Adhesion, Penetration and Intracellular Replication of *Legionella pneumophila*: an *in vitro* Model of Pathogenesis

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*Legionella pneumophila* attached to, penetrated and replicated within the three eukaryotic cell lines, MRC-5, HEp-2 and Vero. Multiplication occurred rapidly in these cells for an initial 48 h after inoculation and declined thereafter. Infected MRC-5 cell monolayers developed lytic-type cytopathic changes, with organisms being readily released. HEp-2 cells showed a more chronic infection, with slowly developing granular changes in the monolayers, and slow release of intracellular bacteria. In Vero cells, organisms were released rapidly along with a more progressively developing granular cytopathic effect in the monolayers. *L. pneumophila* was unable to grow in cell-free culture fluids. Uptake and intracellular development was similar for each cell type, and was initiated by 'bacteriopexis', a process in which the organisms bound via receptors and were surrounded by cellular microvilli which eventually fused, leading to bacterial engulfment. Replication of organisms in vacuoles within the cytoplasm of infected cells was confirmed by thorium labelling. These vacuoles were lined with ribosomes and, at the early stages of intracellular development, were found in close proximity to mitochondria, cytoplasmic filaments and banded enclosures. Ruthenium red staining showed that acid mucopolysaccharide capsular material was not present on these organisms during the attachment process or intracellular phase. Organism release was by lysis of the infected cells.

**INTRODUCTION**

*Legionella pneumophila*, the most commonly occurring aetiological agent of legionellosis in man, is a facultative intracellular pathogen. Dividing forms of the organism have been observed in the cytoplasm of alveolar macrophages in post-mortem human lung tissues from cases of Legionnaires' disease (Rodgers *et al.*, 1978; Chandler *et al.*, 1979; Glavin *et al.*, 1979; Rodgers, 1979). The intracellular location of the bacteria has also been reported in laboratory animals including guinea-pigs and rats (Davis *et al.*, 1982; Winn *et al.*, 1982). Intracellular replication of *L. pneumophila in vitro* has been shown in cultured cell lines (Wong *et al.*, 1980; Daisy *et al.*, 1981), animal derived alveolar and peritoneal macrophages (Kishimoto *et al.*, 1979, 1981) and human monocyte cultures (Horwitz & Silverstein, 1980). Conversely, it has been shown that organisms were taken up by human polymorphonuclear leucocytes, but failed to replicate (Horwitz & Silverstein, 1981). Although pili (Rodgers *et al.*, 1980) and numerous other structures (Flesher *et al.*, 1979; Rodgers & Davey, 1982) have been visualized on the surfaces of *L. pneumophila*, the nature of attachment and penetration of these organisms to eukaryotic cells has remained unclear (Rodgers, 1983). Intracellular survival and subsequent replication probably depend upon a bacterial cytotoxin-induced interruption of the phagolysosomal process (Friedman *et al.*, 1982).

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**Abbreviations:** PBH, poly-β-hydroxybutyrate; SEM, scanning electron microscopy; TEM, transmission electron microscopy.

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To evaluate the interaction of *L. pneumophila* with cells in culture, the sequence of events from attachment to penetration, intracellular growth and release of organisms in mammalian cells was studied by optical and electron microscopy. The structural findings were correlated with attachment to penetration, intracellular growth and release of organisms in mammalian cells.

**METHODS**

*Bacteria and cultivation.* *L. pneumophila* serogroup 1, strain Nottingham N7, a clinical isolate from sputum, was passaged once on buffered enriched blood agar containing a low concentration of sodium chloride (LBA; Dennis et al., 1981). Cultures were grown at 36 °C for 48 h. Organisms were harvested and suspended in Hanks' balanced salt solution without antibiotics (HBSS) to give a concentration of 2 × 10^7 c.f.u. ml^-1.

*Cell cultures.* Three laboratory cell lines were used: MRC-5, a semi-continuous cell line of human embryonic lung fibroblasts, and the two transformed cell lines, Vero (African Green Monkey) and HEP-2 (human epithelial laryngeal carcinoma). Cells were grown as monolayers in tubes and on glass coverslips at 36 °C until confluent. After rinsing in HBSS, the MRC-5 and HEP-2 cultures were maintained on antibiotic-free Eagle's minimum essential medium (MEM) and the Vero cells on 199 medium. Both were supplemented with 5% (v/v) foetal calf serum, 2 mm-glutamine and 0.5 mM-NaHCO₃. Aliquots of 0.1 ml *L. pneumophila* suspension were added to the washed cells and incubated at 37 °C for 2 h to allow adsorption to occur. The cells were washed free of unattached organisms, reincubated with the appropriate fresh medium and examined daily for the development of cytopathic effects.

*Viable counts.* Attachment of *L. pneumophila* to each cell type was determined on LBA from counts made from washed cell monolayers lysed in distilled water after the 2 h adsorption period by serial dilution of lysate aliquots. Intracellular replication of the organisms in the eukaryotic cells was similarly assessed at 24 h intervals. To assay release of bacteria from cells into the suspending menstruum, extracellular fluids were harvested daily. Each cell line was then washed three times in HBSS and the fluids and washings from each cell type and time were pooled for viable counts. In addition, aliquots of supplemented cell-free MEM and 199 culture media, previously inoculated with *L. pneumophila*, were assayed daily to evaluate extracellular replication.

*Electron microscopy.* At daily intervals, infected and uninoculated control monolayers were fixed in situ for 1 h in 3% (v/v) glutaraldehyde in 0.1 M-cacodylate buffer containing 10 mM-MgSO₄, pH 7.2. For scanning electron microscopy (SEM), samples were dehydrated, critical-point dried and coated with 40 nm of gold in a Balzers sputter coater. For thin-section transmission electron microscopy (TEM), cells were pre-embedded in molten 2% (w/v) agarose, post-fixed in 1% (w/v) OsO₄, dehydrated in ethanol and embedded in Epon-Araldite. In addition, infected and control monolayers were treated with ruthenium red to stain for acid polyanions such as mucopolysaccharides (Rodgers, 1979). Further monolayers were washed in 3% (v/v) acetic acid after postfixation, and surface labelled with 0.5% (w/v) thorium colloid (Polysciences Ltd, Northampton, UK) in 3% acetic acid for 16 h at 4 °C. Unattached thorium was removed by washing with acetic acid followed by cacodylate buffer. The monolayers were then embedded for TEM. Ultra-thin sections were obtained on an LKB Ultratome III and stained with uranyl acetate and lead citrate. All samples were examined in a Jeol 100C Temscan electron microscope used at 40 kV in the SEM mode and 100 kV for TEM.

**RESULTS**

*Viable counts*

Attachment of *L. pneumophila* to each of the three cell lines occurred rapidly, but with low efficiency, and by 2 h of incubation approximately 1% of the bacteria in the inoculum were bound to the eukaryotic cell receptors. Such organisms could not be removed from the cell monolayers by repeated washings. Although the difference in the final uptake of *L. pneumophila* by each cell type was not large, uptake was greatest for HEP-2 and least for Vero cells (Fig. 1a). In Vero cells, intracellular replication progressed rapidly for an initial period of 48 h after inoculation, and slowed thereafter, with a peak at 72 h. This was followed by a decline in the number of intracellular organisms present over the remaining 72 h. The highest titres of intracellular organisms were recorded with these cells. Similar results were obtained for HEP-2 cells, but the slower phase of intracellular multiplication of organisms occurred after only 24 h. The lung fibroblasts showed a peak of intracellular replication of organisms at 48 h after inoculation and this was followed by a sharp decline in bacterial numbers. This finding was in keeping with the more rapidly lytic infection of these cells leading to the liberation of organisms into the extracellular fluids (Fig. 1a).
Attachment and replication of *L. pneumophila*

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Release of organisms from each of the three cell types into the suspending menstruum occurred progressively over the 6 d study period, but was most pronounced with the MRC-5 and Vero cells. The gradual release of the bacteria from the infected HEp-2 cells reflected the slower development of the cytopathic effect and the more chronic-type infection which occurred in these cells (Fig. 1b). *L. pneumophila* inoculated into either MEM or 199 culture medium in the absence of cells failed to replicate (Fig. 1b).

**Morphological studies: light microscopy**

Inoculation of cell cultures with *L. pneumophila* was followed by the development of a cytopathic effect in all three cell lines. Although granulation of cells and eventual lysis were common to all three cell types, differences were noted in the distribution and rate of cell destruction. The lung fibroblasts developed rapidly progressive cytolytic changes distributed throughout the entire cell monolayer within 24 h of inoculation, with cell destruction complete by 96 h. In contrast, the transformed human cells showed occasional rounded cells 48 h after bacterial inoculation and either a spreading focal or generalized granular cytopathic effect of the monolayers by 96 to 120 h. After 48 h of infection, the organisms had induced changes in the Vero cells. These appeared as clumps of granular cells surrounding circumscribed zones of cell lysis, the size of which increased with incubation time.

**Morphological studies: electron microscopy**

Within 1 h of inoculation, legionellae were observed in loose combination with the surfaces of each of the three cell types. Binding of the organisms followed this initial contact and adherence bridging between the membranes of the bacteria and the eukaryotic cells was evident (Fig. 2). Attachment of organisms was often seen associated with thickening of the eukaryotic cell membrane in apposition to the region of bacterial contact. By SEM, this process was invariably associated with areas of the cell membranes possessing numerous microvilli and occasional filopodia. The presence of these cellular projections, which were rarely seen either in control cultures or areas of inoculated cells lacking attached legionellae, was in direct response to the organisms. Indeed, the microvilli appeared to extend towards the attached bacteria (Fig. 3a), were often in direct contact with them (Fig. 3b) and apparently initiated the process of cellular uptake into phagocytic-like vacuoles (Fig. 3c). Similar observations were made in thin sections, with engulfment of organisms mediated by microvilli by a process of ‘bacteriopexis’ which resulted in intracellular organisms (Fig. 3d-g).
In the early stages of intracellular development, single organisms were observed in association with dense clusters of cytoplasmic filaments (Fig. 4a), often associated with vesiculation of the cytoplasm (Fig. 4b) and followed by dividing organisms within close fitting cytoplasmic vacuoles which were often lined with ribosomes (Fig. 4c). At this stage of uptake, organisms in vacuoles were frequently seen surrounded by cross-banded strands of cell cytoplasm (Fig. 4d). Thorium labelling demonstrated that organisms occurred within the cytoplasm of infected cells, and not in invaginations of the cytoplasmic membrane (Fig. 5a). The appearance of morphologically dividing organisms coincided with a decline in the number of organisms containing poly-β-hydroxybutyrate (PBH) granules together with an increased dilation of both the endoplasmic reticulum of infected cells (Fig. 5b) and the vacuoles containing organisms, often in the absence of ribosomes (Fig. 5c). The electron-lucent zones surrounding such intracellular organisms lacked acid polyanions as shown by ruthenium red labelling (Fig. 5d).

In the late stages of infection, distended vacuoles containing many organisms rich in PBH granules occupied most of the cell cytoplasm (Fig. 6a), and this, together with a paucity of cytoplasmic organelles in the infected cells, suggested end-stage replication of the bacteria with depletion of cell-derived nutrients. Although organisms were not found within the nuclei of infected cells, karyorrhexis and karyolysis were common as were fatty degeneration of the mitochondria and dilation of the perinuclear spaces and endoplasmic reticulum, suggestive of toxin-induced damage. Release of organisms into the extracellular fluids was by cell lysis (Fig. 6b). The sequence of development of legionellae in MRC-5 cells is outlined in Table 1.

**DISCUSSION**

*L. pneumophila* adhered to, penetrated and replicated within intracellular vacuoles in MRC-5, Vero and HEp-2 cells *in vitro*, inducing cytopathic changes. The outward spreading granular foci seen in the transformed cell lines by light microscopy, and the slow release of organisms, particularly from HEp-2 cells, were suggestive of a more chronic type of infection in these cells with some cell-to-cell spread of the bacterium. Electron microscopy confirmed the slow chronic-type infection, in which all the stages of organism uptake and development were seen simultaneously throughout the study. In contrast, the rapidly lytic generalized infection of the MRC-5 cells demonstrated their sensitivity to infection with the organism. The fastidious nature of this facultative intracellular pathogen was confirmed by its failure to replicate in the extracellular culture fluids *per se*. It does, however, replicate readily on selected artificial media (Edelstein, 1984) as well as in enriched broth (Rodgers *et al.*, 1980). In supplying essential
Attachment and replication of *L. pneumophila* in MRC-5 cells *in vitro* mediated by microvilli and filopodia. (a, b) SEM of organisms attached to cell surfaces by microvilli at 2 h post-inoculation. Note microvilli appear to enfold bacteria. (c) SEM of similar organism at 4 h post-inoculation showing engulfment of one end of the bacterium by a mass of microvilli. (d–g) TEM sequence showing stages of uptake by 'bacteriopexis'. Bars, 0.5 μm.
Fig. 4. TEM of the early stages of intracellular development of *L. pneumophila*. (a) Uptake associated with cytoplasmic filaments. (b) Organisms associated with vesiculation of the cytoplasm. (c) Replication within vacuoles lined with ribosomes (arrows). (d) Organisms enclosed within cross-banded cytoplasmic extensions (arrows) within vacuoles lined with ribosomes. (a, b, d) MRC-5 cells; (c) HEP-2 cells. Bars, 1 μm.
Attachment and replication of *L. pneumophila*

Fig. 5. (a) Thorium labelling of membranes (arrow) indicated that *L. pneumophila* multiplied within the cytoplasm of infected cells and not in invaginations of the cytoplasmic membrane. Note lack of electron-dense thorium label surrounding organisms in vacuoles. (b) Organism-induced dilation of the endoplasmic reticulum. (c) Late stage infected cells with dilated vacuoles containing numerous dividing legionellae. (d) Infected cell labelled with ruthenium red shows no acid mucopolysaccharide material surrounding the organisms either within vacuoles or in the process of release. (a, b, d) MRC-5 cells; (c) HEp-2 cells. Bars, 1 μm.
nutrients, the cellular ribosomes and mitochondria located at the periphery of early infected vacuoles may be crucial to initiating intracellular multiplication. However, these structures were not found associated with vacuoles later in infection, despite continued bacterial multiplication. The precise role of eukaryotic organelles in the intracellular development of legionellae remains unclear.
Attachment and replication of *L. pneumophila*

Table 1. Morphological features involved in the infection of MRC-5 cells by *Legionella pneumophila*

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<th>8</th>
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* Feature commonly (+), rarely (±) and not (-) observed.

Attachment of organisms to the cells was mediated by thickening of the eukaryotic cell membrane followed by the production of microvilli. That such structures occurred only in areas of bacterial attachment indicated that they appeared in direct response to organisms or organism-bound products rather than to soluble toxin. Bacterial uptake was completed by a process of 'bacteriopexis' following engulfment of organisms by microvilli in association with intracellular cytoplasmic filaments. The coiling phagosome described by Horwitz (1984) for human monocytes, primarily phagocytic cells, was not observed for the cells in this study. Both cross-banded cytoplasmic projections and the numerous vesicles enclosing organisms, which appeared during the active phase of bacterial multiplication rather than during uptake, probably reflected cellular responses to the production and release of toxins. It is also probable that it was the presence of toxins which eventually induced such cellular changes as dilation of the perinuclear spaces and endoplasmic reticulum, fatty degeneration of the mitochondria and breakdown of the nuclei. These toxin-induced changes together with physical distortion due to extensive intracytoplasmic bacterial replication led to the release of organisms by cell lysis. At this stage the reappearance of PBH granules within the legionellae indicated a depletion of intracellular nutrients.

The unresponsive nature of clinical legionellosis to treatment with many antibiotics is primarily due to the intracellular nature of the organisms *in vivo* and the inability of such antibiotics to permeate within cells. Further work is required to elucidate the virulence factors associated with *L. pneumophila* uptake and intracellular survival in eukaryotic cells. Current investigations involve characterization of the cellular and bacterial receptors involved in the attachment of legionellae and hence initiation of the infectious process. These cell culture studies serve as useful models of infection at the cellular level.

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REFERENCES


