Detection and identification of gonococcal L-Forms using a direct immunofluorescence test

SHEENA A. WAITKINS*† AND I. GEARY**

From the Neisseria Department*, Statens Seruminstitut, Copenhagen, Denmark, Department of Medical Microbiology,** University of Sheffield Medical School, Sheffield, and Central Public Health Laboratory,† London

SUMMARY A fluorescent antibody test was used to identify L-Forms of *N. gonorrhoeae* induced *in vitro*. It was possible to differentiate the Large Bodies of the L-Forms from parent gonococci and the fluorescent reaction remained specific in the presence of tissue culture cells. A possible method to identify L-Forms of gonococci from patients presenting with postgonococcal urethritis is described.

Introduction

In the last few years there has been considerable interest in the L-Forms of *Neisseria gonorrhoeae*. Their presence in the genital secretions of patients who attended with suspected gonorrhoea was demonstrated by Gnarp *et al.* (1972) and by Gnarp and Wallin (1973). Although these authors isolated L-Forms from patients who were infected with *N. gonorrhoeae*, it is difficult to determine whether these L-Forms were present in the patient, or had been accidentally induced by the cultural techniques. The causative agents in postgonococcal urethritis are still unknown, but the presence of gonococcal L-Forms was suggested by Holmes *et al.* (1967a) and in another study the same group (Holmes *et al.*, 1967b) proposed that inadequate penicillin treatment might transform the gonococcus to its L-Form. These views have been partially substantiated by the findings of Barile *et al.* (1959) who demonstrated that of seven strains of penicillin-resistant *N. gonorrhoeae*, one strain transformed very easily to a penicillin-resistant gonococcal L-Form. L-Forms may also be responsible for clinical complications—such as gonococcal arthritis. Holmes *et al.* (1971) reported a case in which gonococcal L-Forms grew from 'sterile' synovial fluid, but their findings were not substantiated by clinical evidence.

There is no longer any doubt that the gonococcal L-Form can occur under *in vitro* conditions. Lawson and Douglas (1973) gave an account of the induction and reversion of the L-Form of *N. gonorrhoeae* on cultural media but the presence of these same organisms in clinical specimens still has to be proved. This lack of proof is mainly due to inability to isolate and identify specifically the gonococcal L-Form without undertaking long and tedious laboratory techniques.

The results presented in this paper show an easy, effective, and specific method of identifying gonococcal L-Forms. Dienes (1942, 1966) described a simple staining method to illustrate the various elementary structures of bacterial L-Forms. He showed that all L-Forms consisted of both vacuolated Large Bodies (Fig. 1), and small granules growing from and/or around them. These observations have been substantiated by many workers, and it is now possible to say that in every case where bacterial L-Forms are developing, these distinct structures of the Large Body can be seen (Dienes, 1942; Weibull, 1963; Fass and Prior, 1974). In the current work, Large Bodies of developing gonococcal L-Forms were shown to be stained specifically with fluorescein-isothiocyanate (FITC) labelled rabbit antigonoococcal globulin. The combination of the distinct structure and the specific fluorescent staining was used to identify the developing gonococcal L-Form, from *in vitro* induced L-Forms.

Materials and methods

**MEDIA**

Basic Growth Medium Difco GC medium base plus 2% defined supplement (White and Kellogg, 1965) was used.

Address for reprints: S. A. Waitkins, Cross-Infection Reference Laboratory, Central Public Health Laboratory, Colindale Avenue, London NW9 5HT

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Fig. 1 Usual arrangement of structures found in mature L-Form colonies, L-Form at all stages can be seen: Large Bodies (LB) interspersed with gonococci (GC), granules (G) can also be seen inside the Large Body. Dienes' stain × 1000.

Induction Medium was prepared using the Basic Growth Medium described above and adding 10% inactivated horse serum, 500 units/ml benzylpenicillin, and 1% polyvinylpyrrolidone (PVP) with a molecular weight of 700 000 (BDH Chemicals Ltd). The Basic Growth Medium plus 1% PVP was autoclaved at a temperature of 121°C for 15 min; when it had cooled to 55°C the penicillin, horse serum, and defined supplement were added.

Reversion Medium was prepared using the Basic Growth Medium with 10% inactivated horse serum and 10% sucrose solution as an osmotic stabiliser.

Five per cent lysed blood agar was prepared by using Columbia agar (Oxoid) 39 g and distilled water 1000 ml. The agar was soaked for 15 min, mixed, and autoclaved at a temperature of 121°C for 15 min. When it had cooled to 55°C, 50 ml of sterile blood (Wellcome), lysed by repeated freezing and thawing, was added.

Organisms
Organisms from Sheffield and Copenhagen were examined. All strains of N. gonorrhoeae from Sheffield were isolated on 5% lysed horse blood agar containing 1% vancomycin (500 units/ml). They were identified by colonial form, Gram stain, a positive oxidase reaction, and the fermentation of glucose but not of maltose or sucrose. Fermentation reactions were carried out using a modified carbohydrate medium containing Difco GC medium base with 1% sugar concentrations (Flynn and Waitkins, 1972).

Isolation and identification of the Danish strains were performed by the methods recommended by Reyn (1969).

After only one subculture Kellogg types 1, 2, 3, and 4 colonies were identified and grown on Basic Growth Medium for 18 hours at a temperature of 37°C in 5% CO₂ atmosphere. The organisms were
harvested by scraping off, suspending, and diluting in saline to a final concentration of $10^8$ gonococci/ml as described by Miles et al. (1938). This suspension was used in the test systems.

**Induction of gonococcal L-Forms**

L-Forms of gonococci were induced by flooding 0.1 ml of the gonococcal suspension on to freshly poured plates of Induction Medium. A control for gonococcal viability was included by culturing on Basic Growth Medium. The inoculated plates were then incubated at 36° in a moist atmosphere containing 10% CO$_2$ and 90% humidity. The surface of the plates was inspected every day for possible L-Form colony development using a stereoscan plate microscope.

**Identification of L-Forms**

As already described, the L-Forms consist of three distinct morphological structures: the Large Body, the elementary corpuscles, and granules. Because the Large Bodies are so distinct we propose to use their presence as an indication of L-formation while using the fluorescent antibody (FA) technique in the identification of the gonococcal L-Form.

**Methods used for identification**

**Block removal technique, using Dienes' stain (1942)**

(Fig. 2)

At intervals in the development of the L-Form, blocks of agar approximately 1 cm$^2$ were removed using a sterile scalpel blade. The block was placed growth downwards on a cleaned microscope slide and fixed in situ with Bouin's fixative overnight in a moist box at 37°C. After fixation the blocks were removed and discarded, the slides were washed with distilled water to remove excess stain, and treated with 70% alcohol for 1 hour. After 1 hour the slides were washed again in water and finally stained using Dienes' stain for 1 hour.

**Fluorescent antibody technique (Fig. 3)**

The technique for fluorescence staining was initially very like that of Dienes' staining; agar blocks were cut out as above and pressed on to slides as before. The imprint that remained was then processed for the FA method.

The imprinted slide was fixed with methanol for 5 min, washed in water for 10 seconds, dried and stained with FITC-labelled rabbit gonococcal globulin for at least 30 min in a moist box at room temperature. After the specimen had been stained, excess conjugate was washed off for 10 min, then allowed to dry and finally mounted in glycerol under a cover slip.

Antigonococcal conjugate was prepared by the Lind method (Lind, 1967, 1975) and used at 1 : 16 dilution. This was made using 0.1 ml of diluted conjugate, 0.2 ml antistaphylococcal serum, and 1.3 ml distilled water.

Large Bodies were found to fluoresce very brightly using this method.
Reversion technique

In the introduction it was explained that some L-Forms are unstable and can revert to their parental form. The L-Forms of *N. gonorrhoeae* are also capable of such reversions and this phenomenon can be used in the specific identification of gonococcal L-Forms. The L-Form is allowed to revert to the parent bacterial organism on suitable non-inducing medium, and in the case of gonococcus, the medium used is similar to the Induction Medium, except that it does not contain any antibiotics and has sucrose as an osmotic stabiliser. Once reversion to the parent gonococcus has occurred, the identification of the organisms is as usual.

Method

Agar blocks were cut out of the Induction Medium and placed growth side downwards on the Reversion Medium. The agar block was then gently pushed forwards and backwards over the entire surface of the new medium, followed by incubation at 36°C in a moist atmosphere containing 10% CO₂ and 90% humidity. When reversion occurred the resulting gonococcal colony was identified by Gram's stain, positive oxidase reaction, and sugar fermentation reactions.

Tissue Culture Cells

Media

Eagle's Minimum Essential Medium (MEM) was obtained as a 10 times concentrated stock solution from Wellcome Reagents Ltd (Eagle, 1959). Eagle's Growth Medium (EGM), 100 ml of MEM, 100 ml calf serum (Biocult), and 20 ml of 4-4% sodium bicarbonate solution were added to 780 ml deionised water.

Osmotically stabilised EGM (for L-Forms) was obtained as above for EGM plus 1% sterile PVP.

Tissue culture cells

Vero cells were grown on cover slips (1 x 4 cm) inside Leighton tubes which were then incubated statically at 37°C at an angle of 5° from the horizontal to ensure that a monolayer of cells would form on the cover slips. The final tissue culture cell concentration was approximately 1 x 10⁸ cells per ml.

Organisms

*N. gonorrhoeae* were prepared as before but the final suspension of organisms was made in EGM.

Gonococcal L-Forms

Mature L-Form colonies were scraped off and washed in osmotically stabilised EGM. The colonies were concentrated by centrifugation x 5000 rev/min for 5 min, and the final concentration was adjusted to approximately 1 x 10⁴ L-Forms per ml with the same medium.

Method of infecting the cells

Cells were infected with either gonococci or their L-Forms as described by Waitkins and Flynn (1973); 0.1 ml of the suspension of organisms was added to each Leighton tube containing tissue culture cells, and then incubated at 37°C for six hours. After this time each cover slip was washed thoroughly in phosphate buffered saline (pH 7.2) to remove any adherent organisms and then stained by the FA method.

Fluorescent antibody technique for tissue culture cells

The infected tissue culture cells were first fixed in cold acetone for 5 min, and then washed in phosphate buffered saline (pH 7.2) to remove excess acetone. After the cover slips were dry they were stained with FITC-labelled rabbit gonococcal immunoglobulin G purified by diethylaminoethyl cellulose (DEAE)
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chromatography (optical density ratio 0.7) and used in a one-step inhibition test at a dilution of 1:2. The fluorescent staining technique was similar to the FA block staining method described above. The staining action took place in a moist box at room temperature for 30 min after which the cover slips were washed with water and allowed to dry in air before they were mounted in glycerol on clean microscope slides and viewed on a Zeiss standard microscope equipped for fluorescence microscopy.

Reading of FA staining results
In both the block stains and the tissue culture cells the method for reading the results was the same. The degree of fluorescence was recorded in values from 0 to ++++++. The values ++++ and ++++++ corresponded to brilliant, apple-green fluorescence. The positive reaction was characterised by the typical morphology of the organisms, the Large Body of the L-Form (Fig. 4), and the diplococci of the gonococcus (Fig. 5), and by the degree of fluorescence (+++ and ++++++).

Control organisms
The L-Form of Staphylococcus aureus strain NCTC 8530 Cowan 1 and Mycoplasma hominis were grown on the same medium as the gonococcal L-Forms and acted as controls for the specificity of the FA method. These control organisms were subjected to the same FA block procedures as the gonococcal L-Forms.

Results
As Table 1 shows, 55 strains of N. gonorrhoeae were induced to form gonococcal L-Forms. These were examined for the production of Large Bodies using Dienes' staining method and when found to possess these structures they were further examined by the FA method for specific identification of the organisms. All Large Bodies derived from gonococci were found to have positive FA reactions similar to

Table 1 Number of strains of N. gonorrhoeae examined for the production of L-Forms

<table>
<thead>
<tr>
<th>Of N. gonorrhoeae</th>
<th>Induced to form L-Forms</th>
<th>Giving a positive FA reaction + to +++++</th>
<th>Reverting to their parental types</th>
</tr>
</thead>
<tbody>
<tr>
<td>55</td>
<td>55</td>
<td>55</td>
<td>40</td>
</tr>
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</table>

Fig. 5 Typical diplococcal gonococci stained with FITC-labelled rabbit gonococcal immunoglobulin ×1000.
their parent gonococci varying in values of between ++ to ++++. Forty of the 55 strains reverted to the parent gonococcal strains and could subsequently be identified using classical bacteriological techniques. Furthermore, the gonococci reverted to their original Kellogg colonial types. Table 2 shows that this FA reaction was specific for L-Forms from gonococci only. The control L-Forms from S. aureus and M. hominis did not react with the anti-gonococcal conjugate.

Table 2  Fluorescent antibody method applied to control micro-organisms

<table>
<thead>
<tr>
<th>Micro-organisms</th>
<th>Bacterial form</th>
<th>L-Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staphylococcus aureus NCTC 8530</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Mycoplasma hominis</td>
<td>—</td>
<td>0</td>
</tr>
<tr>
<td>Neisseria gonorrhoeae</td>
<td>++++</td>
<td>++++</td>
</tr>
</tbody>
</table>

Figures 1, 6, and 7 show the distinct differences in the microscopical appearance of parent gonococci and their L-Forms. Figure 6 illustrates the classical arrangement of gonococci using Dienes' block staining technique, whereas Fig. 7 shows the one Large Body isolated on its own showing the cytoplasmic content of this structure. Figure 1 illustrates more clearly the usual arrangement of structures found in the mature L-Form colonies, L-Form at all stages can be seen; Large Bodies were interspersed with gonococci and other L-phase structures in the process of L-Form production. It is clear from these figures that the potential L-Form colonies can be recognised by the presence of the Large Body, because in all cases when Large Bodies appear typical L-Form colonies as seen in Fig. 8, will eventually develop either in days or weeks.

The specific identification of these Large Bodies can be achieved using FA methods. Figure 5 illustrates the control gonococci giving a specific positive reaction characterised by the typical diplococcal morphology of the micro-organism and the degree of fluorescence of ++++. The corresponding FA reaction of the gonococcal Large Bodies can be seen in Fig. 4, the same degree of specific reaction was characterised this time by the distinct morphological structure of the Large Body.

The FA technique was then used to identify gonococci and their L-Forms which had been
inoculated on to tissue culture cells. Vero cell sheets were prepared and inoculated with the organisms as described above; however, it was necessary to use the highly purified fluorescent conjugate because of gross non-specific fluorescence by the tissue culture cells. Using this purified conjugate the non-specific reactions from the tissue cells were eliminated while the gonococcus and their L-Forms gave a strongly positive fluorescence.

Figures 9 and 10 illustrate the resulting fluorescent reactions. Figure 9 shows the control gonococcus adhering to the tissue culture cells and Fig. 10 demonstrates the Large Body adhering to the background culture cells.

**Discussion**

Observations on the morphology of gonococcal L-Forms induced in vitro showed the presence of Large Bodies in all the cultures. They were easily identified, and are considered characteristic in the morphology of the developing L-phase cultures. The Large Body may or may not be viable in its own right, but it is certainly morphologically distinct and appears regularly whenever gonococci are induced to the L-phase; because of this we believe that it can be used as an identification structure for the presence of L-Forms.

In all strains tested, the Large Bodies gave strongly positive fluorescent reactions and were easily distinguished from the granules and other elements of the L-phase cultures.

The FA test was then applied to a mixed culture of L-Forms and gonococci, and both the Large Bodies and gonococci produced bright fluorescence. However, it was possible to distinguish the Large Bodies of the L-Forms from the parent gonococci by the distinct morphological difference. Also there was a high level of specificity in the detection of gonococci and their L-Forms as neither the mycoplasmal nor the staphylococcal L-Form was stained by the antigonococcal conjugate.

Having ascertained that it was possible to use the FA technique to identify the L-phase *N. gonorrhoeae*, we then used a highly purified conjugate to demonstrate this specific fluorescent reaction in the presence of tissue culture cells.

The strong degree of fluorescence exhibited by gonococcal L-phase Large Bodies and the absence of fluorescence from the tissue culture cells lead us to believe that this test may have potential in identifying gonococcal L-Forms in specimens from patients presenting with suspected postgonococcal urethritis.
Fig. 9  Control gonococci which have been stained by FITC-labelled rabbit gonococcal immunoglobulin adhering to Vero tissue culture cells ×1000.

Fig. 10  A gonococcal Large Body stained with FITC-labelled rabbit gonococcal immunoglobulin adhering to Vero tissue culture cells ×1000.
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References


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