Morphogenesis of Mycoplasma and Bacterial L-form Colonies

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SUMMARY

The development and mechanism of formation of Mycoplasma and bacterial L-form colonies were studied. The micro-organisms penetrate into the agar within a few hours after inoculation. It is suggested that penetration is caused by capillary forces which draw the minute plastic organisms into the dried agar gel, together with the water surrounding them. Penetration does not take place when the agar surface is very moist. The organisms appear to develop in the interstices of the fibrillar network of the agar gel and form a firm and elastic ball-like colony growing in all directions. When reaching the agar surface the growth spreads into the thin free water film which covers the agar, forming the peripheral zone typical of mature colonies. Factors which decrease the thickness of the free water film, like drying the medium or increasing the agar concentration, cause a decrease of the peripheral zone dimensions. Factors which retard growth, such as high concentrations of inorganic salts or hydrogen ions, inhibit the formation of the peripheral zone by limiting the initial ball-like growth inside the agar, rendering it incapable of reaching the agar surface. Typical colonies composed of both central and peripheral zones appear also on other fibrillar media such as the cellulose pellicle of Acetobacter xylinum and gelatin.

INTRODUCTION

Mycoplasma organisms (PPLO) and the L-forms of bacteria share a typical colony morphology, the 'fried-egg' colony. This colony is made up of a central zone, embedded in the agar and a peripheral zone on the agar surface (Klieneberger-Nobel, 1960; Dienes, 1960). The morphology of the colony, unique for these organisms, serves as one of the most important criteria in distinguishing them from other bacteria. Although much work has been done on the morphology of Mycoplasma and L-forms, the information concerning the mode of formation of the colonies is scanty (Klieneberger, 1984; Dienes, 1945). Our aim was, therefore, to study the development of these peculiar colonies and the mechanisms involved in their formation.

Two features shared by the Mycoplasma and L-forms seem to be responsible for the formation of the 'fried-egg' colonies: the very small dimensions (100–800 μm) of the minimal reproductive units (Klieneberger-Nobel, 1956) and their plasticity due to the absence of a rigid cell wall (Klieneberger-Nobel, 1960).
METHODS

Organisms. Mycoplasma laidlawii strain A (PG8), M. laidlawii strain B (PG9), M. bovigenitalium (PG11), M. gallisepticum (PG81), M. mycoides var. mycoides (PG1), M. neurolyticum (PG 28) and the stable L-form of Proteus were kindly given by Dr D. G. ff. Edward (The Wellcome Research Laboratories, Beckenham, Kent). M. mycoides var. capri and the stable L-form of Streptobacillus moniliformis were the gift of Dr E. Klieneberger-Nobel (The Lister Institute for Preventive Medicine, London). M. hominis was isolated in our laboratory from a case of vaginitis.

Media. The medium used for stock cultures and most experiments was a modified Edward medium (Butler & Knight, 1960a) where pooled inactivated human serum replaced horse serum. Beef heart for infusion, peptone and yeast extract used for compounding Edward medium were products of Difco Laboratories Inc. (Detroit 1, Michigan, U.S.A.). Following the suggestion of Klieneberger-Nobel (1956) the medium was enriched by addition of a nutrient broth culture filtrate of Staphylococcus aureus in a final concentration of 12.5% (v/v). Edward medium was solidified with 1.25% (w/v) Bacto-agar (Difco, certified) except for some experiments where 12% (w/v) Bacto-gelatin (Difco) was used.

Coagulated serum medium (Löffler's medium) was prepared by mixing three parts of bovine serum with one part of Difco nutrient broth containing 1% (w/v) glucose. The medium was poured into Petri dishes and sterilized by inspissation at 85° for 2 hr.

Cellulose medium. Acetobacter xylinum was grown on the medium of Hestrin & Schramm (1954) for 2 days at 80°. The thick pellicle formed was removed, washed thoroughly with sterile distilled water and immersed in sterile distilled water at 6° for 8 days. The water was changed daily. The white cellulose pellicle was then soaked in liquid Edward medium, transferred to a filter paper disk to remove excess liquid and finally placed on the surface of solid Edward medium.

Conditions of growth. Test tubes (6 x ½ in.) containing 5 ml. of liquid Edward medium were inoculated with 0.1 ml. of the stock cultures and incubated statically in air at 37°. Optimal growth of all strains occurred usually after 48 hr. incubation. Decimal dilutions of these broth cultures were made in a solution of 1% (v/v) Edward medium in 0.01M-K₂HPO₄ buffer (pH 8.0; Butler & Knight, 1960b). Standard drops (0.02 ml. each) of these dilutions were placed on solid medium plates which had previously been dried by leaving them open in the 37° incubator for 1-1.5 hr. The dilutions used for inoculation were chosen so as to give no more than 500 colonies per drop area. The plates were placed in closed tins containing moist cotton wool and incubated at 37° for 8 days.

Measurement of colony and central zone diameters. The growth which appeared on plates was inspected by means of an ordinary light microscope, using slightly oblique transmitted light. Measurement of colony and central zone diameters was carried out with a ×10 objective and a ×5 ocular fitted with a micrometer disk. The drop area was divided by a line through its centre cut by means of a razor blade, and 50 colonies were measured along this line, assuring the representation of colonies growing in the periphery and centre of the drop area.

Vertical sections of colonies. Large colonies were selected for this purpose. The section was cut through the centre of the colony by means of a razor blade. For good
The form of PPLO colonies

results the thickness of the section had to be not greater than 0.2-0.8 mm. The sections were transferred to slides and examined by the light microscope.

Technique for demonstrating penetration of micro-organisms into the agar. Standard drops of diluted suspensions of the organisms tested were placed on the surface of dried Edward medium plates. Immediately after the drops had dried and at various time intervals later, small agar squares (larger than a drop area) were cut from the Edward medium and placed on the drop areas, care being taken not to form air bubbles. The agar plates and slices were then incubated at 37° for 3 days in a moist atmosphere. The number of colonies appearing in the drop areas on the plates and on the corresponding agar slices was determined by using the light microscope (magnification x 25).

RESULTS

The development of Mycoplasma and bacterial L-form colonies

Edward medium plates were inoculated with suspensions of Mycoplasma laidlawii strain A and the L-form of Streptobacillus moniliformis. Visible colonies appeared after 24-hr. incubation at 37° (Pl. 1, figs. 1, 4). At this phase of growth almost all colonies in both organisms appeared to be small spheres embedded in agar. This was proved by the circular shape of vertical sections through the colonies and by the inability to remove the colonies on scraping the agar surface with a razor blade.

Further incubation for another 24 hr. caused enlargement of Mycoplasma laidlawii strain A colonies, some of which had already developed the peripheral zone (Pl. 1, fig. 2). The L-form grew faster than the Mycoplasma and almost all colonies developed into the 'fried egg' type after 48 hr. incubation (Pl. 1, fig. 5). The peripheral zone of the colonies, located on the agar surface, was easily removed by scraping. Mature colonies of both organisms may be seen in Pl. 1, figs. 3, 6. These colonies had big peripheral zones.

The development of colonies was also followed quantitatively by measuring the diameter of the colonies and their central zones (Fig. 1). The central zone was the first to develop, followed by the formation of the peripheral zone later on. The central zone grew only a little after 48-hr. incubation, whereas the peripheral zone continued to grow, until it occupied the greater part of the colony after incubation for 96 hr.

Vertical sections of mature colonies

All Mycoplasma and L-form strains used in this investigation were examined. The colony sections showed the same basic structure in all organisms: a hemispherical central zone which is embedded in the agar and a thin peripheral zone which spreads over the agar surface (Pl. 2, figs. 7-10). The central zone was a firm and elastic structure in comparison with the very soft peripheral zone.

Penetration of Mycoplasma and L-forms into the agar and formation of the central zone

In order to understand the mechanism of formation of Mycoplasma and L-form colonies we thought it worth while to investigate why the growth of these organisms begins in the agar. Experiments to this purpose showed that the organisms penetrated into the agar medium before multiplication. As may be seen from
Table 1, immediately after the drops had dried about 50% of the Mycoplasma organisms could be transferred to the agar slices. Further incubation gradually decreased the number of organisms that could be transferred and at 6–8 hr. after inoculation almost all organisms remained on the agar plate.

The sum of the number of colonies which developed on the drop area on the plate and the corresponding agar slice remained constant within the first 8 hr. of experiment, indicating that no mortality or multiplication took place during this period. After 24-hr. incubation the number of colonies which appeared on the agar slices increased and became equal to the number of colonies on the corresponding drop areas. However, in most experiments the growth was confluent. This was due to the smear of the surface peripheral zones which began to appear at this time (Pl. 1, figs. 2, 4).

The four Mycoplasma strains tested behaved similarly; almost all organisms penetrated into the agar within 6–8 hr. after inoculation. The L-forms of *Proteus* sp. and *Streptobacillus moniliformis* differed in this respect since the number of organisms which could be transferred to the agar slice did not change significantly during the first 8 hr. of the experiment.

For comparison the same experiment was repeated with *Staphylococcus aureus* and *Streptococcus pyogenes*. More than 50% of the organisms inoculated could be transferred to the agar slices after 8-hr. incubation, indicating that these organisms did not penetrate into the agar.

Mycoplasma penetration into the agar did not occur when the surface of the agar was moist. Moist Edward agar plates were prepared by pouring different amounts of sterile distilled water on the agar surface. With *Mycoplasma laidlawii* strain A as
Table 1. *The penetration of Mycoplasma and L-forms of bacteria into agar*

All experiments were done in duplicate and some in triplicate. The table gives typical results of some of the experiments. For technical details see text.

- **a** = number of colonies found on the drop area on the plate.
- **b** = number of colonies found on the corresponding agar slice.

<table>
<thead>
<tr>
<th>Hr. after inoculation</th>
<th>Mycoplasma laidlawii strain A</th>
<th>Mycoplasma laidlawii strain B</th>
<th>Mycoplasma mycoides var. capri</th>
<th>Mycoplasma homogenitilum</th>
<th>L-form of Proteus</th>
<th>L-form of Streptobacillus moniliformis</th>
<th>Staphylococcus aureus*</th>
<th>Streptococcus pyogenes*</th>
</tr>
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<tr>
<td>0</td>
<td>110 80</td>
<td>55 44</td>
<td>187 160</td>
<td>55 45</td>
<td>152 215</td>
<td>88 78</td>
<td>6 16</td>
<td>29 68</td>
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<td>45 48</td>
<td>166 122</td>
<td>68 28</td>
<td>201 152</td>
<td>70 71</td>
<td>11 16</td>
<td>25 68</td>
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<td>244 164</td>
<td>135 165</td>
<td>10 14</td>
<td>120 280</td>
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<tr>
<td>6</td>
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<td>108 12</td>
<td>250 45</td>
<td>58 25</td>
<td>216 162</td>
<td>129 128</td>
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<td>200 80</td>
<td>90 8</td>
<td>270 18</td>
<td>70 35</td>
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* in this experiment thallium acetate and penicillin were omitted from the Edward medium.
** in confluent growth.
inoculum, growth was inspected after incubation at $37^\circ$ for 3 days in a moist atmosphere. The colonies which appeared on the surface of the moist plates were large, flat, and had either minute central zones or none at all (Pl. 2, figs. 11, 12).

Suspensions of Mycoplasma or L-forms were mixed with melted Edward agar at 45° and poured into plates. Most colonies developed deep inside the agar, forming ball-like structures, in contrast to the lens-like deep agar colonies of Eubacteria such as *Staphylococcus aureus* (Pl. 3, figs. 18–15).

**Factors influencing the form and dimensions of the peripheral zone**

It was assumed that the peripheral zone of Mycoplasma and L-form colonies develop in the thin free water film which is present on the agar surface (Knaysi, 1951). This view was supported by the demonstration of the spreading of peripheral zones along cotton-wool fibres placed on the agar (Pl. 3, figs. 16, 17).

![Graph](image)

**Fig. 2.** The effect of pre-drying the Edward medium on the morphology of colonies of *Mycoplasma laidlawii* strain A and the L-form of *Streptobacillus moniliformis*.

The need of a humid atmosphere for optimal growth of Mycoplasma and L-forms has been stressed by many authors (Edward, 1954; Klieneberger-Nobel, 1954). Drying of the plates and subsequent incubation in a dry atmosphere decreases the free water film on the agar surface, possibly affecting the formation of the peripheral zones. To test this assumption, solid Edward medium in Petri dishes was dried for different periods of time by opening the plates in a $37^\circ$ incubator. After the inoculation with suspensions of *Mycoplasma laidlawii* strain A and the L-form of *Streptobacillus moniliformis* the plates were placed in tins without moist cotton wool and incubated at $37^\circ$ for 8 days. The dimensions of the peripheral zones were found to be markedly decreased in the case of the dried plates (Fig. 2). Drying for 5 hr. caused complete disappearance of the peripheral zones in all colonies of *M. laidlawii* strain A
The form of PPLO colonies

and in almost all colonies of the L-form of S. moniliformis. The rate of reduction of the central zone diameter due to drying of the medium was much smaller than that of the peripheral zone diameter.

Fig. 3. The effect of agar concentration in Edward medium on the morphology of Mycoplasma laidlawii strain A colonies.

Fig. 4. The effect of agar concentration in Edward medium on the morphology of colonies of the L-form of Streptobacillus moniliformis.

The effect of agar concentration on the morphology of the colonies

The effect of changes in the agar concentration in Edward medium on the morphology of colonies of Mycoplasma laidlawii strain A and the L-form of Streptobacillus moniliformis is illustrated in Figs. 3 and 4. With low agar concentrations (0.1-0.8 %, w/v), the medium was semisolid and M. laidlawii strain A grew in small irregular ball-like colonies at various depths of the medium. A hazy surface growth was also present. Typical colonies, consisting of both central and peripheral zones, appeared at a concentration of 0.5 % (w/v). Increasing the concentration of the agar to 1.25 % (w/v) increased the diameter of the colonies, mainly that of the peripheral zones. Further increase in agar concentration caused a gradual decrease of the colony diameter, affecting primarily the peripheral zone. At a concentration of 5 % (w/v) agar, almost all colonies were without peripheral zones. Similar results were obtained with the L-form of S. moniliformis (Fig. 4). The diameter of the central zone of the L-form colonies did not change significantly by changing the agar concentration of the medium. The optimal agar concentration for the development of the peripheral zone was about 1.5 % (w/v); higher concentrations inhibited its formation. A hazy surface growth was observed with low agar concentrations (0.1-0.2 %, w/v).

Morphology of Mycoplasma colonies on solid media other than agar

Mycoplasma laidlawii strains A and B and M. mycoides var. capri were grown on Edward medium solidified with 12 % (w/v) gelatin instead of agar. The plates were incubated for 8 days at 25°. Growth appeared mostly in the shape of ball-like colonies at various depths of the medium. Some colonies near the surface also
developed peripheral zones. Liquefaction of the gelatin by the organisms after further incubation masked this typical colony morphology.

*Mycoplasma laidlawii* strain A gave rise to colonies composed of central and peripheral zones when grown on the cellulose pellicle of *Acetobacter xylinum*. Many colonies, however, had no peripheral zones since they had developed deep inside the pellicle, resembling the deep agar colonies of *Mycoplasma* (Pl. 8, fig. 18). *Mycoplasma laidlawii* strain A gave flat surface colonies without central zones when grown on coagulated serum medium. The same type of surface colonies, not penetrating into the medium, appeared when this organism was grown on the surface of a cellophan dialysis membrane placed on solid Edward medium.

![Graph of pH value vs. colony growth](image)

*Fig. 5.* The effect of pH value on the growth of *Mycoplasma laidlawii* strain A and the L-form of *Streptobacillus moniliformis* in Edward medium.

**The effect of inorganic salts and hydrogen ion concentration on the morphology of *Mycoplasma* and L-form colonies**

Sodium chloride or Li$_2$SO$_4$.H$_2$O were incorporated into solid Edward medium in concentrations of 0.5, 1.0, 2.0 and 3.0% (w/v). The plates were inoculated with *Mycoplasma laidlawii* strain A and the L-form of *Streptobacillus moniliformis* and the growth inspected after 8 days of incubation at 37°C. Normal growth of *M. laidlawii* strain A occurred at concentrations 0.5 and 1.0% (w/v) with both salts. At a salt concentration of 2.0% (w/v) a partial inhibition of growth was noted; the colonies were much smaller and many lacking in peripheral zones. With 3.0% (w/v) of Li$_2$SO$_4$.H$_2$O none of the colonies had peripheral zones, while NaCl at a similar concentration inhibited growth completely. The growth of the L-form of *S. moniliformis* was completely inhibited with 2.0% and 3.0% concentration of both salts. At a salt concentration of 1% (w/v) most colonies appeared with central zones only, while 0.5% (w/v) of NaCl or Li$_2$SO$_4$.H$_2$O gave normal colonies.
The pH value of the medium has a marked influence on the growth of Mycoplasma organisms (Edward, 1954; Peoples, Morton & Feo, 1957). The growth and colony morphology of *Mycoplasma laidlawii* strain A and the L-form of *Streptobacillus moniliformis* were tested at various pH values of Edward medium (Fig. 5). *M. laidlawii* strain A did not grow at pH values below 6.85; optimal growth occurred at pH 7.8. Increasing the pH above 8.0 decreased growth, causing most colonies to appear without peripheral zones. At pH 8.55 peripheral zones were absent from all colonies. The L-form of *S. moniliformis* was less sensitive to pH changes than the Mycoplasma, but inhibition of growth also occurred at low or high pH values, accompanied by the appearance of small colonies without peripheral zones.

DISCUSSION

On the basis of our findings and those already reported by others, we are able to propose a possible mechanism for the formation of Mycoplasma and L-form colonies (shown schematically in Fig. 6). Under normal conditions the multiplication of these organisms begins inside the agar. The inoculated organisms are drawn into the agar gel together with the water surrounding them. The dried agar surface has strong imbibition properties, and water is absorbed quickly by capillary forces (Porter, 1946). This can be demonstrated by placing a drop of water on the surface of a dry agar plate; the drop is absorbed within a few minutes, resulting in a slight swelling of the agar at the drop area. When the agar surface is moist the absorption of additional water is decreased or even abolished. Therefore penetration of the organisms is very slight, or absent altogether, resulting in the formation of surface

![Diagram](https://example.com/diagram.png)
colonies with small or no central zones (Pl. 2, figs. 11, 18). Hence, the forces responsible for drawing the micro-organisms into the agar are the capillary forces of the agar gel. The determinant factors of the actual penetration, however, are the very small dimensions and the plastic nature of the Mycoplasma and L-forms, allowing these organisms to enter between the agar fibrils and to move in the interstices of the fibrillar network of the agar gel.

Klieneberger-Nobel (1956) showed by filtration experiments and electron microscopy that L-form cultures contain viable particles of various sizes, the smallest of which have a diameter of about 800 μm and occur only in small numbers. In contrast, cultures of Mycoplasma are composed mainly of very small viable particles, which have an average diameter of about 100–150 μm. These differences between L-forms and Mycoplasma might explain the results shown in Table 1. Most viable particles of Mycoplasma penetrated into the agar, whereas only a part of the viable particles of the L-forms showed this ability.

The absence of motility in Mycoplasma and L-forms (Klieneberger-Nobel, 1954, 1960) excludes the possibility of active penetration of the organisms into the agar. Gravity forces do not appear to play any role in the penetration since this process is not affected by incubating the inoculated plates upside down. Actinomycetes are known to penetrate into the agar by their filamentous growth (Knaysi, 1951). Some Mycoplasma species also produce filaments and Freundt (1960) has expressed the opinion that this is a common feature of all Mycoplasma species. The possibility that the Mycoplasma penetrate into the agar through the formation of filaments is, however, disproved by the fact that penetration is accomplished within a few hours after inoculation, before the occurrence of any growth and multiplication (Table 1).

It is further shown by our results that the multiplication of the Mycoplasma and L-forms begins inside the agar at various distances from the surface, depending on the initial depth of penetration of the viable particle giving rise to the colony. During the process of multiplication the organisms do not push aside the agar, because they can penetrate and occupy the interstices between the agar fibrils. This is demonstrated by the finding that the growth of Mycoplasma and L-forms in the agar results in the formation of firm ball-like colonies instead of the lens-shaped colonies formed by the usual Eubacteria inside the agar (Pl. 3, figs. 13–15). The formation of lens-shaped deep agar colonies is explained by the inability of the usual bacteria to penetrate into the interfibrillar spaces, therefore causing the agar to split during the multiplication; the colony then spreads along this split (Knaysi, 1951). Staphylococci and streptococci do not penetrate into the agar, as shown in Table 1. It may be concluded that the central zone with Mycoplasma is a ball composed of a network of agar fibrils, with the soft micro-organisms occupying the interstices. The agar fibrils are responsible for the firmness of the central zone.

The ball-like growth inside the agar continues to grow and expand in all directions until it reaches the agar surface which is usually covered by a thin film of free water (Knaysi, 1951). At this stage the organisms spill into this free water film, forming the peripheral zone. The surface tension of the water is responsible for the circular shape of the peripheral zone. The latter is very soft because it does not contain agar, in contrast to the firm central zone. Cotton-wool fibres on the agar cause the peripheral zones of adjacent colonies to lose their round form and to spread along the fibres due to changes in the surface tension of the water surrounding the
The form of PPLO colonies

fibres. This is further evidence for the location of the peripheral zone in the free water film of the agar.

The dimensions of the peripheral zone are determined by the thickness of the free water film. The thinner the film, the smaller will be the peripheral zone, and the colony as a whole. This explains, at least partly, the importance of a fresh medium and a humid atmosphere for optimal growth of Mycoplasma (Edward, 1954).

It is interesting to note that the large bodies, typical of L-forms and Mycoplasma, appear only at the peripheral zone of the colony (Liebermeister, 1960). This phenomenon might be explained by remembering that inside the agar the organisms are pressed together by the agar fibrils, whereas in the peripheral zone they are free to expand either by growth of the cytoplasm, or by swelling due to low osmotic pressure of the free water film.

The necessity of a fibrillar structure of the medium for the formation of the central zone was demonstrated with the cellulose pellicle of Acetobacter xylinum, known to be composed of a network of cellulose fibrils (Mühlethaler, 1949). The typical morphology of Mycoplasma and L-form colonies on gelatin medium apparently depends also on the fibrillar nature of this gel. Hayflick & Stinebring (1960) reported the formation of typical Mycoplasma colonies on the fibrillar plasma clot. On the other hand coagulated serum, which is not fibrillar, does not allow penetration of the organisms and the typical 'fried egg' colony does not appear with this medium (Dienes, 1960).

Apparently the wider interstices between the cellulose and gelatin fibrils, compared to agar at a concentration of 1.25 % (w/v), enable the organisms to penetrate deeper into the cellulose and gelatin media. Thus, the inside growth frequently cannot reach the surface and grows into a ball-like colony inside the medium. The same may happen when the concentration of the agar in Edward medium is decreased. The wider interstices between the fibrils at low agar concentrations explain also the smallness and the irregular boundaries of the central zones observed at these concentrations. The larger viable particles of the L-forms (Klieneberger-Nobel, 1956) apparently do not penetrate quite so deeply into the agar (Table 1), thus explaining the development of large central zones and the appearance of peripheral zones at lower agar concentrations than with the Mycoplasma. The decrease of the peripheral zone dimensions observed at high agar concentrations might be explained by the diminution of the free water film under these conditions (Knaysi, 1951). Increasing the agar concentration also causes decrease of the central zone diameter, apparently due to the inhibition of penetration of the organisms into the medium.

It is noteworthy that when the medium becomes unfavourable for growth, as in the presence of high concentrations of inorganic salts or unsuitable hydrogen ion concentrations, the peripheral zone is the first to disappear, presumably because the initial ball-like growth inside the agar is not big enough to reach the surface and form the peripheral zone. This observation might explain the formation of the T-colonies of Mycoplasma (Shepard, 1960), which are composed apparently only of central zones embedded in the agar (Hayflick & Stinebring, 1960). This is due to the unsuitability of the medium to support optimal growth of these exacting Mycoplasma strains. This was also our experience while testing defined media for Mycoplasma. Poor media only gave rise to this type of agar-embedded colonies without peripheral zones.
Our thanks are due to Dr M. Benzman for supplying us with the cellulose pellicle of *Acetobacter xylinum*.

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REFERENCES


EXPLANATION OF PLATES

**PLATE 1**

Figs. 1–8. The development of *Mycoplasma laidlawii* strain A colonies on Edward medium after 24, 48 and 96 hr. incubation at 37°C. × 50.

Figs. 4–8. The development of the L-form of *Streptobacillus moniliformis* colonies. Medium and incubation periods as above. × 50.
S. RAZIN AND O. OLIVER

(Pacing p. 230)
S. RAZIN AND O. OLIVER
The form of PPLO colonies

PLATE 2

Figs. 7–10. Vertical sections of colonies.

Fig. 7. Mycoplasma laidlawii strain A. ×150.

Fig. 8. Mycoplasma mycoides var. capri. ×150

Fig. 9. Mycoplasma gallisepticum. ×600.

Fig. 10. L-form of Proteus. ×150.

Figs. 11, 12. Colonies of Mycoplasma laidlawii strain A grown on moist Edward agar. × 50.

PLATE 3

Fig. 13. Deep Edward agar colonies of Mycoplasma laidlawii strain A. Including one surface colony with a peripheral zone. × 50.

Fig. 14. Deep Edward agar colonies of the L-form of Streptobacillus moniliformis. One small surface colony is seen at the right end of the photograph. ×50.

Fig. 15. Deep Edward agar colonies of Staphylococcus aureus. ×50.

Fig. 16. Mycoplasma laidlawii strain A colonies touching cotton-wool fibres on the surface of Edward medium. ×50.

Fig. 17. L-form of Streptobacillus moniliformis colonies touching cotton-wool fibres on the surface of Edward medium. ×50.