

## *icmT* Is Essential for Pore Formation-Mediated Egress of *Legionella pneumophila* from Mammalian and Protozoan Cells

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The final step of the intracellular life cycle of *Legionella pneumophila* and other intracellular pathogens is their egress from the host cell after termination of intracellular replication. We have previously isolated five spontaneous mutants of *L. pneumophila* that replicate intracellularly similar to the wild-type strain but are defective in pore formation-mediated cytolysis and egress from mammalian and protozoan cells, and the mutants have been designated *rib* (release of intracellular bacteria). Here, we show that the *rib* mutants are not defective in the activity of enzymes secreted through the type II secretion system, including phospholipase A, lysophospholipase A, and monoacylglycerol lipase, although they are potential candidates for factors that lyse host cell membranes. In addition, the *pilD* and *lspG* mutants, which are defective in the type II secretion system, are not defective in the pore-forming toxin. We show that all five *rib* mutants have an identical point mutation (deletion) following a stretch of poly(T) in the *icmT* gene. Spontaneous revertants of the *rib* mutants, due to an insertion of a nucleotide following the poly(T) stretch in *icmT*, have been isolated and shown to have regained the wild-type phenotype. We constructed an *icmT* insertion mutant (AA100kmT) in the chromosome of the wild-type strain by allelic exchange. The AA100kmT mutant was as defective as the *rib* mutant in pore formation-mediated cytolysis and egress from mammalian and protozoan cells. Both the *rib* mutant and the AA100kmT mutant were complemented by the *icmT* gene for their phenotypic defect. *rtxA*, a gene that is thought to have a minor role in pore formation, was not involved in pore formation-mediated cytolysis and egress from mammalian and protozoan cells. We conclude that the *icmT* gene is essential for pore formation-mediated lysis of mammalian and protozoan cells and the subsequent bacterial egress.

*Legionella pneumophila*, the etiologic agent of Legionnaires' disease, is a gram-negative bacterium, ubiquitous in the aquatic environment, where it survives as an intracellular parasite of amoebae and ciliated protozoa (21, 31). In humans, *L. pneumophila* replicates within alveolar macrophages and possibly epithelial cells (1, 19) and utilizes similar mechanisms to parasitize the protozoan host (18, 36). After phagocytosis by mammalian and protozoan cells, the bacteria modulate the biogenesis of their vacuole into a niche that permits it to escape the classical endosomal-lysosomal degradation pathways (9, 23). This vacuole is subsequently surrounded by mitochondria and rough endoplasmic reticulum (2, 22). A group of 23 genes designated *dot* (defect in organelle trafficking)/*icm* (intracellular multiplication) are implicated in modulating the formation of the replicative vacuole and subsequent intracellular replication (33, 38). The *dot/icm* complex is thought to constitute a type IV-like secretion system capable of transferring effector molecules into the host cell to evade the endocytic pathways.

A fundamental step in the pathogenic life cycle of intracellular pathogens is their ability to lyse and egress from the host cell after termination of intracellular replication, which subsequently leads to infection of uninfected neighboring cells or to transmission to a new host. The mechanisms by which intracellular pathogens egress from the host cell are not well un-

derstood. It has been presumed that the physical and metabolic burden on the host cell by a large number of intracellular bacteria is sufficient to rupture the host cell by nonspecific means. Recent data about *L. pneumophila* have shed some light on a specific and pathogen-regulated mechanism for bacterial egress from the host cell. After termination of intracellular replication within mammalian and protozoan cells, *L. pneumophila* induces cytolysis of the host cell, which is mediated by a pore-forming activity (6, 17). This pore formation-mediated cytolysis is triggered in vitro and in vivo upon growth transition into the postexponential phase (6, 10). Five mutants of *L. pneumophila* defective in the pore-forming toxin are not defective in modulating biogenesis of their replicative vacuole and replicate intracellularly similar to the wild-type strain. However, these mutants are defective in egress from mammalian and protozoan cells upon termination of intracellular replication, and the mutants have been designated *rib* (release of intracellular bacteria) (6). However, the mutants are released at a later time, most likely due to apoptotic changes in the infected cell (6). We propose to designate the pore forming-toxin Rib toxin. The *rib* mutants are also defective in acute cytotoxic lethality to mice (6). We have shown that the miniTn10::Kan insertions used to generate these five mutants are located in different chromosomal regions distinct from the two regions of the *dot/icm* genes (6). However, cosmid clones harboring the chromosomal regions defective in the mutants failed in complementation studies for the defect in the pore-forming toxin. In addition, reconstruction of the original five mutants by allelic exchange of the Kan-inserted loci into the

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wild-type chromosome showed that the newly constructed mutants are indistinguishable from the wild-type strain in intracellular replication, cytotoxicity, and pore formation-mediated egress from mammalian and protozoan cells. Therefore, the mutations in the five mutants are spontaneous (6).

In this report, we show that the *rib* mutants are not defective in any of the enzymes secreted through the type II secretion system, which are potential factors that disrupt host cell membranes. This was further confirmed by the *pilD* and *lspG* mutants, both of which are defective in the type II secretion system and both of which are not defective in the pore-forming toxin. We show that the *icmT* gene is the defective locus in the *rib* mutants and is essential for egress of intracellular bacteria from both U937 macrophages and *Acanthamoeba polyphaga* upon termination of intracellular replication. The defect in *icmT* is a point deletion after a stretch of poly(T) in the carboxy terminus of the gene. In addition, we show that the *rtxA* gene that has been thought to be required for pore formation (12) is not required for the pore formation-mediated egress from macrophages and protozoa.

#### MATERIALS AND METHODS

**Bacterial strains and media.** *L. pneumophila* serogroup I strain AA100 was the parental strain used for all experiments in this study. *L. pneumophila* strains were grown on buffered charcoal-yeast extract (BCYE) plates or in buffered yeast extract broth. All the strains and the plasmids used in this study are described in Table 1. For *L. pneumophila*, antibiotics were added to the media at the following concentrations: kanamycin, 50 µg/ml; chloramphenicol, 2.5 and 5 µg/ml. *Escherichia coli* strains were cultured on Luria-Bertani (LB) agar plates or in LB broth. Antibiotics were added to the media at the following concentrations: kanamycin, 50 µg/ml; chloramphenicol, 20 µg/ml.

**Cell culture.** U937 cells were cultured in RPMI 1640 containing 10% fetal bovine serum (Gibco). U937 cells were differentiated with phorbol 12-myristate 13-acetate (Sigma) for 48 h before use as previously described (18). *A. polyphaga* cells were maintained in peptone yeast glucose medium. Glucose was omitted from the medium to prevent growth of amoebae during infection with *Legionella*.

**DNA manipulations.** Transfections, restriction enzyme digestions, and DNA manipulation were performed as previously described (5). Restriction enzymes and T4 DNA ligase were purchased from Promega (Madison, Wis.). *L. pneumophila* chromosomal DNA was prepared by using the Puregene DNA isolation kit (Gentra Systems, Minneapolis, Minn.). Plasmid and cosmid DNA preparations were performed with the Bio-Rad (Hercules, Calif.) Quantum miniprep kit and by the polyethylene glycol DNA extraction procedure as described before (32). Electroporations were performed with a Bio-Rad Gene Pulser, as recommended by the manufacturer. Purification of DNA fragments from agarose gels for subcloning was carried out with a QIAquick gel purification kit (Qiagen Inc., Chatsworth, Calif.). Oligonucleotide synthesis for PCR and sequencing were performed by Integrated DNA Technologies, Inc. (Coralville, Calif.). Sequencing was carried out by ACGT Inc. (Northbrook, Ill.). Sequence analysis and comparisons were carried out with DNAMAN software and NCBI molecular biology software (www.ncbi.nlm.nih.gov).

**Enrichment of *rib* transconjugants in *A. polyphaga*.** A cosmid library of *L. pneumophila* in *E. coli* was mobilized into the *rib* mutants GN229 and GL208 by conjugation. The selection was carried out on BCYE containing kanamycin (50 µg/ml) and chloramphenicol (5 µg/ml). The resulting transconjugants were pooled and frozen at -80°C until use. The pools were used for infection of *A. polyphaga* cells to enrich for clones that were able to egress from *A. polyphaga* after intracellular replication. Briefly, *A. polyphaga* was infected with the pool of transconjugants of the *rib* mutant GN229 at a multiplicity of infection (MOI) of 10 in six-well tissue culture dishes containing 10<sup>6</sup> *A. polyphaga* per well, and the dishes were incubated for 1 h at 37°C. At the end of the infection period, the monolayers were treated with gentamicin (50 µg/ml). The bacteria released into the culture medium of amoebae after 48 h incubation were used to infect freshly grown *A. polyphaga* as described above, and the supernatant of the infection was obtained after 24 h of incubation. Three rounds of enrichment were carried out. Representatives of the transconjugant clones that egress from amoebae were used in a contact-dependent hemolysis assay and cytotoxicity assay for U937 cells.

TABLE 1. Strains and plasmids used in this study

Strain or plasmid	Genotype or sequence	Reference or source
<b>Strains</b>		
<i>L. pneumophila</i>		
AA100		3
GN229	Kan <sup>r</sup>	6
GR159	Kan <sup>r</sup>	6
GL208	Kan <sup>r</sup>	6
GP247	Kan <sup>r</sup>	6
GP263	Kan <sup>r</sup>	6
AA100kmT	AA100; <i>icmT</i> :: Kan <sup>r</sup>	This study
AA100kmS	AA100; <i>icmS</i> :: Kan <sup>r</sup>	This study
ψlp24	ΔRtxA	13, 14
NU259	130b; LpsG::Kan <sup>r</sup>	30
NU243	130b; PilD::Kan <sup>r</sup>	28
ribrev1		This study
ribrev2		This study
ribrev3		This study
<i>E. coli</i> DH5α	<i>supE44</i> Δ <i>lacU169</i> (ψ80 <i>lacZ</i> ΔM15) <i>hsdR17 recA1 endA1 gyrA96 thi-1 relA1</i>	BRL
<b>Plasmids</b>		
pBC SK+	ColE1 origin, <i>lacZ</i> α, Cm <sup>r</sup>	Stratagene
pMW100	pMMB207αβ; Cm <sup>r</sup> ; <i>icmGCDJB</i> , <i>tphA</i> , and <i>icmF</i> region	29
pGS-LC-32	pMMB207αβ; Cm <sup>r</sup> ; <i>icmTSRQPO</i>	34
pGS-LC-47	pMMB207αβ; Cm <sup>r</sup> ; <i>icmNMLKEG</i>	33
pTSRQ	pBC SK+; Cm <sup>r</sup> ; <i>icmTSRQ</i>	This study
pTS	pBC SK+; Cm <sup>r</sup> ; <i>icmTS</i>	This study
pT	pBC SK+; Cm <sup>r</sup> ; <i>icmT</i> gene	This study
p <sub>kan</sub> T	pBC SK+; Cm <sup>r</sup> ; ; <i>icmT</i> :: Kan <sup>r</sup>	This study
pS	pBC SK+; Cm <sup>r</sup> ; <i>icmS</i> gene	This study
p <sub>kan</sub> S	pBC SK+; Cm <sup>r</sup> ; ; <i>icmS</i> :: Kan <sup>r</sup>	This study
pR	pBC SK+; Cm <sup>r</sup> ; <i>icmR</i> gene	This study
p <sub>kan</sub> R	pBC SK+; Cm <sup>r</sup> ; <i>icmR</i> :: Kan <sup>r</sup>	This study
pQ	pBC SK+; Cm <sup>r</sup> ; <i>icmQ</i> gene	This study
p <sub>kan</sub> Q	pBC SK+; Cm <sup>r</sup> ; <i>icmQ</i> :: Kan <sup>r</sup>	This study

**Genetic characterization of the *rib* mutants.** Three plasmids (pMW100, pGS-LC-47, and pGS-LC-32) containing different regions of *dot/icm* genes (kind gifts of H. Shuman) (Table 1) were used to transform the *rib* mutants by electroporation. The *icmTSRQ* locus was amplified from AA100 genomic DNA, and *SalI* and *NorI* sites were introduced in the forward (P1) and reverse (P2) primers (Table 2), respectively, by PCR using long-template *Taq* (Boehringer Mannheim). The annealing temperature used was 50°C, with the extension temperature of 68°C for 25 cycles. The PCR product was purified from the agarose gel and ligated to the *SalI*- and *NorI*-digested pBC sk+ plasmid. The resulting ligation mixture was used to transform DH5α. *icmR* and *icmQ* genes were cloned in a similar fashion as described above to create pR and pQ, respectively. *AvaII* digestion was used to delete the *icmRQ* region from the plasmid pTSRQ, hence resulting in pTS. Plasmid pT contains an in-frame deletion of the *icmS* gene created by inverse PCR of the plasmid pTS using the primers P3 and P4 (Table 2) with the *NheI* restriction enzyme sequence linked to the 5' end of the primers. The same strategy was utilized in creating in-frame deletion of the *icmT* gene with the resulting plasmid pS, using primers P5 and P6 (Table 2). The following primer combinations were used for amplification of different *icm* genes: P7 and P8 for the *icmR* gene and P9 and P2 for the *icmQ* gene.

In vitro transposon mutagenesis was carried out on the plasmid of choice using an EZ::TN<KAN-2> insertion kit as described by the manufacturer (Epicenter Technologies, Madison, Wis.). A transposon library of the plasmid was used to transform *E. coli* DH5α. Transformants were selected on LB agar plates containing kanamycin-chloramphenicol and screened by PCR for location of insertion. The resulting plasmid extracted from *E. coli* DH5α was used to transform the *rib* mutant. Some of the plasmid constructs containing the transposon kanamycin insertion were used for allelic exchange to *L. pneumophila* AA100 chromosome by natural transformation (37).

TABLE 2. Primers used in this study

Primer	Sequence <sup>a</sup>	Restriction enzyme
P1 (salIIcmTS)	ACGCGTCGACCACAGTAAACTTCAAGCTGA ACC	<i>SalI</i>
P2 (notIRO2)	ATAAGAATGCGGCCGCTGCTCAGAGCTATTTTT	<i>NotI</i>
P3	CTAGCTAGCACACTTGCTAATATCTCGCT	<i>NheI</i>
P4	CTAGCTAGCCGTTTGTATGTATGACGG	<i>NheI</i>
P5	CTAGCTAGCCAGTGGCTCGTTGCT	<i>NheI</i>
P6	CTAGCTAGCGTTCAGTTAGATCTGCTAG	<i>NheI</i>
P7	ACGCGTCGACATGAGCTTATTCAAGGGGGC	<i>SalI</i>
P8	ATAAGAATGCGGCCGCGGAACCAAGAATTAG GGG	<i>NotI</i>
P9	ACGCGTCGACAATGTTTGGTGCTGCTGAATC	<i>SalI</i>

<sup>a</sup> Underlined letters are the sequences of restriction enzymes used for cloning.

Complementation of the defects was carried out for the resulting transformants by contact-dependent hemolysis assay, cytolysis of *A. polyphaga* and U937 cells following intracellular replication, and cytotoxicity assay as described below.

**Contact-dependent pore formation assay.** Contact-dependent pore formation in plasma membrane was determined by examining hemolysis of sheep red blood cells (sRBCs) by *L. pneumophila* at an MOI of 25 following 2 h of bacterium-sRBC contact, as previously described (6, 26).

**Cytotoxicity of *L. pneumophila* to U937 macrophages and *A. polyphaga*.** *L. pneumophila* strains were grown on BCYE plates for 3 days prior to infection of U937 macrophages or amoebae. Infection was performed, in triplicate, in 96-well plates containing  $10^5$  cells/well at an MOI of 1 for 1 h for U937, and in 24-well plates with coverslips containing  $5 \times 10^5$  cells/well at an MOI of 10 for *A. polyphaga*. The infection period was followed by gentamicin (50 µg/ml) treatment for 1 h, which was followed by three washes to remove extracellular bacteria. At several time points, the monolayers of macrophages were treated for 4 h with 10% Alamar Blue dye (Alamar Bioscience Inc., Sacramento, Calif.) as previously described (4). Viability of the monolayers of macrophages was determined by measurement of optical density (OD) of Alamar Blue-treated monolayers by using a VMAX Kinetic Microplate reader (Molecular Devices, Menlo Park, Calif.) and expressed as percent cell death compared to uninfected cells by using the formula  $[1 - (\text{mean OD of treated cells} / \text{mean OD of untreated cells})] \times 100\%$ .

In *A. polyphaga* infection experiments, the cytolysis of amoebae was documented by images of the infected monolayers at different time intervals to provide a qualitative assessment of the cytolysis. Cells were observed with an Axiophot Photomicroscope (Carl Zeiss, Inc., Oberkochen, Germany).

**Growth kinetics of released *L. pneumophila* strains from U937 macrophages.** Infections of U937 macrophages by *L. pneumophila* strains were performed, in triplicate, in 96-well plates containing  $10^5$  cells/well at an MOI of 1 for U937 macrophages. At the end of the infection period, the monolayers were treated with gentamicin (50 µg/ml) for 1 h as described above. The number of released bacteria in the monolayers at several time intervals after washing of the gentamicin was determined.

**Lipolytic activities in supernatants and cell lysates.** The bacteria were grown in buffered yeast extract broth to an OD at 600 nm ( $OD_{600}$ ) of 1.85 (late-logarithmic phase). The bacteria were pelleted and lysed with Triton X-100-lysozyme, and the volume of the cell lysate was adjusted to the volume of the supernatant. Due to their ability to release high levels of fatty acids from dipalmitoylphosphatidylglycerol (DPPG) and monopalmitoylphosphatidylcholine, dilutions of 1:2 or 1:10 of the lysates were made, prior to the enzymatic assays.

Phospholipase A (PLA), lysophospholipase A (LPLA), and monoacylglycerol lipase activities were determined as we described previously (7, 16). Briefly, PLA activity was determined by hydrolysis of the substrates DPPG and dipalmitoylphosphatidylcholine (final concentrations, 5 mg/ml). The LPLA activity was determined by hydrolysis of the substrate monopalmitoylphosphatidylcholine (1-MPG) (final concentration, 3.4 mg/ml). Mono-acylglycerol lipase activity was determined by hydrolysis of the 1-monopalmitoylglycerol substrate (final concentration, 2.2 mg/ml).

## RESULTS

**Enrichment for cosmid clones to complement the *rib* mutants for their defect in egress from *A. polyphaga*.** Since the *rib*

mutants are completely defective in egress from *A. polyphaga*, we exploited this phenotype to isolate cosmid clones that complement the mutants for egress from amoebae. The cosmid library of *L. pneumophila* was mobilized into the *rib* mutants GN229 and GL208, and the two pools of transconjugants were used, separately, to infect *A. polyphaga*. The bacteria released into the culture medium after 48 h incubation were used to infect freshly grown *A. polyphaga* to enrich for transconjugants that are capable of egress. The two mutants (without the cosmid library) were used as controls in these infections to monitor the increase in the number of GN229 and GL208 transconjugants that are able to egress compared to the mutants. The defect in egress for the two *rib* mutants was completely complemented to the wild-type level after three enrichments for transconjugant clones that were able to egress from *A. polyphaga*. In contrast, the defect in egress of the two mutant controls was stable during the three enrichments (data not shown). Surprisingly, several trials have failed to isolate cosmids from 30 random transconjugants of GL208 and 15 random transconjugants of GN229, which regained the wild-type phenotype for egress from *A. polyphaga*. We concluded that these transconjugants were spontaneous revertants that have been able to repair the genetic lesion responsible for the *rib* phenotype.

**Lipolytic activities of enzymes secreted through the type II secretion system by the *rib* mutants.** Many lipolytic enzymes secreted by the type II secretion system of *L. pneumophila* (*lsp*, for *Legionella* secretion pathway) are potential candidates for enzymes that may lyse the host cell membrane by the hydrolysis of lipid constituents or generation of the hemolytic agent lysophosphatidylcholine. Therefore, we tested the *rib* mutants for these activities (7, 16). Our data show that none of the previously described enzyme activities of PLA, LPLA, and monoacylglycerol lipase were reduced in the *rib* mutant GN229 (Fig. 1A and 1B). On the contrary, all activities were slightly higher in the *rib* mutant than in the wild-type strain. The acid phosphatase enzyme, which is also secreted by the type II secretion system (30), was also elevated in culture supernatants of the *rib* mutants (data not shown). It is not clear why the activities are elevated in the *rib* mutant.

The data were further confirmed by examination of the pore-forming activity, using contact-dependent hemolysis of sRBCs, of the two mutants defective in the type II secretion system. These mutants are NU259, which carries a Kan insertion in *lspG*, which encodes an inner membrane structural protein, and the NU243 mutant, which is disrupted in *pilD*, the prepilin peptidase gene. The result showed that these two mutants were indistinguishable from the wild-type strain in pore formation activity (data not shown). Thus, the type II secretion system is not required for the pore-forming activity, and the *rib* mutants are not defective in lipolytic enzymes secreted by the type II secretion system.

**The *icmTSRQPO* locus complements the phenotype of the five *rib* mutants.** Since the *dot/icm* genes have been shown to be involved in the pore-forming activity of *L. pneumophila* (26), we decided to introduce different regions of the 23 *dot/icm* genes to the *rib* mutants for possible complementation. The *icmTSRQPO* region; the *icmGCDJB*, *tphA*, and *icmF* regions; and the *icmNMLKEG* region were tested for the ability to restore the pore-forming toxin activity to the *rib* mutant

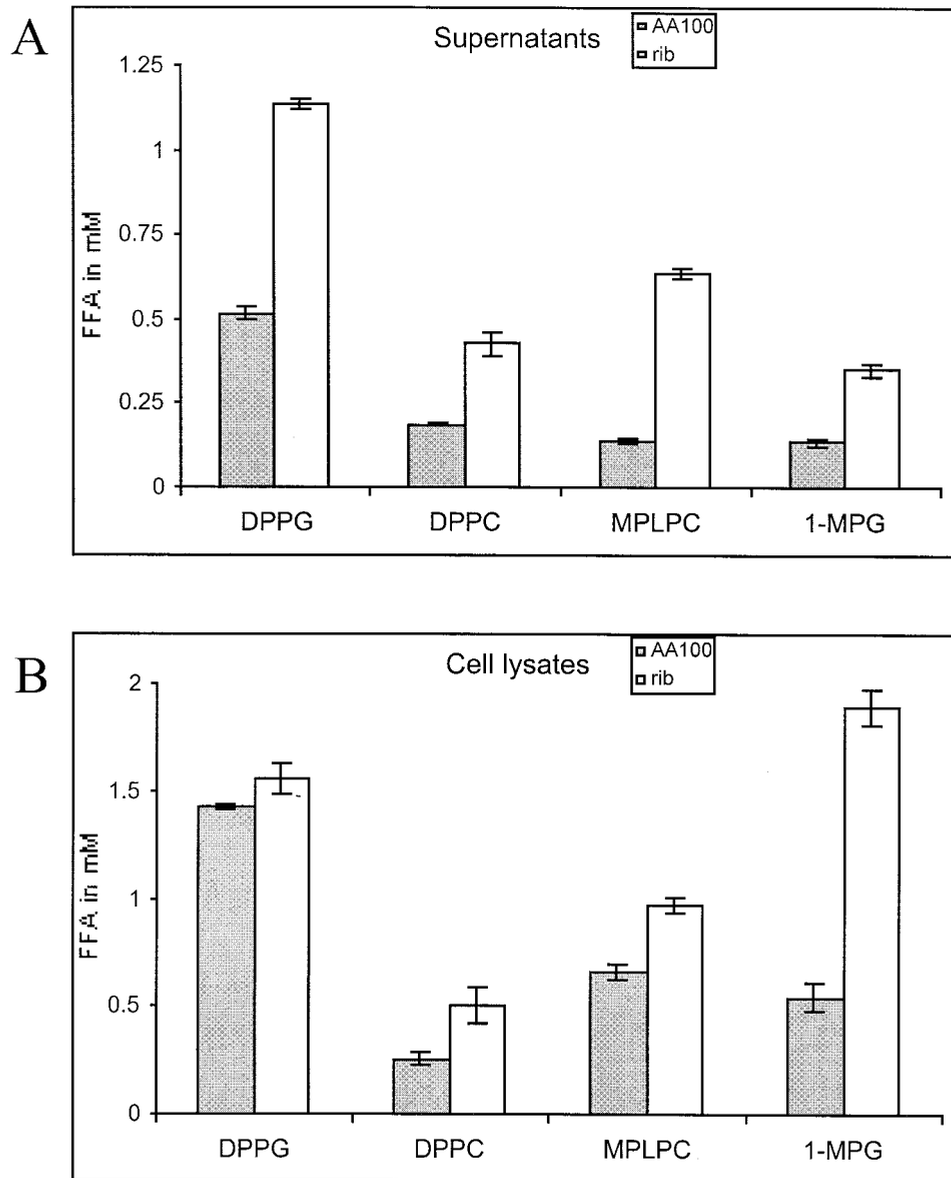


FIG. 1. The *rib* mutants are not defective in lipolytic enzymes secreted through the type II secretion system. Lipid hydrolysis by supernatants (A) and cell lysates (B) of AA100 and the *rib* mutant was tested for the ability to release free fatty acids (FFA) from dipalmitoylphosphatidylcholine (DPPC), DPPG, monopalmitoylphosphatidylcholine (MPLPC), and 1-MPG. These results are representative of two additional experiments, with the exception of a variation observed between experiments for the cell lysates activity of the wild-type strain and the mutant on 1-MPG (data not shown). Error bars represent standard deviations.

GN229, using contact-dependent hemolysis of sRBCs. The data showed that only the plasmid harboring *icmTSRQPO* complemented the GN229 phenotype (Fig. 2). Interestingly, the other four *rib* mutants were also complemented by the same *icmTSRQPO* locus. The data showed that the *rib* mutants' genetic lesion was within *icmTSRQPO*.

**The *icmT* gene is essential for the pore-forming activity.** In order to determine which gene(s) was defective in the *rib* mutants, two approaches were adopted. The first approach was to clone various regions of the *icmTSRQPO* and test them for complementation of the *rib* mutants (Fig. 3). As *icmPO* has homologies to proteins involved in the DNA transfer by conjugation and essential for intracellular replication (33), the

*icmTSRQ* region was cloned first from AA100 and tested for complementation of the GN229 mutant. The contact-dependent hemolysis assay (Fig. 4) showed that pTSRQ complemented the *rib* mutant GN229. The plasmid pTS, obtained by deletion of *icmRQ* from the plasmid pTSRQ, and the plasmids pT and pS, obtained by in-frame deletions in *icmS* and *icmT*, respectively, were tested for complementing the defect in the GN229 mutant. In addition, the plasmids pR and pQ, which contain *icmR* and *icmQ*, respectively, were also tested for complementation. The data showed that only *icmTS* and *icmT* complemented the defect of GN229 in the pore-forming toxin (Fig. 4). Plasmids harboring *icmS*, *icmR*, or *icmQ*, failed to complement GN229.

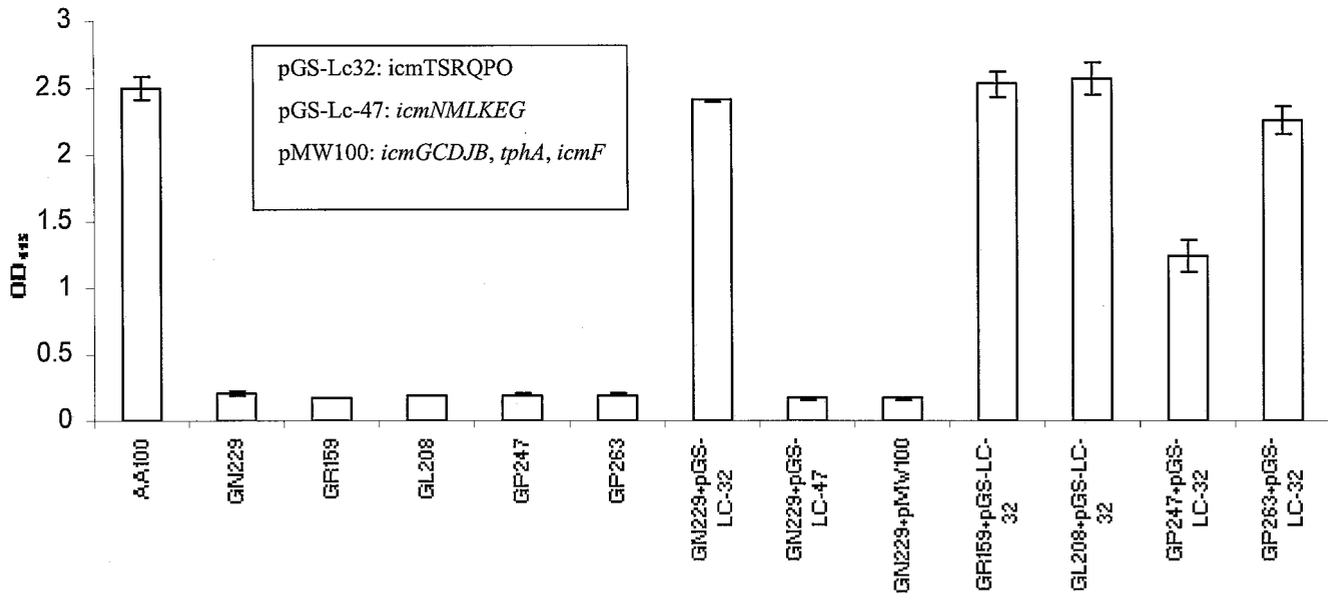


FIG. 2. The *icmTSRQPO*-containing plasmid (pGS-Lc32) complements the *rib* mutants. Contact-dependent hemolysis of sRBC was performed at MOI of 25 for 2 h at 37°C and measured by OD<sub>415</sub> for the release of hemoglobin. The genes harbored by each plasmid are indicated in the box insert. These data are representative of three different experiments, each done in triplicate, and error bars represent standard deviations.

The second approach was to disrupt the individual genes within pTSRQ by transposon mutagenesis to create new plasmids for complementation and also to facilitate construction of *L. pneumophila* AA100 isogenic mutants within this region (Table 1). The *rib* mutant GN229 was transformed with the pTSRQ-derived plasmids, p<sub>kan</sub>T, p<sub>kan</sub>S, p<sub>kan</sub>R, and p<sub>kan</sub>Q (Kan indicates insertion within the gene), and the hemolysis assay was performed to test for complementation for the pore-

forming toxin (Fig. 4). The mutation in GN229 was complemented by p<sub>kan</sub>S, p<sub>kan</sub>R, and p<sub>kan</sub>Q but was not complemented by p<sub>kan</sub>T.

To further confirm the role of *icmT*, we performed allelic exchange of p<sub>kan</sub>T into the chromosome of the wild-type strain AA100 to create an isogenic insertion mutant (AA100kmT). The AA100kmT mutant was defective for the pore-forming toxin and was complemented with *icmTS* and *icmT*, similar to

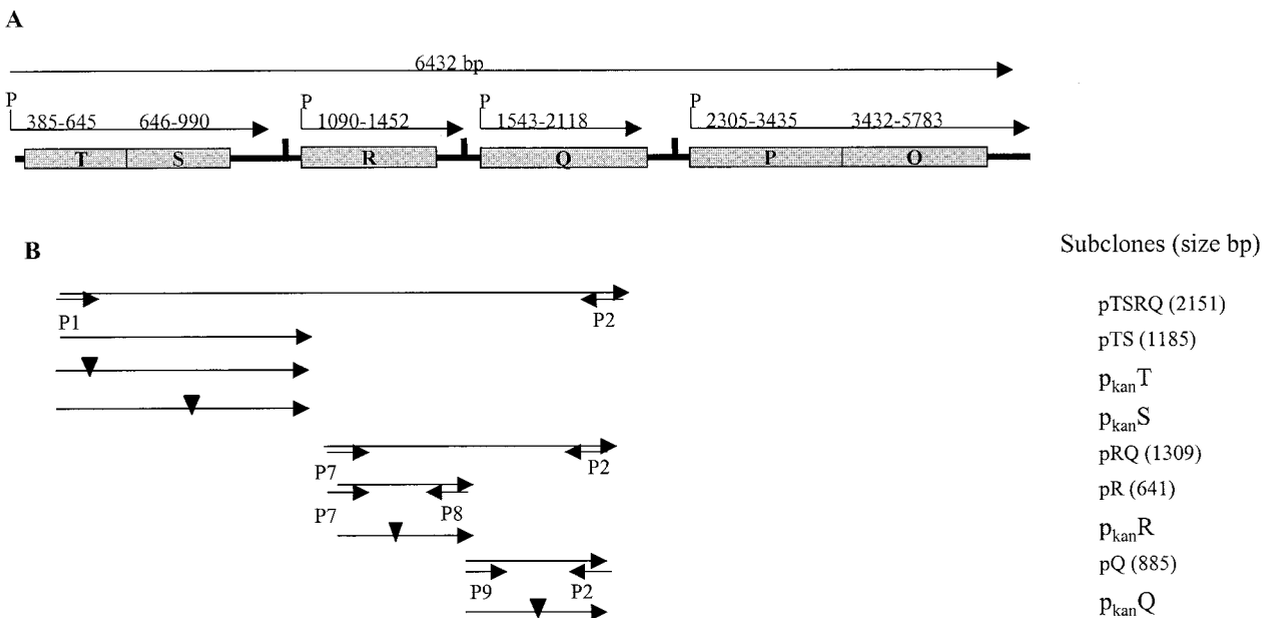


FIG. 3. Genetic organization of the *icmTSRQPO* locus (EMBL accession no. Y12705). (A) Size and predicted transcriptional units of the *icmTSRQPO* locus are indicated by the arrows, while the numbers above the arrows indicate the nucleotide range of the sequence. (B) The genetic regions contained in several subclones and the primers used for their PCR amplification are shown. Inverted filled triangles indicate Kan insertions, and the small arrows indicate the primers used for PCR.

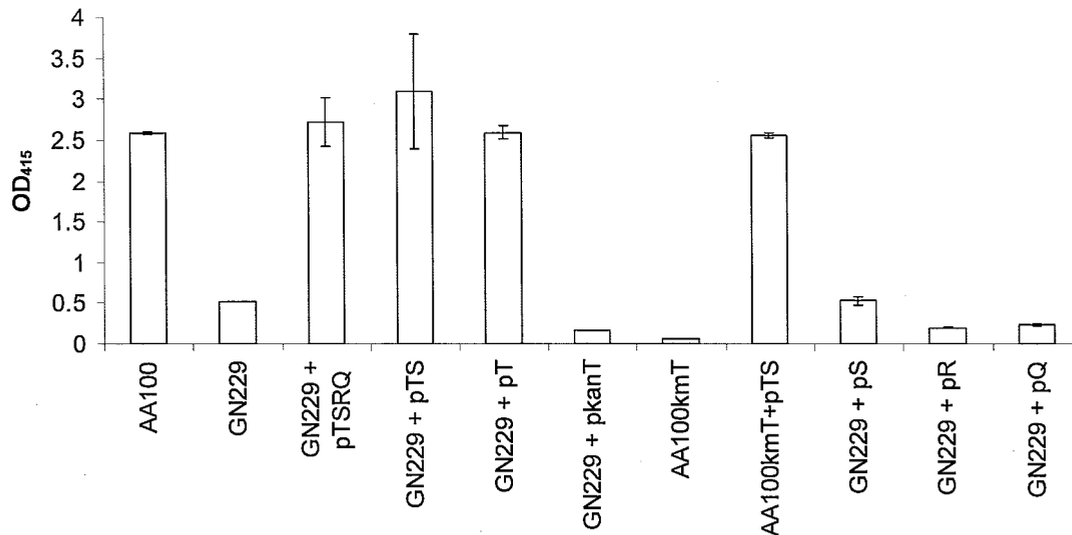


FIG. 4. The *icmT* gene is essential for pore-forming activity. Contact-dependent hemolysis of sRBCs was performed using the strain GN229 or AA100kmT transformed by plasmids carrying different subclones of *icmTSRQ* or by plasmids harboring Kan insertions in *icmT*. In each experiment, AA100 and GN229 were used as positive and negative controls, respectively. These data are representative of three different experiments, each done in triplicate, and error bars represent standard deviations.

GN229 (Fig. 4). Taken together, the data clearly showed that *icmT* was defective in GN229 and was essential for the pore-forming toxin.

**Genetic lesion within *icmT* of the *rib* mutants.** To ascertain the fact that the *rib* mutant phenotype was the result of a mutation in *icmT*, the *icmTS* region of the wild-type strain AA100 and of the five *rib* mutants was sequenced. The results showed that the only difference in the sequence between the *rib* mutants and the wild-type strain was located in the *icmT* gene (Fig. 5). All five *rib* mutants had an identical deletion of a single nucleotide (G) at position 544. This deletion in *icmT* occurred after a stretch of poly(T) regions and is predicted to result in the expression of a truncated protein of 54 amino acids instead of a native protein of 86 amino acids. Since all five *rib* mutants were complemented for the defect in the pore-forming toxin by *icmT* alone, this point deletion in *icmT* of the *rib* mutants is responsible for their defect in the pore-forming toxin-mediated lysis and egress from the host cells.

**Genetic bases of reversion of the *rib* mutants phenotype.** The first section of our results showed that our trial to isolate cosmid clones for complementation of two of the *rib* mutants (GN229 and GL208) for the defective egress from *A. polyphaga* resulted in spontaneous revertants that had no cosmids but regained the wild-type phenotype. This prompted us to analyze the *icmTS* region in the revertants. Sequencing of the *icmTS* region in three GN229 revertants (Fig. 5) showed an insertion of a T at nucleotide 544 in *icmT*, which is the exact location of the point deletion in the *rib* mutants. Although the insertion was a T in the revertants and not a G as in AA100, this point insertion resulted in a codon for the same amino acid as the wild-type strain (GGG for wild type and TGG for revertants; both encode glycine).

**The *icmT* gene is essential for cytolysis of macrophages and *A. polyphaga* by *L. pneumophila* and subsequent bacterial egress.** Cytotoxicity assays were carried out in U937 macrophages and *A. polyphaga* to examine the *rib* mutants and the

complemented strains for cytotoxicity to the host cells. The Alamar Blue assays, carried out after 72 h of infection at an MOI of 1 (Fig. 6) showed that the *rib* mutant GN229 or AA100kmT had minimal cytotoxicity compared to the wild-type strain. The microscopic observation of the infected monolayers showed that the bacteria were trapped in the cells and failed to lyse and egress from the host cells (data not shown), consistent with our previous observations (6). The GN229 and AA100kmT complemented by *icmTS* or *icmT* showed a phenotype similar to that of the wild-type strain in cytolysis and egress from the host cell.

We have previously shown that the *rib* mutants are defective in egress from the host cell after termination of intracellular replication but are subsequently released 24 to 48 h later, which is most likely due to apoptosis of the infected cell (6). The kinetics of bacterial egress from U937 macrophages was performed at an MOI of 1 to test the ability of the mutants and the complemented strains to egress from the host cell after intracellular growth. As expected, there was an ~100-fold decrease in the egress of GN229 ( $10^5$  CFU/ml) compared to that



FIG. 5. The mutation of the five *rib* mutants and their revertants is located in *icmT*. The sequences of *icmT* of the five *rib* mutants and three of the *rib* revertants are aligned. *L. pneumophila* (strain AA100) from our laboratory and that from the sequence deposited by Segal and Shuman (34), designated here lp(db), were also included.

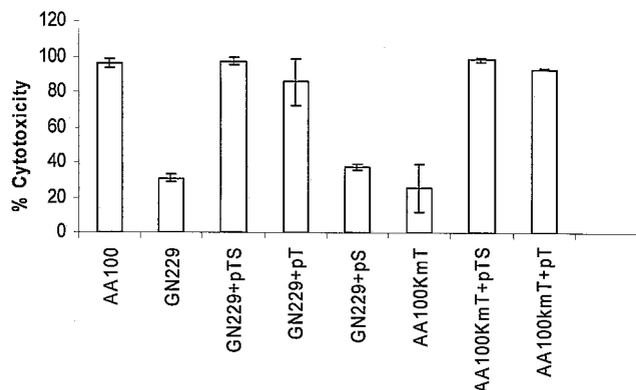


FIG. 6. The *icmT* gene of *L. pneumophila* is essential for cytotoxicity to U937 macrophages. Cytotoxicity to U937 macrophages infected at an MOI of 1 by *L. pneumophila* strains was measured at 72 h postinfection for the reduction of the Alamar Blue dye, and results were compared to those for noninfected cells. These data are representative of three different experiments, each done in triplicate, and error bars represent standard deviations.

of AA100 ( $10^7$  CFU/ml) at 24 h. At 48 h postinfection, there was a 500-fold reduction in egress of GN229 compared to the parental strain (Fig. 7). Importantly, GN229 complemented by *icmTS* or *icmT* egressed from U937 macrophages similar to AA100, while GN229 complemented with *icmS* was similar to the GN229 mutant (Fig. 7). These data showed that *icmT* was essential for egress from macrophages upon termination of intracellular replication.

The infection of *A. polyphaga* by the wild-type strain, the

mutants, and the complemented strains (Fig. 8) showed that GN229 and AA100KmT were defective in cytolysis of the host cell, in sharp contrast to the wild-type and the complemented strains. As with macrophages, microscopic observation of cells infected by the *rib* mutant GN229 and AA100KmT showed that the mutants failed to lyse the protozoan cells (Fig. 8). The defect in egress of the mutants was complemented by plasmids carrying *icmTSRQPO* (data not shown), *icmTSRQ*, *icmTS*, and *icmT*. We concluded that the *icmT* gene was essential for cytotoxicity and lysis of both human and protozoan cells by *L. pneumophila*.

**The *rtxA* gene is not involved in pore-forming toxin-mediated egress from macrophages and *A. polyphaga*.** The *rtxA* gene has been shown to be involved in entry into macrophages and is thought to have a minor role in pore formation when tested for enhanced permeability of the plasma membrane to ethidium bromide (13, 14). Since it was proposed that RtxA might play a role in pore formation, we examined its role in pore-forming toxin-mediated egress from macrophages and protozoa. The role of this gene in egress, contact-dependent hemolysis, and cytotoxicity to macrophages and amoebae was examined using an isogenic mutant obtained by an in-frame deletion in *rtxA* ( $\psi$ lp24) (13, 14) and compared to the wild-type strain AA100. Amplification by PCR of the *rtxA* gene confirmed the deletion of *rtxA* within  $\psi$ lp24 (data not shown). In all assays (Fig. 9 and data not shown), the  $\psi$ lp24 mutant had an indistinguishable phenotype from that of AA100. Thus, *rtxA* was not required for cytolysis or for pore formation-mediated egress of *Legionella* from mammalian and protozoan cells.

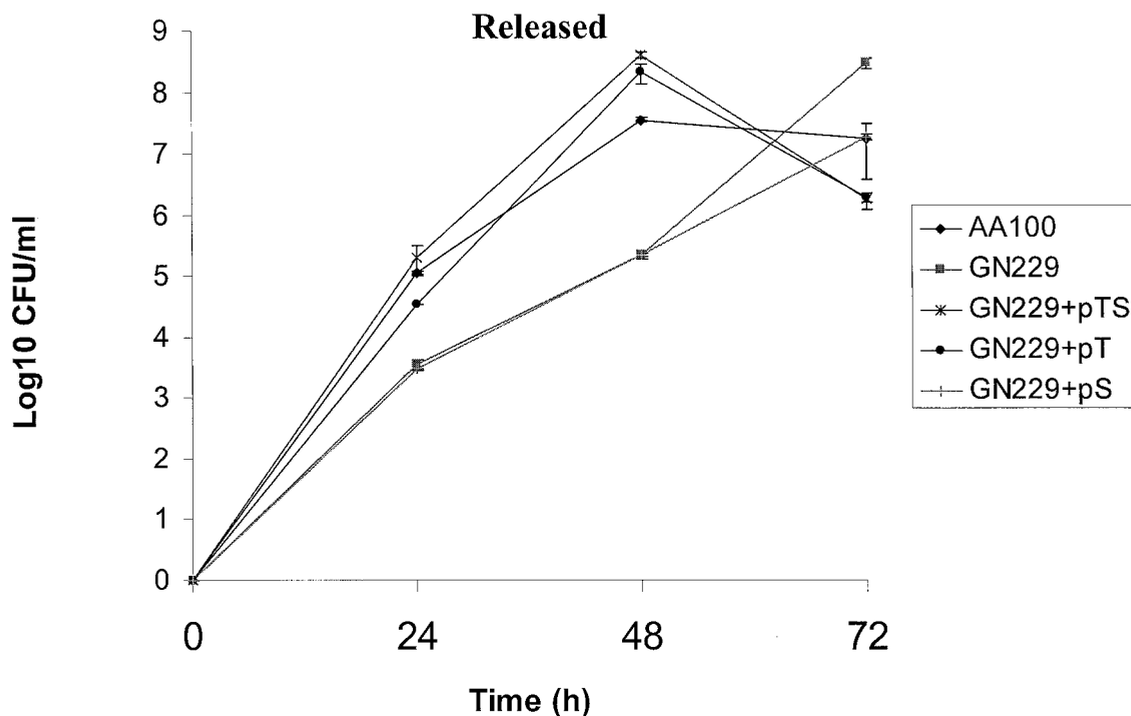


FIG. 7. The *icmT* gene is essential for bacterial egress from macrophages. U937 macrophages were infected by strains of *L. pneumophila* at an MOI of 1 for 1 h followed by gentamicin treatment for 1 h to kill extracellular bacteria. The bacteria released from the supernatants were plated for colony enumeration. These data are representative of three different experiments, each done in triplicate, and error bars represent standard deviations.

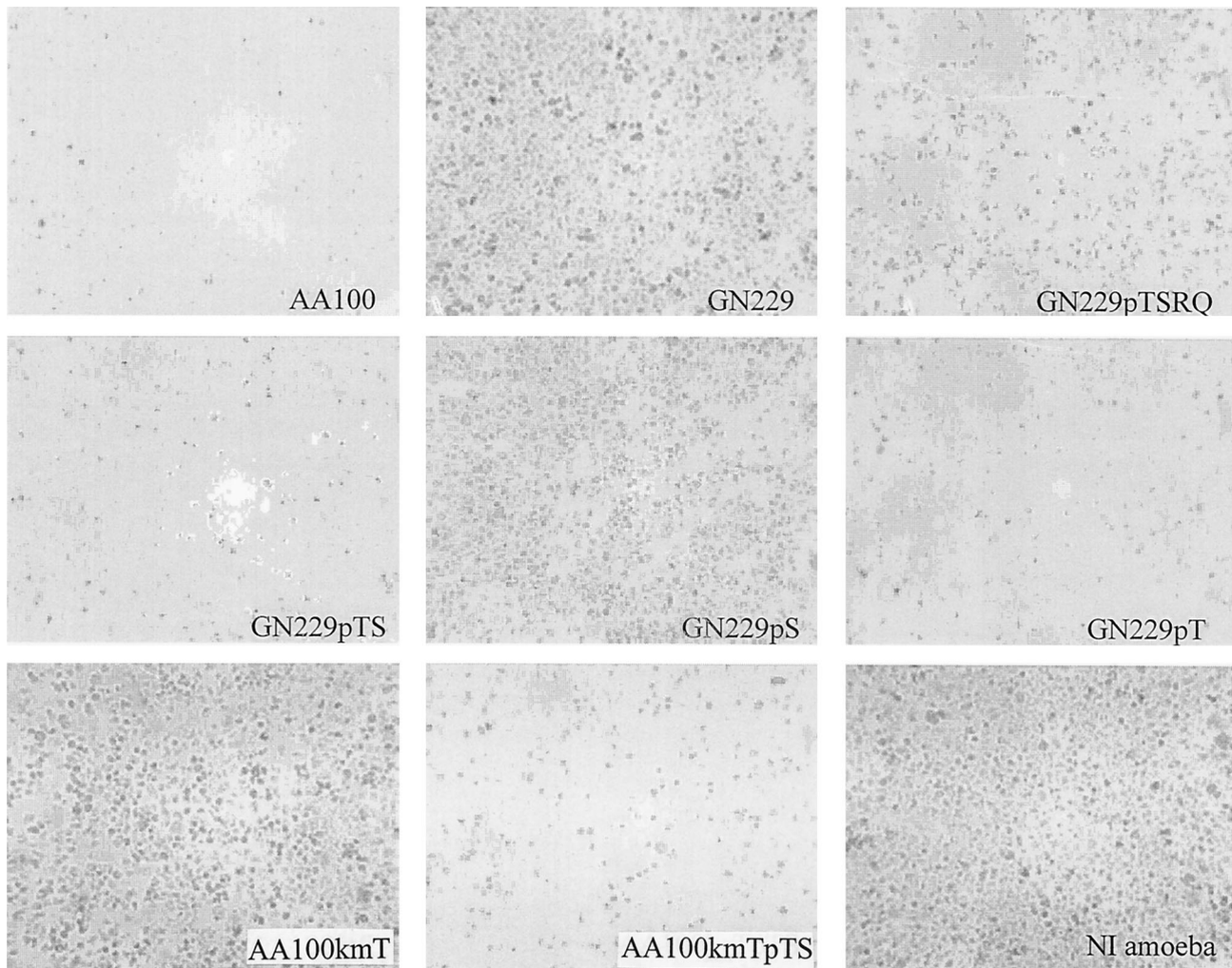


FIG. 8. The *icmT* gene of *L. pneumophila* is essential for cytolysis of *A. polyphaga*. Shown are representative phase-contrast images of *A. polyphaga* infected by *L. pneumophila* AA100, GN229, and GN229 harboring different plasmids at an MOI of 10 and examined at 96 h postinfection. NI, noninfected *A. polyphaga*.

## DISCUSSION

We have shown recently that the five *rib* mutants replicate intracellularly similar to the parental strain but are defective in pore formation-mediated cytolysis and egress from the host cell upon termination of intracellular replication (6, 17). The data show that *icmTS* and *icmT* all complement the *rib* mutants for the defect in pore formation-mediated cytolysis and egress from host cells upon termination of intracellular replication. Since *icmS* is required for intracellular growth of *L. pneumophila* in macrophages (15) and does not complement the *rib* mutants, our studies have been focused on *icmT*.

Although the defective phenotype of the *rib* mutants is due to a spontaneous point deletion in *icmT*, an insertion mutation in *icmT* has a similar phenotype to the *rib* mutants, in the defect in pore formation-mediated egress upon termination of intracellular replication. Importantly, since this insertion mutant is also complemented by *icmT*, the kanamycin insertion does not have a polar effect on the expression of the downstream *icmS*. This is rather surprising, since the termination codon of *icmT* is followed immediately downstream with the

initiation codon of *IcmS*. Our observations suggest that *icmS* may be regulated separately by its own promoter. However, promoter and transcriptional studies would be required to determine this speculation.

Complementation of the pore-forming activity and its role in cytolysis and egress from both mammalian and protozoan cells by *icmTSRQPO* are only partial (34), in contrast to the complete complementation by *icmTSRQ*, *icmTS*, and *icmT*, all of which are cloned in the pBC plasmid. The pGS-Lc-32 plasmid harboring *icmTSRQPO* contains the mobilization region, *mob*, which is essential for conjugation. It has been previously shown that the presence of the mobilization genes on a complementing plasmid is also responsible for the low level of complementation of other *dot/icm* mutants (35). It is thought that the Mob proteins compete with export of bacterial effectors required for cytotoxicity and intracellular multiplication (35). Thus, the partial complementation by the pGS-Lc-32 plasmid is most likely due to the interference of the Mob proteins in export of the pore-forming toxin.

The *rib* mutants are spontaneous (6), and our trials to com-

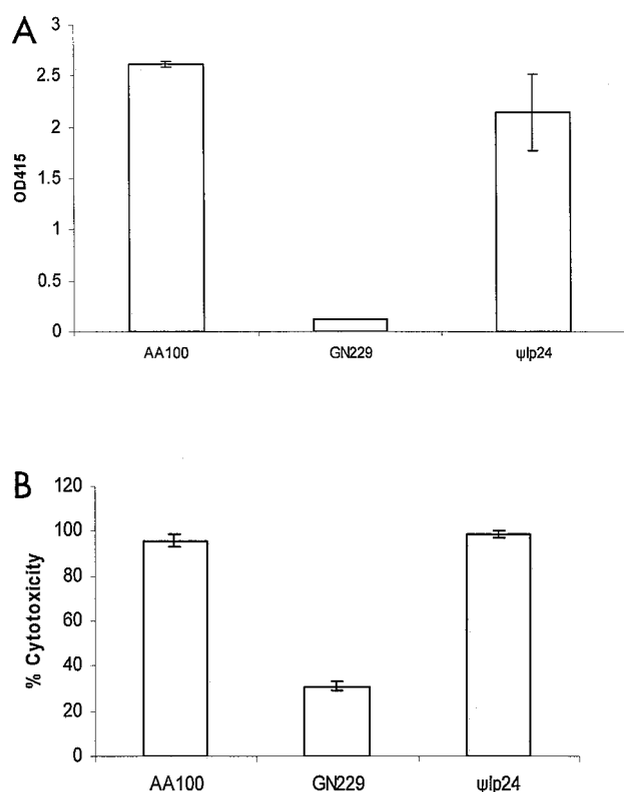


FIG. 9. The *rtxA* gene is not required for pore formation-mediated hemolysis of sRBCs and for cytotoxicity to U937 macrophages. (A) Contact-dependent hemolysis assays of sRBCs; (B) cytotoxicity assays to U937 macrophages. Strains tested are the wild-type strain AA100, GN229, and  $\psi$ lp24 (in-frame *rtxA* deletion mutant). These data are representative of three different experiments, each done in triplicate, and error bars represent standard deviations.

plement them for egress from *A. polyphaga* with the cosmid library have also resulted in isolation of spontaneous revertants that regained the wild-type phenotype but do not contain cosmids. Interestingly, two of the mutants (without being transformed by the cosmid library) that have been used as controls in the amoebal enrichment do not revert to the wild-type phenotype. The spontaneous revertants do not have the wild-type sequence of *icmT*, suggesting that there has been no recombination with the cosmid harboring the *icmT* region. However, it is not clear how harboring the cosmid library would enhance the spontaneous reversion process, which occurred in the two tested mutants. Sequencing of the *icmT* gene of the five *rib* mutant strains and three of the *rib* revertants has shown that a single nucleotide deletion or insertion, respectively, is responsible for the phase variation in the pore-forming activity. This deletion or insertion in the *rib* mutants and their revertants, respectively, occurs after a poly(T) stretch and is most likely due to slipped-strand mispairing during DNA replication. This mode of phase variation has been described for virulence genes of many pathogens such as the *pilC* of *Neisseria gonorrhoeae*, in which a slipped-strand mispairing after a stretch of G nucleotides results in a frameshift mutation (24). It would be interesting to determine the rate of phase variation of *icmT* in vitro and intracellularly. It would be also interesting to examine phase variation at this locus and its role

in the defect in the pore-forming toxin of other *Legionella* species, such as *L. micdadei* (8), that have been shown to have the *dot/icm* loci (25).

In addition to its essential role in pore formation-mediated lysis of the host cell, IcmT is also involved in the DNA conjugation (35). The IcmT protein sequence is similar to the TraK protein encoded by the transfer region of IncI plasmid (27). Although our data clearly show an essential role for IcmT in pore formation-mediated egress from the host cell upon termination of intracellular replication, it is not known how IcmT is involved in this process at the molecular and biochemical level. It is unlikely that IcmT is a common regulator of several virulence traits that are induced upon termination of intracellular replication (10), including the pore-forming toxin (6), evasion of phagolysosomal fusion, and motility, since the *rib* mutant evades phagolysosomal fusion and is motile (6). It is possible that IcmT is the structural toxin or is a cofactor that is specifically required for export of the pore-forming toxin, as a cofactor or a chaperon. Future studies are designed to determine whether IcmT is the pore-forming toxin or is a cofactor required for export and/or activity of the pore-forming toxin.

Whether the Lsp type II and the Dot/Icm type IV secretion systems of *L. pneumophila* interact in export of effector molecules through the cytoplasmic and outer membrane, respectively, is not known. The pertussis toxin of *Bordetella pertussis* is exported through the type II secretion systems into the periplasm, where the holotoxin is assembled and is subsequently exported by the type IV secretion system through the outer membrane (11). We have shown that mutants of *L. pneumophila* defective in the type II secretion system are not defective in the pore-forming toxin. Therefore, in contrast to the pertussis toxin, export of the pore-forming toxin is independent of the type II secretion system. We have also recently shown that export of the effector molecule responsible for the induction of apoptosis occurs through the Dot/Icm system and its export is also independent of the type II secretion system (Zink et al., submitted for publication). Since the type II secretion system is not required for intracellular replication of *L. pneumophila* within mammalian cells (20), it is most likely that it is also not required for export of effector molecules responsible for modulation of the phagosome to evade maturation along the endosomal-lysosomal degradation pathway. Therefore, export of many virulence factors through the Dot/Icm type IV secretion system is independent of the type II secretion system.

The RtxA-like protein of *L. pneumophila* (repeats in structural toxin) has been shown to be involved in entry into macrophages and to play a minor role in pore formation when examined by the increase in permeability to ethidium (13). We have shown that RtxA plays no role in pore formation when detected by hemolysis of RBCs. We propose that there are at least two pores of different sizes generated by *L. pneumophila* in host cell membranes. One pore can be detected by the increase in permeability of the host cell membrane to ethidium bromide and is mediated by RtxA upon bacterial entry. The second and larger pore is generated by the pore-forming toxin that is required for bacterial egress and is detected by hemolysis of RBCs. This hypothesis is supported by our data, which show that *rtxA* is not required for pore formation-mediated bacterial egress and that the *rtxA* mutant is not impaired for

pore formation-mediated hemolysis of RBCs. However, the *rtxA* mutant is partially defective in causing an increased permeability of the host cell membrane to ethidium bromide (13). Kirby et al. have shown that polyethylene glycol (PEG) 3350 (average molecular weight, 3350) prevents blebbing of macrophages infected by *L. pneumophila*, whereas PEG 1000 does not. Interestingly, PEG 3350 does not protect infected cells from becoming permeable to ethidium bromide (26). It is likely that the pore used for bacterial entry is the pore detected by increased permeability of the host cell to ethidium bromide, and RtxA may partially play a role in this process. This smaller pore is likely to be distinct from the larger pore utilized for cytolysis and bacterial egress from the host cell and is detected by hemolysis of RBCs.

In summary, we have shown that the *rib* mutants, which replicate intracellularly but are defective in pore formation-mediated egress from the host cell, are defective in *icmT* due to a point deletion after a stretch of poly(T) in the sequence encoding the C terminus of the protein. We have also shown that the type II secretion system of *L. pneumophila* is not involved in export of the pore-forming toxin.

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