Preliminary report on the pathogenicity of *Legionella pneumophila* for freshwater and soil amoebae

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**SUMMARY** *Legionella pneumophila*, the causative organism of Legionnaires’ disease, is pathogenic for free living, ubiquitous, freshwater, and soil amoebae of the genera *Acanthamoeba* and *Naegleria*. Some species support the growth of strains from serogroups 1 to 6, others only strains from certain serogroups. Initial studies with seeded material indicate that amoebal enrichment could be utilised for the isolation of legionellae from clinical specimens and natural habitats. It is suggested that a vacuole, or amoeba, full of legionellae, rather than free legionellae, could be the infective particle for man.

*Legionella pneumophila*, the causative organism of Legionnaires’ disease, has been isolated from the water of cooling towers,1–3 evaporative condensers,4 ponds,5 a thermally polluted stream, the mud of that stream,1 2 and various other aquatic habitats. It will survive several months, or longer, in water for injection,6 and in distilled,7 8 tap,7 stream,9 and pond water.10 However, laboratory media for the growth of *L. pneumophila* require supplementation with L-cysteine and ferric iron, usually ferric pyrophosphate. The main carbon sources for *L. pneumophila* are L-amino acids. Arginine, cysteine, isoleucine, leucine, methionine, threonine, valine, and phenylalanine or tyrosine are required for growth. Most strains require serine, some also require proline. Serine and, to a lesser extent, threonine can be used as sole sources of carbon and energy.11 12 Carbohydrates11 13 and many organic acids are not utilised, while some organic acids, for example, citric and oleic acid, are inhibitory.11 Growth on solid media is initiated only within a narrow pH range, around 6.9 (6.3–7.28).14 15 The optimum incubation conditions are 35°C in air +2·5% (v/v) CO2; no growth occurs anaerobically. The temperature range over which growth can occur is approximately 25°C–40°C. On first isolation, growth is usually very slow; only pin-point colonies are seen after three days. Growth of *L. pneumophila* on solid media is inhibited by many bacteria, for example, certain members of the oral flora, a *Bacillus* species16 and *Pseudomonas aeruginosa*, also by a purified pyocin (unpublished data). These cultural characteristics indicate that *L. pneumophila* is not a free living aquatic bacterium. Recently, a relationship between *L. pneumophila* and a *Fischerella* sp. has been described. Growth of *L. pneumophila* occurred only when the cyanobacteria were photosynthesising.17

In man, *L. pneumophila* infects macrophages, in which large numbers of legionellae have been observed intracellularly.16–20 Extracellular legionellae appear to be closely associated with lysed macrophages.18 *In-vitro* studies with cyanolmolgus monkey macrophages21 showed that *L. pneumophila* can grow in their cytoplasm and that, although rat and mouse alveolar macrophages ingest only antibody-coated *L. pneumophila*,22 this is not a necessary prerequisite for phagocytosis by cyanolmolgus monkey macrophages.21 Intracellular *L. pneumophila* in polymorphonuclear leucocytes appears to be rare.18 23 Macrophages are amoeboid cells and, because of the similarity between macrophages and amoebae, this study was initiated to determine the interactions of *L. pneumophila* and similar organisms with amoebae. Free living amoebae of the genera *Naegleria* and *Acanthamoeba* are ubiquitous in soil and water. The commonest species, *A. polyphaga*, *A. castellanii*, and *N. gruberi*, have been isolated from humidifiers and may be present there in large numbers.24

**Methods**

**Culture**

Ninety millimetre diameter plates of Stoianovitch’s malt extract, yeast extract agar,26 prepared with Page’s amoebal saline,28 pH 6·9, were flooded with a turbid suspension of *Klebsiella aerogenes* (NCTC

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7427) in amoebal saline, drained, and incubated aerobically overnight at 35°C. The agar was divided into two semicircular sections by the removal of a diagonal 10 mm wide strip. To kill the klebsiellae, the open plates were exposed for 30 minutes to the UV light of a microbiological safety cabinet. One end of each semicircular lawn was inoculated with a suspension of amoebal trophozoites, and approximately 15-20 amoebae were drawn with a wire across the semicircle so that the advancing amoeba would have to cross it. The wire was inoculated with legionellae from an enriched blood agar26 (with soluble starch 1·5 g/l, and L-serine 1·0 g/l) plate, incubated at 35°C, in air + 2·5 % v/v CO2, for three to five days. On the control side, UV-killed klebsiellae were used. Amoebal cultures were incubated aerobically at 30°C, the sump temperature of the type of humidifiers from which L.pneumophila has been isolated.27

MICROSCOPY
The amoebae were examined microscopically in situ on the agar surface with a × 40 12 mm working distance objective (Vickers Instruments, Haxby Road, York). Isolation of single cysts and amoebae was facilitated by the long working distance of this objective and the absence of image inversion and reversion. For higher magnifications, amoebae were suspended in one drop of amoebal saline on a 22 × 64 mm cover slip. Small drops of glycerol were placed at each end, and a cavity slide was inverted over the cover slip. The mount was left in that position for at least 20 minutes to allow the amoebae to extend and adhere to the cover slip; the slide was inverted for examination. After examination of the live material the slide was removed, and 3 drops of hot (≈80°C) 10% (v/v) formalin were added to the amoebal drop just before it dried. After 2 minutes the cover slip was immersed for 2 minutes in unheated 10% (v/v) formalin, rinsed in tap water, air dried, and stained by a modification of the Giménez technique,28 in which the buffered carbol fuchsin was warmed to 50°C, filtered on to the cover slip, and left for 10 minutes. After thorough washing in tap water, the cover slip was stained with aqueous malachite green (8 g/l) for 5-10 seconds, washed in tap water, stained again with the malachite green for 5-10 seconds, washed in tap water, and finally stained with aqueous methylene blue (40 mg/l) for 60-90 seconds, washed, and dried. With this technique klebsiellae (coccobacilli) were blue to pale pink, rarely maroon, legionellae (bacilli) pink to maroon, amoebal cytoplasm pale blue, and nucleus blue-grey. The identity of the extracellular bacteria can be confirmed by immunofluorescence but not of those intracellularly, as antibody does not enter intact amoebae.

Results and discussion
The Oxford-129-31 strain of L.pneumophila isolated from a patient in an Oxford renal transplant unit29 was chosen for special study because it was a recent isolate and thus less likely to be laboratory-adapted, and because similar strains had been isolated from the sieves of the transplant unit showers (JO'H Tobin and RG Mitchell, personal communication). The slowest growing amoeba, Acanthamoeba castellanii, took about five days to reach the line of legionellae. After two days in contact with the line there was no advance of the amoebae beyond the line. The amoebae in contact with the line were usually rounded up, smaller, and vacuolate; some were lysed. Heavily infected amoebae (Fig. 1) occurred in the vicinity of the line. The infection was confined to vacuoles, often up to 10-15 vacuoles/amoeba. The legionellae in the vacuoles were non-motile. In some rare amoebae, the infection was not confined, and these amoebae developed into bags of non-motile legionellae. This latter type of infection was more common with the other
L. pneumophila strains and is similar to that seen in human\textsuperscript{18} and monkey\textsuperscript{21} macrophages. In preparations of live material, some amoebae contained numerous large vacuoles, but even heavily infected amoebae usually produced pseudopodia. On the control side, the amoebae advanced beyond the line of dead klebsiellae; no legionellae could be demonstrated in those amoebae. Small groups of legionellae and single legionella in recently infected amoebae could be demonstrated only by staining. Some amoebae were so heavily infected (Fig. 1) that the infection could only be fatal. Infected amoebae were easily ruptured. Similar results were obtained with Oxford-1 and A. polyphaga.

An Acanthamoeba sp. isolated from a local humidifier grew much faster and encysted more rapidly than the axenically maintained culture collection acanthamoebae. Growth across the entire plate occurred in three to four days and was not prevented by the line of legionellae, which were phagocytosed. Numerous faecal vesicles, 2-4 µm in diameter, were present in the area of the legionella inoculum; these did not contain recognisable legionellae. A heavy lawn of mature cysts was laid down on the inoculum line. Particularly where the inoculum was heavy, cysts were present before and after the line. On the control side, many scattered cysts were present, but few were arranged on the position of the klebsiella inoculum line. On the test side, after 48 hours malformed cysts, some empty, and grotesquely malformed amoebae were seen behind the line and between the line and the advancing edge of amoebal growth. No cysts or malformed amoebae were seen at the advancing edge of amoebal growth. In hanging drop preparations, the amoebae usually appeared normal, but some rare amoebae had a single large endoplasmic vacuole containing many rapidly motile bacilli. These infected amoebae were still moving, and apparently feeding, explaining how infected amoebae and failed cysts could be found behind the inoculum line. Large vesicles presumably from ruptured or lysed amoebae were rare, but some of these also contained rapidly moving bacilli. When stained, some of the smaller amoebae were also found to be infected. The results for other strains are given in the Table. In some rare amoebae, one to three intracellular filaments were seen; occasionally a filament would be almost coiled. Similar coiled forms of legionella have been reported in human lung.\textsuperscript{32} Mature cysts appear to be free from, and resistant to, infection. No legionellae were observed intracellularly in the flagellate phase of Naegleria. Studies on the temperature limits for infection are still in progress, but at room temperature (\textasciitilde 21°C) it is more difficult to find infected amoebae.

Amoebae of the genera Acanthamoeba and Naegleria are ubiquitous in soil and fresh water. These studies show that they are possible natural hosts for L. pneumophila. The addition of UV-killed klebsiellae to humidifier sediments at room temperature greatly increased the number of amoebal

<table>
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<th>L. pneumophila strain</th>
<th>Serogroup</th>
<th>Acanthamoeba castellanii AC L1501/2A</th>
<th>Acanthamoeba polyphaga AP L1501/3A</th>
<th>Acanthamoeba sp. humidifier strain*</th>
<th>Naegleria gruberi 1518/1E</th>
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*Clone derived from a single cyst. CPE = cytopathic effect (rounding up, vacuolation, lysis); + Slight; + + Moderate; + + + Marked; advance of amoebae often stopped by the legionella inoculum line; ♂ Grotesquely deformed amoebae; C = an increase in the number of cysts on the legionella inoculum line, particularly where the inoculum was heavy.

I = infection; + rare; + + several; + + + most amoebae; FV = large numbers of faecal vesicles on the legionella inoculum line; M = legionellae motile inside an endoplasmic vacuole of a live amoeba, rare; m = legionellae motile inside an extracellular vesicle, uncommon.

Named amoebae from the Culture Collection of Algae and Protozoa, Cambridge; NCTC = National Collection of Type Cultures, Colindale, London.
trophozoites. Rotifers, if present, selectively grazed these (Fig. 2) until only rare amoebal cysts were left; the rotifers themselves then declined. A further addition of dead klebsiellae enabled the cycle to be repeated. These initial studies show that a population of suitable feeding amoebae at a temperature above 21°C, probably in the region of 30-35°C, would be necessary for large numbers of legionellae to occur in a humidifier. Cooling towers and evaporative condensers, implicated so often in Legionnaires' disease outbreaks, 3 operate at, or near, the optimal growth temperatures for amoebae and L. pneumophila. Infected amoebae are fragile and thus likely to burst when forced through the spray nozzles of such equipment. With the continual recycling of the water, most amoebae could quickly become infected. Antifouling chemicals might inhibit rotifers, and filters used to prevent the spray nozzles from clogging may also remove them, allowing the amoebal population to increase above the levels found in natural habitats. Infected amoebae and vesicles containing legionellae would be present in the drift from contaminated equipment. The larger amoebae might be caught by the baffle plates, but smaller amoebae and vesicles would be less likely to be removed.

Fig. 2  Live rotifer feeding on amoeba (a). From this material out of a local humidifier the Acanthamoeba sp. in the Table was isolated. A trace of methylene blue was added to the preparation to stain the amoeba.

It is conceivable that infection is acquired not by inhaling free legionellae but by inhaling a vesicle or an amoeba full of legionellae. The potential host would then receive up to 50 or, in the case of an amoeba, probably up to 1000 or more legionellae at once. The surrounding vesicle membrane, or amoeba, would help to prevent rapid dehydration of the legionellae. Such a mechanism would explain the lack of patient spread in Legionnaires' disease and the occurrence of the Broad Street cases in the 1976 Philadelphia outbreak, 3 and the four cases in people who passed by in an adjacent street to a Memphis hospital during an outbreak. 3 Trials with seeded material suggest that amoebal enrichment could perhaps be utilised for the recovery of legionellae from natural habitats and clinical specimens. Certain serogroups might be selectively enriched (Table). The WIGA (ALLO-I) strain of Legionella bozemanii, 36 and the legionella-like organism Tex-KL, 37 isolated from cases of fatal human pneumonia, also infect amoebae (in preparation). Further studies, including electron microscopy, on the interactions of L. pneumophila and similar organisms with amoebae are in progress.

I am indebted to Dr LR Hill, of the NCTC, for the WIGA, Tex-KL, and NCTC L. pneumophila strains, and to Mr PG Greaves, of the Nottingham Public Health Laboratory, for the other legionella strains.

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