well with the observation of Lancefield (1934) that when the strain of group B streptococcus which she was investigating lost its haemolytic properties, it also lost the property of pigment production. Significantly, however, this strain did not lose its virulence so that it is unfortunate that the present system will not assist in the detection of non-haemolytic group B streptococci in situations where these may be acting as pathogens. Under the name of Strep. agalactiae group B streptococci have long been recognized as having pathogenic potential being the cause of bovine mastitis (which may assume epidemic proportions). Following the description of group B by Lancefield (1933) occasional reports of the isolation of group B streptococci from human disease were made but it was not until the study of Hood, Janney, and Dameron (1961) was published that the significance of group B streptococci in perinatal morbidity and mortality in man was appreciated. Since that time other workers have confirmed and extended these findings (Eickhoff, 1972; Baker, Barrett, Gordon, and Yow, 1973; Barton, Reigin, and Lins, 1973; Francisoi, Knoostman, and Zimmerman, 1973). It now seems that it is important to recognize group B streptococci from the human genital tract so that decisions may be taken regarding any action or surveillance to be undertaken on any given patient. The method described enables such recognition to be made easily and rapidly without the need, in many instances, for time-consuming and expensive definitive serological grouping of Strep. agalactiae. Hence, in routine diagnostic practice, a vaginal swab is inoculated onto a half plate of Columbia agar which is incubated overnight in an atmosphere of H₂ and CO₂ together with a blood agar plate. If orange pigmented colonies are seen on the Columbia agar plate and typical large, butyrous β haemolytic colonies on the blood agar plate they can be identified presumptively as belonging to Lancefield's group B. If there are
several types of haemolytic colony present it is naturally important to identify the strain not producing pigment by further examination using routine methods. The procedure can be applied to β haemolytic streptococci isolated from other sites and in this instance many strains can be streaked on one Columbia agar plate, any pigmented strains being presumptively identified as group B.

Finally, it is important to note that the pigment may not be visible if plates are examined in fluorescent light. The pigment is best seen in daylight or in light from a tungsten filament lamp.

I wish to thank Dr M. T. Parker, for examining some of our group B strains, for providing strains of known serotype, and for supplying Hartley digest agar, and Miss E. P. Laird for secretarial assistance.

References


The rapid recognition of Lancefield group B haemolytic streptococci

R. J. Fallon

doi: 10.1136/jcp.27.11.902

Updated information and services can be found at: [http://jcp.bmj.com/content/27/11/902](http://jcp.bmj.com/content/27/11/902)

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
[http://group.bmj.com/group/rights-licensing/permissions](http://group.bmj.com/group/rights-licensing/permissions)

To order reprints go to:
[http://journals.bmj.com/cgi/reprintform](http://journals.bmj.com/cgi/reprintform)

To subscribe to BMJ go to:
[http://group.bmj.com/subscribe/](http://group.bmj.com/subscribe/)