Electron microscopic evidence of flagella and pili on *Legionella pneumophila*

**FG RODGERS, PW GREAVES, AD MACRAE, AND MJ LEWIS**

*From the Public Health Laboratory, University Hospital, Queen's Medical Centre, Nottingham NG7 2UH, UK*

**SUMMARY** Twenty-one strains of *Legionella pneumophila*, representing the six known serotypes of the organism, cultured on various bacteriological media and in the yolk sacs of fertile hens' eggs were examined by negative stain electron microscopy for flagella and pili. These appendages were usually observed after cultivation on media capable of inducing an early profuse growth of the organisms.

*Legionella pneumophila*, a recently reported bacterial species,1 is the causative organism of an acute form of lobar pneumonia known as Legionnaires' disease, so called after an outbreak that followed an American Legion Convention in Philadelphia in 1976.2 From postmortem lung tissue or culture on bacteriological media the organism appeared characteristically to be a small bacillus with properties resembling those of other Gram-negative bacteria.3-8 Its size is approximately 0.3 to 0.5 μm by 1 to 3 μm, but longer filamentous forms are common. Organisms can be specifically identified from postmortem tissues, infected yolk sacs of fertile hens' eggs, and bacteriological media by immunofluorescent antibody staining8 and immunoferritin electron microscopy.7 By gas-liquid chromatography the bacteria show a characteristic fatty acid profile.8 Strains of legionellae have also been recovered from environmental sources.9 10

Early studies of the organism after growth on less satisfactory bacteriological media such as Mueller-Hinton agar did not reveal the presence of flagella or pili (fimbriae)11 or indicate motility. On new media,12 13 which give a more rapid and profuse growth of the organisms, some strains were shown by optical microscopy14 to possess flagella. Similarly, flagella and fimbriae or pili were evident by electron microscopy.15 This report confirms and amplifies the finding of such appendages on strains of legionellae cultured on various bacteriological media and in egg yolk sacs.

**Material and methods**

**Organisms and culture media**

Twenty-one strains of *L. pneumophila* were examined. These consisted of (a) serogroup 1: Philadelphia 1 and 2, Pontiac 1, Cambridge 1, Washington 1, Nottingham N/P/1, N/M/2, N3, N/P/4, N5, N6, N7, N8, N9, Bloomington 1, and Corby 1; (b) serogroup 2: Togus 1; (c) serogroup 3: Bloomington 2; (d) serogroup 4: Los Angeles 1; (e) serogroup 5: Cambridge 2; and (f) serogroup 6: Oxford 1. The strains from environmental sources were Bloomington 1 and 2 and Corby 1, the others having been recovered from clinical material. In addition, the Oxford 1 strain was isolated from both environmental and clinical specimens.10

Strains were cultured on Mueller-Hinton agar supplemented with 1% haemoglobin and 2% Iso Vitalex, CYE agar,18 enriched blood agar,19 and enriched broth medium containing: (Difco) proteose peptone 1.5% (w/v); yeast extract 0.5% (w/c); panmede liver extract 0.25% (w/v); sodium chloride 0.5% (w/v); l-cysteine hydrochloride 0.04% (w/v); and soluble ferric pyrophosphate 0.0125% (w/v). In addition, a modified CYE agar containing 0.3 g (% w/v) Lab Lemco beef extract was used. Cultures were incubated at 36°C in a humid atmosphere containing 5% CO₂ until, on solid media, individual colonies appeared or, in broth, a visible opacity occurred. The duration of incubation varied with the culture medium, being longest with Mueller-Hinton and CYE agar (five to seven days) and shortest with enriched blood agar (two to three days). Cultures on

Received for publication 8 May 1980
Electron microscopic evidence of flagella and pili on Legionella pneumophila

Fig. 1 Flagella on Legionella pneumophila, negatively stained with 1% phosphotungstic acid, pH 6.5. (a) Part of Cambridge 2 organism from enriched blood agar medium. Note insertion of single flagellum in the polar region of the organism. (x 66 000). (b) Cambridge 1 organism from enriched blood agar medium showing single subpolar flagellum. (x 20 000). (c) Partially plasmolysed organism of Nottingham N/1 strain from enriched broth medium. Note two polar flagella, one curved (broken), one straight. (x 14 400). (d) Dividing organism of Nottingham N/8 strain from enriched broth medium showing single undulating flagellum. (x 14 400). (e) Flagellum with attachment hook (arrow) from Togus 1 strain grown on modified CYE agar. (x 80 000). (f) Filamentous organism of Cambridge 1 strain with single polar, slightly curved, flagellum. From enriched blood agar. (This flagellate organism was 49 μm in length.) (x 22 500).
the modified CYE agar and enriched broth medium were examined after three to five days' incubation. Certain strains were also inoculated into the yolk sacs of 6- to 7-day-old fertile hens' eggs, which were reincubated until death of the embryo was observed, usually after four to six days. The yolk sac membranes were then harvested.

**Electron Microscopy**

Samples taken from discrete colonies and as pieces of infected yolk sac membrane were placed in 0.1 ml 1% (v/v) formal saline; 0.1 ml volumes of broth were removed by pipette and added to 0.1 ml 2% (v/v) formal saline. All samples were left undisturbed for 1 hour and were thenPrepare preparations to disperse the organisms. Drops of approximately 25 µl were removed from all preparations by pipette, placed on 400 mesh formvar-carbon-coated electron microscope grids, and allowed to stand for 30 seconds before the excess was drained with filter paper. Grids were negatively stained by the addition, for a few seconds, of 1 drop 1% (v/v) phosphotungstic acid, pH 6.5. After they had been blotted dry, the prepared grids were examined in an AEI Corinth 500 or JEOL 100C electron microscope; a magnification of ×5000 was used to identify the organisms, and ×25000 to search for flagella and pili.

**Results**

Twenty-one strains of Legionella from six serogroups were examined by negative stain electron microscopy after cultivation, as indicated. Flagella were observed on all strains grown in one or more of the media. Although they were not found on every occasion, the incidence of flagellation was higher in those media giving early profuse growth (Table 1). The

<table>
<thead>
<tr>
<th>Culture medium</th>
<th>Incubation period (days)</th>
<th>Serogroup</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1 2 3 4 5 6</td>
</tr>
<tr>
<td>Mueller-Hinton agar</td>
<td>5-7</td>
<td>3(10) 1 1 0 NT 0</td>
</tr>
<tr>
<td>CYE agar</td>
<td>5-7</td>
<td>4(10) 0 0 0 NT 1</td>
</tr>
<tr>
<td>CYE agar + beef extract</td>
<td>3-5</td>
<td>8(10) 1 1 1 NT 1</td>
</tr>
<tr>
<td>Enriched blood agar</td>
<td>2-3</td>
<td>14(16) 1 1 1 1</td>
</tr>
<tr>
<td>Enriched broth medium</td>
<td>3-5</td>
<td>13(16) 0 1 1 0 1</td>
</tr>
<tr>
<td>Yolk sac membranes</td>
<td>4-6</td>
<td>2 (3) 1 1 1 NT 0</td>
</tr>
</tbody>
</table>

Numbers in parentheses indicate the number of strains tested; serogroups 2-6 one strain of each examined. NT = not tested.

Nottingham N/M/2 and Washington 1 strains were deficient in flagella from the enriched blood agar but positive from enriched broth medium. The reverse effect was observed for the Philadelphia 1, Notting-
In Gram-negative bacteria, the flagellum is linked to the inner and outer membranes of the bacterial cell wall by means of the basal granule comprising a hook-like structure and four protein rings, one of which is incorporated into the peptidoglycan layer of the cell wall. That peptidoglycan is present in legionellae, though in a barely discernible form, has been questioned. The presence of flagella with an attachment mechanism similar to that for other Gram-negative organisms adds further evidence to the existence of this cell wall component.

Though the epidemiology of sporadic cases of Legionnaires’ disease is not clear, there is evidence that the organism is present in the environment, particularly in water. It has been found in cooling tower water and streams in the USA and in bathroom showers in the UK. If the organism exists primarily in water, and its pathogenicity for man is coincidental, the production of flagella under suitable cultural conditions is not unexpected. Failure to observe flagella or pili in organisms from clinical material may reflect the advanced phase of bacterial replication, for example within the lung, a modification to the organism cell wall, or conditions in vivo adverse to flagella production. That pili were observed only on strains grown on enriched blood agar and in broth medium is not surprising, since this type of appendage, thought to be an integral cell wall component, is found on many species in the logarithmic phase of growth and disappears thereafter. Although flagella and pili were not found in clinical specimens, it is possible that these antigens may play a part in stimulating antibodies in the early phase of disease during exponential growth of the bacteria in vivo.

The presence of such appendages on legionellae is very significant in the determination of the taxonomic relation of the new group to other bacterial species. Cultural and structural studies are in progress to characterise these remarkable microorganisms further.

We thank Mrs Joan Casson for help in preparing the electron micrographs.

References


4. Chandler FW, Cole RM, Hicklin MD, Blackmon JA,


12 Feeley JC, Gorman GW, Gibson RJ. Primary isolation media and methods. In: Jones GL, Hebert GA, eds. Legionnaires' the Disease, the Bacterium and Methodology. Atlanta, Georgia: Centre for Disease Control, 1978:107-17.


Requests for reprints to: Dr FG Rodgers, Public Health Laboratory, University Hospital, Queen's Medical Centre, Nottingham NG7 2UH.
Electron microscopic evidence of flagella and pili on Legionella pneumophila.

F G Rodgers, P W Greaves, A D Macrae and M J Lewis

doi: 10.1136/jcp.33.12.1184

Updated information and services can be found at:
http://jcp.bmj.com/content/33/12/1184

These include:

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

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/