MODELS OF INFECTION

Suckling CD1 mice as an animal model for studies of Legionella pneumophila virulence

M. CASTELLANI PASTORIS, E. PROIETTI*, C. MAURI, P. CHIANI and A. CASSONE

Laboratories of Bacteriology and Medical Mycology and *Virology, Istituto Superiore di Sanità, Viale Regina Elena, 299-00161 Rome, Italy

On the assumption that specific host defences are lower in newborn and infant animals, the susceptibility of CD1 suckling mice to Legionella pneumophila was studied with the hypothesis that this model could detect consistent differences in virulence among Legionella isolates from various clinical and environmental sources. Mice 3–14 days old were indeed markedly susceptible to intraperitoneal challenge with fresh clinical isolates, but not to serially subcultured or type collection strains of L. pneumophila. For example, intraperitoneal inoculation of $10^7$ cells of a fresh clinical isolate of L. pneumophila (strain Monza 3) caused 60% mortality of suckling mice in 1 day whereas the same number of cells of a culture-attenuated derivative (strain Monza 3p50) caused <10% mortality in >15 days. Lethal infection by the ‘virulent’ Monza 3 strain was strictly dependent on mouse age (no death was observed in mice >26 days old), required the inoculation of viable cells and was not related to endotoxin production. The ‘virulent’ L. pneumophila strain was cleared from mouse lungs less rapidly, while adhering to, and being internalised into the peritoneal exudate cells (PEC) of suckling mice to a greater extent, than the avirulent derivative, as shown by immunofluorescence and confocal microscopy. The Monza 3 strain also induced the production by PEC in vivo of 5–10 times more tumour necrosis factor-alpha (TNF-α) mRNA than the Monza 3p50 strain. Overall, suckling CD1 mice appear to provide a promising, easily handled, highly reproducible and relatively inexpensive animal model for studies of the virulence of L. pneumophila, and possibly, of the role of pro-inflammatory cytokine production in this phenomenon.

Introduction

Legionellae are ubiquitous, facultative intracellular micro-organisms that can cause severe pneumonia in man [1]. In particular, Legionella pneumophila is the aetiological agent of Legionnaire’s disease [2]. The status of the host’s immune system plays a critical role in this disease, as demonstrated by the fact that infection occurs mostly in immunocompromised patients, in patients with underlying diseases and can be induced in immunocompromised experimental animals [3–8]. Nonetheless, legionellae probably possess an array of pathogenicity factors, conferring upon the micro-organisms different degrees of virulence that can be demonstrated by different, complementary in-vivo and in-vitro assays [9–13].

A common in-vivo assay for the evaluation of bacterial virulence is the direct challenge of susceptible animals. Although various animals have been proposed for the assessment of L. pneumophila virulence [13–16], the most useful appears to be the guinea-pig, either for parenteral or aerosol infection [15]. Normally, adult mice are highly resistant to extensive legionella pneumonia and lethal disease, although not to infection. However, immunocompromised and A/J mice have been reported to be more susceptible [4, 7, 17–19]. The proposed models are usually characterised by difficulty in handling and high costs, coupled with inconsistencies and large inter-laboratory variations in the results obtained. To find an easily handled and inexpensive experimental animal model, and on the assumption that specific host defences are lower in newborn animals, the susceptibility of suckling mice to legionellae was studied, with the principal aim of examining whether these animals could be useful in detecting consistent differences in virulence among clinical and environmental isolates of Legionella spp. [20, 21].

Materials and methods

Micro-organisms

L. pneumophila strains isolated from patients and from water supplies associated with human infection were used in this study. A few isolates were also serially
passaged on an artificial culture medium with the aim of selecting avirulent derivatives, one of which, referred to hereafter as Monza 3p50 strain, was re-inoculated four times into suckling mice to check for the definitive loss of virulence. The reference strains of L. pneumophila (Philadelphia 1, ATCC 33152), L. micdadei Tatlock (ATCC 33218), and L. bozemanii Toronto 3 (ATCC 33545), from type collection cultures, were also used. L. pneumophila strain Corby of known virulence [12] was provided by Dr R. B. Fitzgeorge, CAMR, Porton Down.

Micro-organisms were maintained at –80°C in skimmed milk; after thawing, they were grown at 36°C in a humidified atmosphere with CO₂ 2.5%.

Media and reagents

Buffered charcoal-yeast extract agar containing alpha-ketoglutarate 0.1% (BCYE-α agar; Unipath) [22] with and without selective supplement (glycine 3 g/L, polymyxin B 100 000 IU/L, vancomycin 5 mg/L and cycloheximide 80 ml/L) [23] were used as culture media for legionellae.

Subtyping of L. pneumophila serogroup 1 strains was performed according to the Oxford scheme [24] with monoclonal antibodies purchased from the John Radcliffe Hospital, Oxford.

Cells were examined by the direct immunofluorescence (DFA) test with a specific fluorescent antiserum against L. pneumophila serogroup 6 (Scimedx Corporation, NJ, USA).

Preparation of inocula

After growth for 2–4 days on BCYE-α agar, the micro-organisms were suspended in sterile distilled water. Cell concentrations were standardised optically with McFarland density standards and the number of viable organisms contained in the inoculum were counted as cfu.

For some strains, legionellae suspended in water were also killed by heating at 65°C for 30 min. In one experiment, bacterial cells were ruptured by sonicaton of the cell suspension. Cells were then filtered through membrane filters (Millipore, pore size 22 μm) and viable counts were made to calculate the number of disintegrated legionellae by comparison with bacterial counts before sonication.

Endotoxin assay

The Limulus amoebocyte rapid test (Whittaker Bioproducts Inc.; sensitivity 0.125 ng/ml) was performed according to the manufacturer’s instructions. Killed legionella suspensions, and cell-free sonicates of the same organisms, were tested to ascertain their endotoxin activity. Sterile pyrogen-free water was used as diluent.

Experimental model

The suckling mice were inoculated intraperitoneally (i.p.) with 10 and 20 μl of cell suspensions by the use of a Hamilton syringe. Intranasal (i.n.) challenge was also performed by depositing 5 μl of the legionella suspension with a micropipette on the external nares of unanaesthetised mice. Deaths were recorded for up to 1 month in the initial experiments, thereafter up to 15 days, and monitored as median survival time (MST, expressed in days) and number of dead over total (D/T) animals. Each experiment was repeated at least three times with different litters. In some experiments, heat-killed legionellae (80°C, 30 min) were used, while in others cell-free sonicates (corresponding to a dose of (2.5–3) × 10⁸ legionellae) of a freshly isolated strain of human origin, were injected i.p. into mice.

To assess the clearance of L. pneumophila from the whole body or internal mouse organs, mice were killed at different days (2–5) post-infection and the contents of the thoracic and abdominal cavities, with the exception of the stomach and the bowel, were removed aseptically together with a peritoneal wash, and homogenised with sterile distilled water. Legiolnella colony counts were done in duplicate on BCYE-α agar after serial dilutions. In other experiments, lung, spleen and liver were removed aseptically, rinsed twice in gentamicin 50 μg/ml solution, and allowed to remain for a further 1 h in gentamicin at room temperature, to kill the external legiolnellae. After rinsing twice with sterile distilled water, the organs were carefully homogenised in a mortar and vortex mixed. Serial dilutions were plated and colonies were counted. Guinea-pigs were also used to check the mortality after i.p. inoculation of a human fresh L. pneumophila isolate and its culture-passaged derivative. For this purpose, groups of four animals were inoculated i.p. with 1 ml suspensions of c. 10⁷, 10⁸ and 10⁹ legionellae from the Monza 3 and the Monza 3p50 strains, and any deaths were recorded up to 1 month.

Animals

Swiss albino CD1 pathogen-free mice (young pregnant females, 2-day-old litters with their mother and young adult males), and Hartley male guinea-pigs weighing c. 250 g were obtained from Nossan or Charles River breeding laboratories, Italy. The animals were housed in individual cages in a laminar flow housing cabinet, with an automatically controlled temperature of 22°C and 12 h of light. Newborn mice were allowed to live with their mother during the experimental period. The mice used were 3–6 days old, unless otherwise stated. The medium weight of a 5-day-old mouse was 3.3–3.5 g.

Mice were killed by exposure to ether or by cervical dislocation.
Mouse age-dependent susceptibility to L. pneumophila

To evaluate the dependency on mouse age of the outcome of L. pneumophila infections, groups of eight mice aged 3–16, 17–19, 20–26 and >26 days were inoculated with 6.5 x 10^7 legionellae from a suspension of L. pneumophila Monza 3 strain. Mortality was recorded for 15 days, and the MST calculated.

Resident peritoneal exudate cells

Resident peritoneal exudate cells (PEC) were obtained by washing the peritoneal cavity of infected or uninfected mice with 200 µl (suckling mice) or 3 ml (adult mice) of RPMI 1640 tissue culture medium (Microbiological Associates Bioproducts, Walkersville, MD, USA). During harvesting the cells were kept in an ice bath to prevent clumping. A small sample was mixed with trypan blue dye solution to check viability, and cells were counted with a haemocytometer. The PEC population was mostly composed of macrophages (Mp) and lymphocytes, the former amounting to >50%, as assessed morphologically. To count L. pneumophila, PEC suspensions were washed three times with RPMI medium by centrifugation at 1500 rpm for 5 min to remove non-internalised legionellae. The packed cells were lysed by addition of sterile distilled water and kept at room temperature for 30 min. The lysates were centrifuged as above to eliminate cell debris, the supernates were plated on BCYE-α agar, and the number of intracellular bacteria was determined as cfu.

Microscopy

Suspensions of washed PEC were distributed on to multi-spot slides, carefully air dried and fixed by depositing a small amount of iced (-80°C) methanol on each spot. DFA staining was performed by standard methods. Preparations were observed under UV light with a SM-Lux Leitz microscope. The three-dimensional localisation of legionellae within the cells was observed with a confocal laser scanning microscope (Phoibos 1000, Serastro Inc., Ypsilanti MI, USA).

RT-PCR analysis of TNFα mRNA expression in PEC of suckling mice

The reverse-transcriptase polymerase chain reaction (RT-PCR) was used to detect tumour necrosis factor (TNF-α) mRNA produced in mice challenged with L. pneumophila. Groups of 15–30 mice were inoculated with the two different L. pneumophila strains, as described above; an equal number of mice were inoculated with diluent alone as a control group. After 24 h, PEC were collected from mice by peritoneal washing with RPMI (0.5 ml/mouse) and total cellular RNA was extracted from PEC as described by Chomezynski and Sacchi [25]. Potential contaminating DNA was digested with DNAase (10 U/µl, 15 min, 37°C). The final RNA pellet was resuspended in RNAse-free H2O and reverse-transcribed for 60 min at 37°C in a 20-µl reaction mixture composed of oligo (dT 12-18) (Pharmacia, Uppsala, Sweden) 0.1 µg/ml, Moloney murine leukaemia virus reverse transcriptase (Gibco BRL, Sautersburg, MD, USA) 20 U and 20 mM dithiothreitol in Moloney murine leukaemia virus reverse transcriptase buffer (Gibco BRL).

cDNAs obtained by this procedure were amplified by PCR in the presence of specific primer pairs for murine TNF-α or β-actin gene sequences. 5' and 3' primer pairs were synthesised by Pharmacia (Uppsala, Sweden) with the following sequences: β-actin: TCCTGTGGCATCACGAAACTE and GAAACATTTGCGGTGGACGAT; TNF-α: ATCCCAGCITGGGAAGACTCCCTCCAG and GATCTCAAAGACAACACC-TAGTG.

PCR mixtures contained 0.05 µM of 5' and 3' primers, 2 mM deoxynucleotides, and 0.5 U of AmpliTaq polymerase in gene Amp PCR buffer (Gene-Amp Kit; Perkin-Elmer, Nonvalk, CO, USA), in a total volume of 20 µl. The reaction was conducted in a Perkin-Elmer thermal cycler for 30 cycles (40 s of denaturation at 94°C, 40 s of annealing at 62°C and 1 min extension at 72°C).

PCR products were visualised by agarose electrophoresis; HaeIII-digested φX174 DNA (Gibco BRL) was run in parallel as the mol.-wt markers. PCR products were identified by their predicted molecular mass.

For semi-quantitative measurements of TNF-α mRNA, serial five-fold dilutions of cDNA samples were subjected to PCR amplification of the β-actin gene sequence; the amount of the resulting PCR product was quantified with a Pharmacia LKB Ultrascan XL densitometer and plotted as a function of the input template cDNA to construct linear curves. For the comparative analysis of TNF-α mRNA expression by PEC of different groups of mice, dilutions of cDNA were normalised with these curves so as to yield an equivalent intensity of β-actin signal in all samples.

Statistical analysis

Mortality differences among the various groups of L. pneumophila-infected mice were assessed for their statistical significance by the Mann-Whitney U test or the Fisher's exact test, as appropriate.

Results

Preliminary infections in adult and newborn mice

Preliminary infections of suckling mice by i.n. or i.p. routes with L. pneumophila Monza 3 or after 50 passages in subculture (Monza 3p50) showed elevated mortality
after i.n. infection, with no difference among the strains. In contrast, mice inoculated i.p. with the serially passaged strain showed a significantly lower mortality than did mice inoculated with the original parent strain. In fact, MST and D/T ratio of animals inoculated with 3 x 10^8 cells of strains Monza 3 and Monza 3p50 were 1 day and 6/6, 15 days and 1/7, respectively.

When animals of different age were examined for their resistance to a challenge with the Monza 3 strain, the results clearly showed a progressive decrease in susceptibility to infection, and no mouse older than 1 month was killed by an inoculum as high as 3 x 10^8 bacterial cells (Table 1).

Viable Legionella cells were needed for lethal infection of suckling mice, as shown by the total lack of animal mortality following inoculation of up to 3 x 10^8 heat-killed or sonicated cells (data not shown). Equal amounts of viable or sonicated cells of both Monza 3 and Monza 3p50 strains gave similar positive results in the Limulus endotoxin assay.

Experimental infection of guinea-pigs confirmed the loss of the apparent 'virulence' of L. pneumophila strain Monza 3p50. In fact, groups of four animals inoculated i.p. with graded cellular suspensions up to 10^9 cfu of the strain Monza 3 showed an inoculum size-dependent mortality that reached 100%, while those receiving the same numbers of Monza 3p50 cells showed a 100% survival and no signs of illness during the 15 days of observation. Notably, the strain Monza 3p50 did not regain the ability to kill either the suckling mice or the guinea pigs after four in-vivo passages in suckling mice.

On the basis of these data, suckling mice aged 5–7 days were used in all further experiments and strains Monza 3 and Monza 3p50 were conventionally defined as virulent and avirulent strains of L. pneumophila, respectively.

**Virulence variations of different L. pneumophila strains**

Several L. pneumophila strains were compared for their virulence in the suckling mouse model. No significant differences were detected among strains of the different serogroups examined, provided that they were freshly isolated from human sources. All of them caused c. 50% mortality and an MST ranging from 1 to 5 days after inoculation of (1–3) x 10^7 cells. The two environmental isolates of L. pneumophila were apparently less lethal than human isolates (see the MST values in Table 2).

**Clearance of L. pneumophila by suckling mice**

Mice were examined for the persistence of L. pneumophila in their organs after i.p. challenge with the virulent or the avirulent strain. As shown in Table 3, spleen and liver of three mice randomly taken from the different groups yielded a high number of legionellae 5 h after inoculation with either the virulent or the avirulent strain, and this trend was generally apparent at 24 h. Seven days after infection, more legionellae of the avirulent strain than of the virulent one were recovered from the above organs. In contrast, more legionellae were recovered from the lungs of mice infected with the virulent strain than from the lungs of those inoculated with the avirulent strain, even

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**Table 1. Age dependence susceptibility of Swiss CD1 mice to intraperitoneal injection of L. pneumophila strain Monza 3 (serogroup 6)**

<table>
<thead>
<tr>
<th>Age of mice (days)</th>
<th>D/T (%)</th>
<th>MST (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–16</td>
<td>7/8 (87.5)</td>
<td>3</td>
</tr>
<tr>
<td>17–19</td>
<td>4/8 (50.0)</td>
<td>4</td>
</tr>
<tr>
<td>20–26</td>
<td>1/8 (12.5)*</td>
<td>&gt;15*</td>
</tr>
<tr>
<td>30–35</td>
<td>0/10 (0.0)*</td>
<td>...</td>
</tr>
<tr>
<td>60†</td>
<td>0/10 (0.0)</td>
<td>...</td>
</tr>
</tbody>
</table>

The inoculum comprised 6.5 x 10^7 cells; MST, median survival time; D/T, dead over total animals within 15 days of observation.

*Statistically significant difference (p < 0.01) between the indicated values and the D/T value of 0–16-day-old mice (as assessed by Fisher's exact test) or the MST value of either the 0–16 or the 17–19-day-old animals (as assessed by the Mann-Whitney U test).
†Inoculated with 3 x 10^8 L. pneumophila cells.

**Table 2. Mortality of suckling mice after intraperitoneal inoculation with fresh isolates of L. pneumophila**

<table>
<thead>
<tr>
<th>Strain (serogroup)</th>
<th>MAb subtype*</th>
<th>Origin</th>
<th>Mortality (1 x 10^7 cells/mouse) D/T</th>
<th>MST (days)</th>
<th>Mortality (3 x 10^7 cells/mouse) D/T</th>
<th>MST (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monza 3 (6)</td>
<td>...</td>
<td>Human</td>
<td>5/10</td>
<td>3</td>
<td>6/6</td>
<td>1</td>
</tr>
<tr>
<td>Messina 2 (3)</td>
<td>...</td>
<td>Human</td>
<td>6/10</td>
<td>1</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Trento 1 (1)</td>
<td>Pontiac</td>
<td>Human</td>
<td>5/6</td>
<td>2</td>
<td>4/4</td>
<td>2</td>
</tr>
<tr>
<td>Trento 2 (1)</td>
<td>Pontiac</td>
<td>Human</td>
<td>3/6</td>
<td>5</td>
<td>8/8</td>
<td>1</td>
</tr>
<tr>
<td>Trento 3 (1)</td>
<td>Pontiac</td>
<td>...</td>
<td>3/11</td>
<td>&gt;30†</td>
<td>5/8</td>
<td>2</td>
</tr>
<tr>
<td>Roma 9 (14)</td>
<td>...</td>
<td>Human</td>
<td>2/5</td>
<td>&gt;30†</td>
<td>ND†</td>
<td>ND‡</td>
</tr>
<tr>
<td>Forio 4 (1)</td>
<td>Olda</td>
<td>Environment</td>
<td>ND</td>
<td>ND</td>
<td>5/9</td>
<td>7†</td>
</tr>
</tbody>
</table>

D/T, dead/total; MST, median survival time; ND, not done.

*Of the Oxford L. pneumophila serogroup 1 monoclonal subtyping scheme.
†Statistically significant (p < 0.01) difference between the indicated values and those of all other groups as assessed by the Mann-Whitney U test.
‡After an inoculum of 2.2 x 10^7, D/T was 6/10, MST was 3.
Presence of L. pneumophila in peritoneal macrophages of suckling mice

Because of the characteristic intramacrophage localisation of L. pneumophila [16,26], the presence of the micro-organism in the PEC of suckling mice was examined, and for comparison, in those of adult mice. The presence of L. pneumophila in suckling mice was demonstrated by immunofluorescence and confocal microscopy. At 24–48 h after challenge with the virulent strain, some Mc contained numerous legionellae within a cytoplasmic vesicle, others appeared completely overwhelmed by the micro-organism (Fig. 1). After infection with the avirulent derivative, fewer Mc appeared to be infected and few legionellae were seen within each infected cell (Fig. 1). Confocal micrographs of PEC from suckling mice taken at different levels of the peritoneal cavity showed intracellular legionellae (Fig. 2).

L. pneumophila cfu counts demonstrated roughly equal numbers of total PEC-associated microbial cells, but it should be remembered that the virulent strain induced a much greater PEC recruitment in the peritoneal cavity than the avirulent strain (about 1 log₁₀ difference 6 h after infection) (data not shown).

A similar pattern of intramacrophage localisation of the two Legionella strains was observed in PEC from adult mice 24–48 h after infection. Five and 10 days after infection the macrophages of adult mice showed fragments and a speckled appearance of immunofluorescence, probably due to digested bacteria.

TNF-α mRNA expression in PEC of suckling mice infected by L. pneumophila

Because L. pneumophila is known to induce TNF-α production by both human and murine Mc [27,28], TNF-α production by PEC of suckling mice challenged by either the virulent or the avirulent L. pneumophila strain was examined. In preliminary experiments, cytokine production was assayed both in the peritoneal fluid (as protein, by ELISA) and in PEC (as mRNA transcript by RT-PCR). As very low and inconsistent amounts of TNF-α were detected in the peritoneal fluid, all subsequent experiments addressed TNF-α mRNA production. As shown in Fig. 3, a strong TNF-α mRNA signal was detected in mice infected with L. pneumophila Monza 3, contrasting with a very weak signal in animals infected with Monza 3p50, and no signal at all in mice given the diluent only (negative control). Densitometric measurements of TNF-α message (expressed as density ratio between TNF-α and β-actin mRNA signals) gave the following results: 1.40 versus 0.30 (first experiment) and 0.70 versus 0.10 (second experiment) for mice challenged with Monza 3 and Monza 3p50, respectively, thus indicating 5–7-fold greater TNF-α mRNA production in mice challenged with the virulent strain.

Discussion

In-vivo studies aimed at evaluating bacterial virulence are essential to unravel host–parasite relationships. In the context of the virulence of Legionella strains for
Fig. 1. Peritoneal exudate cells (PEC) from Swiss CD1 suckling mice infected with *L. pneumophila*, stained by direct immunofluorescence 48 h after infection. Fluorescence (left) and light (right) micrographs show the same microscopic fields. **a**, PEC infected with the virulent *L. pneumophila* strain Monza 3: some cells exhibit numerous fluorescent bacteria, others appear to be completely engulfed. **b**, PEC infected with the avirulent derivative Monza 3p50: few cells appear to be infected and few legionellae are seen within the cells. Magnification ×570.

Fig. 2. Peritoneal monocytic cell from Swiss CD1 suckling mouse infected with the virulent *L. pneumophila* strain Monza 3, stained by direct immunofluorescence. Confocal scanning micrographs taken at different levels of the cell: **a**, tangential to the cell membrane; **b**, cross-sectional to cell nuclear substance, showing intracellular legionellae. Magnification ×2350.

normal laboratory animals, only guinea-pigs have so far been demonstrated to be suitable experimental models [15]. However, these animals are not easily amenable to immunological investigations. Normal adult mice are usually highly resistant to experimental legionellosis [15, 16, 29], even though immunocompromised mice [4, 7, 17, 19] and the genetically susceptible A/J mice [18, 30] have been shown to support growth of
Legionellae in the lung, particularly after intra-tracheal infection [31]. L. pneumophila replicates well in humans and guinea-pigs MΦ as well as in those of A/J mice [29, 32–35]. Growth is limited in MΦ from other mouse species, while it is supported by those of golden hamster and rat, although these animals are resistant to infection [29]. In general, there is conflicting evidence about replication in mouse MΦ and, more generally, on the relative validity of inhalatory versus non-inhalatory routes of infection in mimicking, at least in part, human disease [11, 36].

On the assumption that adult mouse resistance to L. pneumophila could be due to the development of the immune system, and that host defences are lower in infant animals, the susceptibility of suckling mice to infection was studied, with the main purpose of verifying whether the model adopted could be suitable and sensitive enough to detect virulence differences among isolates of L. pneumophila.

These studies indicate that suckling CD1 mice are susceptible to lethal i.p. challenge with viable L. pneumophila strains of fresh clinical isolation, but not with those from long-maintained and repeatedly passaged cultures, in keeping with the observed low mortality rate after inoculation into mice of laboratory strains from stock cultures. However, it was also observed that i.n. inoculation of mice with L. pneumophila brought about an elevated mortality regardless of the type of the strain used for challenge. It is difficult to explain why virulence differences are not evidenced by i.n. infection but are shown by i.p. infection.

Hedlund and co-workers [17] demonstrated replication of legionellae only in the spleen of adult AKR/J susceptible mice challenged i.p. with as high as $2 \times 10^8$ virulent legionellae. The histopathology of the liver of infected animals was suggestive of toxic or anoxic damage rather than bacterial infection. Hedlund [37] also reported the death of animals infected with a cell-free sonicate of legionellae. In the present study, the death of CD1 suckling mice after i.p. inoculation required viable legionellae, as high inocula of heat-inactivated cells ($>10^8$) or sonicates of L. pneumophila cultures were totally ineffective. Endotoxin production appears not to be involved, as the Limulus test was equally positive for suspensions of both virulent and avirulent L. pneumophila strains, and for the crude toxin, but all mice inoculated with the filtrate from c. $3 \times 10^8$ sonicated legionellae survived.

The basis for virulence differences, as detected in the described animal model, are unknown. It should be stressed here that the above model cannot be considered as truly representative of a reproductive infection by L. pneumophila. Given the high inoculum used for infection, what was essentially measured was a difference in the rate of clearance of the legionellae by the mouse. However, the above difference appears to be a sensible correlate of virulence, as it was stably reproduced in different experiments, required viable L. pneumophila cells and was also observed in control experiments with guinea-pigs, which are ordinarily susceptible to L. pneumophila infection. In this context, a potentially relevant observation is that the virulent L. pneumophila strain is less rapidly cleared from lungs (but not from liver and spleen) than the avirulent strain. It would be important in future studies to assess alveolar MΦ of suckling and adult mice for intracellular growth permissiveness of strains Monza 3 and Monza 3pS0. PEC of suckling mice were found to be more heavily parasitised by Monza 3 than by Monza 3p50 cells. Whether this implies some permissiveness of the MΦ of these animals to intracellular growth of L. pneumophila cannot be stated at the moment. However, the data would indicate that, in contrast to MΦ of adult mice, those of the infant mice take up and internalise the virulent strain to a greater extent than the avirulent strain or (alternatively or concurrently) they kill the latter more efficiently than the former.

Relevant to this is the fact that L. pneumophila may exert a direct toxic effect on host cells without necessarily invading or multiplying in them [38]. As pro-inflammatory cytokines, in particular TNF-α, are strongly induced by legionellae [27], TNF-α production by PEC of animals challenged with virulent or avirulent strain was investigated. TNF-α mRNA was found to be expressed more in PEC from mice challenged with the former strain, although this could not be confirmed by the actual cytokine detection in the peritoneal fluid (probably because of its low quantity or rapid binding, neutralisation or degradation). It is possible that – together with the production of other toxic shock-related cytokines, such as, for instance, IL-1β [39] – TNF-α production could rapidly bring the animal to toxic shock and death. TNF-α is currently considered to be an important
factor facilitating the resolution of *L. pneumophila* infection [40] but, as in other biological situations, localised and rapid induction of this cytokine in a newborn animal may also exert an intense toxic effect and cachexia [41].

Although no specific virulence factor or mechanism has been sought in this study, these results suggest that suckling CD1 mice are a promising, easily handled and inexpensive animal model for studies of *Legionella* virulence. With this model, virulence of fresh clinical isolates of *L. pneumophila* was demonstrated, and seen to be attenuated or even totally lost after repeated subcultures on artificial media, a classic biological parameter of bacterial virulence attenuation. Interestingly, virulence was accompanied by a longer persistence of a greater burden of legionellae in the lungs, and a greater capacity for stimulating TNF-α production in the peritoneal cavity, both observations being potentially relevant to human pathology. Finally, the model showed a very low variability in the susceptibility of different litters of mice and thus has good reproducibility. Studies are in progress to determine specific virulence factors of *Legionella* spp. and modulation of the host cytokine response in mouse organs.

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**References**

VIRULENCE OF LEGIONELLA SPP. IN SUCKLING MICE