

Mutational Analysis of the *Chlamydia muridarum* Plasticity Zone

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Pathogenically diverse *Chlamydia* spp. can have surprisingly similar genomes. *Chlamydia trachomatis* isolates that cause trachoma, sexually transmitted genital tract infections (chlamydia), and invasive lymphogranuloma venereum (LGV) and the murine strain *Chlamydia muridarum* share 99% of their gene content. A region of high genomic diversity between *Chlamydia* spp. termed the plasticity zone (PZ) may encode niche-specific virulence determinants that dictate pathogenic diversity. We hypothesized that PZ genes might mediate the greater virulence and gamma interferon (IFN- γ) resistance of *C. muridarum* compared to *C. trachomatis* in the murine genital tract. To test this hypothesis, we isolated and characterized a series of *C. muridarum* PZ nonsense mutants. Strains with nonsense mutations in chlamydial cytotoxins, *guaBA-add*, and a phospholipase D homolog developed normally in cell culture. Two of the cytotoxin mutants were less cytotoxic than the wild type, suggesting that the cytotoxins may be functional. However, none of the PZ nonsense mutants exhibited increased IFN- γ sensitivity in cell culture or were profoundly attenuated in a murine genital tract infection model. Our results suggest that *C. muridarum* PZ genes are transcribed—and some may produce functional proteins—but are dispensable for infection of the murine genital tract.

Acquisition of genetic islands encoding the machinery necessary to invade eukaryotic cells was a key event in the evolution of many bacterial pathogens from free-living ancestors. For those pathogens that subsequently lost the ability to replicate outside host cells, a dearth of foreign microbial genetic material and Muller's ratchet may have tilted the balance of niche adaptation toward genomic decay. Reductive evolution appears to have played an especially pivotal role in niche adaptation of the obligate intracellular bacterial pathogens in the family *Chlamydiaceae* (1). Understanding why extant *Chlamydia* spp. retained different combinations of the metabolic pathways encoded by their common ancestors could reveal insights into chlamydial pathogenesis and cognate immune defenses against these intracellular pathogens.

Whole-genome comparisons provided early clues concerning the importance of genomic streamlining in chlamydial evolution but less insight into how loss of specific genes and metabolic pathways is adaptive (2–5). What is clear is that the genomes of pathogenically diverse *Chlamydia* spp. are small, have limited biosynthetic capacity, and are remarkably similar. For example, the murine pathogen *Chlamydia muridarum* and pathogenically diverse *Chlamydia trachomatis* isolates share 99% of their gene content (4). Various hypotheses have been put forth to explain this paradox, which was recently described as “divergence without difference” (6). One idea is that polymorphisms in conserved genes are key chlamydial virulence determinants. Some *C. trachomatis* trachoma strains have no obvious novel genes that differentiate them from genital strains (7), and some trachoma strains that vary in sensitivity to gamma interferon (IFN- γ) differ by only a few polymorphisms in conserved “core” genes (8). Alternatively, but not exclusively, open reading frames (ORFs) that have accumulated in a variable genomic region termed the plasticity zone (PZ) may be determinants of chlamydial pathogenic diversity (4). Comparative genomic and biochemical studies have implied that a partial tryptophan operon in the PZs of some *C. trachomatis*

strains could reflect niche adaptation to immune-regulated tryptophan availability (9–12). Functionality of chlamydial tryptophan synthase, the product of this operon, in cell culture was recently confirmed using a reverse genetic strategy (13). Whether other PZ ORFs similarly mediate chlamydial niche adaptation or the PZ primarily houses inactive and decaying genes remains unclear.

The PZ is consistently located near the region of replication termination in *Chlamydia* spp. but varies considerably in size and gene content (4). Some PZ ORFs have no homologs outside *Chlamydia*, whereas others have been grouped into families based upon the homology of their predicted protein products to characterized bacterial and mammalian proteins. Some *Chlamydia* spp. encode a protein that resembles eukaryotic membrane attack complex (MAC) proteins and perforin (MACPF) in the PZ (4, 14). Various functions have been ascribed to chlamydial MACPFs, including immune evasion by molecular mimicry, protein secretion

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tion, and bacterial entry/exit (14, 15). Chlamydial MACPF is also purported to assist in lipid modification because of its proximity to a cluster of phospholipase D (PLD)-like genes (14, 16). Indirect evidence also suggests that PZ PLDs, like their mammalian homologs, are sensitive to primary alcohols and play roles in lipid acquisition or transfer (17, 18). The *C. muridarum* PZ contains three putative cytotoxins that have similarity to the large clostridial toxins (LCTs) and yersinial YopT (4). The cytotoxin ORFs of *C. trachomatis* serovars are truncated, disrupted, or absent altogether (2, 19, 20). A *guaBA-add* operon found in *C. muridarum*, but not *C. trachomatis*, could render the mouse pathogen less dependent on host nucleotides (4). These purine interconversion genes are replaced by *trpRBA* in *C. trachomatis* (2, 4). Limitations of available animal models and a historic lack of genetic tools have prevented the testing of hypotheses concerning the functions of PZ ORFs in chlamydial virulence and tropism. However, recent breakthroughs in genetic manipulation of *Chlamydia* now make it possible to address some of these questions directly (13, 21, 22).

Determining the functions of PZ ORFs could provide insight into the molecular basis of chlamydial host and tissue tropism and direct the design of improved animal models of human chlamydial disease. *C. muridarum* infection of the murine genital tract (GT) is a leading model for the study of adaptive immunity to genital infection because innate immunity is sufficient to clear murine genital infection with human strains of *C. trachomatis* (23). We and others have speculated that *C. muridarum* PZ genes play roles analogous to that of *C. trachomatis* tryptophan synthase in the circumvention of niche-specific innate immune responses (4, 24). If correct, this also implies that specific effectors were retained by *C. muridarum* because the cognate targets are relevant barriers during the natural course of infection. Murine modeling of *C. trachomatis* human urogenital tract infection might be improved by use of *C. muridarum* PZ-null mutants and mice lacking the corresponding targets of these effectors (25). Alternatively, if PZ ORFs are dispensable for GT infection, it would suggest either that these ORFs do not encode functional proteins or that the appropriate targets of the proteins are not relevant during experimental GT infection.

In this study, we used a combination of transcriptional profiling, reverse genetic analysis, and mouse modeling to assess if *C. muridarum* PZ ORFs are expressed and necessary for murine GT infection. Our results suggest that although PZ ORFs are transcribed, they are dispensable for infection of the murine GT.

MATERIALS AND METHODS

Cell lines, chlamydial culture, and infection. *C. muridarum* and *C. trachomatis* serovar L2 strain 434/Bu (*C. trachomatis*) were kind gifts from Harlan Caldwell (Rocky Mountain Laboratories, NIAID, Hamilton, MT). *Chlamydia* organisms were routinely propagated in McCoy mouse fibroblast cells (American Type Culture Collection CRL-1696), and infectious elementary bodies (EB) were purified by density gradient centrifugation (26). McCoy cells and HeLa human cervical epithelial cells (American Type Culture Collection CCL-2) were maintained in Dulbecco's modified Eagle's medium (DMEM) containing high-glucose HEPES, glutamine, and sodium pyruvate (HyClone) and supplemented with 10% fetal bovine serum albumin (Atlanta Biologicals) and 100 μ M nonessential amino acids (Gibco). Infections were performed in tissue culture flasks or plates with appropriate dilutions of chlamydial stock in SPG buffer (0.25 M sucrose, 10 mmol/liter phosphate, 5 mmol/liter L-glutamic acid, pH 7.2). The infected flasks were rocked for 2 h at 37°C, whereas infected plates were centrifuged at room temperature for 1 h, followed by rocking

at 37°C for 30 min. During large-scale propagation experiments, SPG was replaced with fresh DMEM containing 0.5 μ g/ml cycloheximide. In some experiments, DMEM containing 20 U/ml murine IFN- γ (R&D Systems) was added to McCoy cells 24 h prior to infection and postinfection. Cycloheximide was excluded from the cell culture media in experiments where IFN- γ was added.

Recoverable IFU (rIFU) assays. McCoy cells in 24-well plates were infected with *C. muridarum* or PZ mutants at a multiplicity of infection (MOI) of 0.25. The infected cell monolayers were frozen in 500 μ l SPG at various intervals postinfection (p.i.). Upon thawing, cells were scraped using a pipette tip and EB were harvested by bead beating the cell suspension in microcentrifuge tubes. Inclusion-forming unit (IFU) assays were performed using serial dilutions of the harvests to infect confluent McCoy cell monolayers in 96-well plates. Cells were fixed with methanol and stained with anti-chlamydial lipopolysaccharide (LPS) monoclonal antibody (MAb) (EVIH1), followed by a secondary Alexa Fluor 488-conjugated anti-mouse IgG MAb (Life Technologies) at 24 h p.i. Chlamydial inclusions were imaged and enumerated using an EVOS FL Auto cell-imaging system (Life Technologies).

RT-PCR and qRT-PCR. McCoy cells in six-well plates were infected with *C. muridarum* at an MOI of 1. Total RNA was extracted at 0, 6, 12, 18, 24, and 30 h p.i. using TriSure reagent (Bioline), treated with DNase I (Qiagen) for 15 min, and then purified on GeneJet RNA purification columns (Thermo Scientific). Reverse transcriptase (RT) PCRs were performed using 100 ng of total RNA and gene-specific primers (see Table S1 in the supplemental material) or primer pairs spanning intergenic regions (see Table S3 in the supplemental material) using the MyTaq one-step RT-PCR kit (Bioline). For quantitative RT-PCR (qRT-PCR) experiments, total RNA was converted to cDNA using the Maxima H Minus first-strand cDNA synthesis kit (Thermo Scientific). The qRT-PCRs were performed in an Eppendorf Realplex4 thermocycler using SensiFast SYBR No-Rox master mix (Bioline) with gene-specific primers (see Table S2 in the supplemental material). The amplification cycle included a 2-min step at 95°C, followed by 40 cycles of 95°C for 5 s, 60°C for 10 s, and 72°C for 20 s. Absolute transcript copy numbers were calculated by normalizing to standard curves of *C. muridarum* genomic DNA amplified in parallel with experimental samples. Control reaction mixtures lacking RT were run in parallel with all RT-PCR and qRT-PCR experiments.

PZ cloning and β -galactosidase assays. Overlapping 500- to 1,000-bp regions of the *C. muridarum* PZ were PCR amplified from *C. muridarum* genomic DNA and were cloned upstream of the *lacZ* gene in pAC-lacZ between the NruI and EagI sites of the vector (27). The sequences of the PCR primers and the genomic coordinates of the cloned PZ regions are listed in Table S4 in the supplemental material. The resulting plasmids were transformed into *Escherichia coli* DH5 α , and the inserts and cloning junctions were confirmed by DNA sequencing. For β -galactosidase activity assays, triplicate samples from overnight cultures of *E. coli* strains carrying various PZ and control plasmids were grown in Luria-Bertani broth containing 50 μ g/ml chloramphenicol and then diluted 1:100 in fresh broth and grown at 37°C until they reached mid-log phase (optical density at 600 nm [OD₆₀₀] \approx 0.6). β -Galactosidase expression analysis was performed as described previously (28, 29) with the following modifications. Cells were diluted in a total volume of 500 μ l Z buffer (60 mM Na₂HPO₄, 40 mM NaH₂PO₄ · H₂O, 10 mM KCl, 1 mM MgSO₄ · 7H₂O) followed by the addition of 50 μ l of chloroform and 50 μ l of 0.1% sodium dodecyl sulfate. The solution was vortexed for 5 s, and the reactions were equilibrated by incubation at room temperature for 5 min prior to the addition of 100 μ l of *ortho*-nitrophenyl- β -galactoside (4 mg/ml). The reactions were stopped by adding 250 μ l of 1 M Na₂CO₃ and then centrifuged at 14,000 rpm for 5 min in an Eppendorf microcentrifuge to separate the cell pellet and the supernatant. The initial OD₆₀₀ of the cells and the OD₄₂₀ of the supernatant were measured and used to calculate β -galactosidase activity.

Preparation of CEL1 endonuclease extract. CEL1 extract was prepared from celery as described previously (30). Fresh celery stalks (25 kg)

were washed, chopped to remove the leaves, and juiced to yield approximately 10 liters of celery juice. Subsequent steps were carried out at 4°C. Five hundred milliliters of buffer A containing 2 M Tris-HCl (pH 7.7) and 0.1 M phenylmethanesulfonyl fluoride (PMSF) was added to the celery juice, and the solution was centrifuged to remove plant debris. Protein was precipitated from the supernatant by ammonium sulfate fractionation in the 25 to 80% saturation range. The protein pellet was resuspended in 100 ml buffer B (0.1 M Tris-HCl, pH 7.7, 0.5 M KCl, 100 μ M PMSF, 0.01% Triton X-100) and dialyzed overnight against a large volume of buffer B. The final CEL1 extract was stored in small aliquots at -80°C.

EMS mutagenesis and library construction. Confluent McCoy cells in a T-75 flask were infected with *C. muridarum* at an MOI of 0.5. At 16 h p.i., the culture medium was replaced with 5 ml of DMEM containing 8 μ g/ml ethyl methanesulfonate (EMS) (Sigma). The cells were exposed to EMS for 1 h and then washed several times with 1 \times phosphate-buffered saline (PBS), after which fresh DMEM without mutagen was added. EB were bead harvested from the cells at 28 h p.i. and resuspended in 2 ml of SPG. An IFU assay estimated chlamydial survival after EMS treatment at less than 0.1% that of the untreated control. Half of the harvest was stored at -80°C, and the remainder was used to infect a fresh T-75 flask of cells. Four rounds of EMS mutagenesis were performed in total, and the final harvest yielded 9×10^5 EB. These EB were divided into small aliquots and stored at -80°C. To construct mutant libraries, 96-well plates of McCoy cells were infected with 10 EB per well of the mutagenized stock. Mutant chlamydiae were scaled up by four rounds of passaging in 96-well plates. Plates from the third passage were stored at -80°C to serve as the library. Genomic DNA was extracted from the final plates by adding 75 μ l of alkaline lysis buffer to each well (0.2 mM EDTA, 25 mM NaOH, pH 12.0) and incubating at 95°C for 10 min, after which the lysis buffer was neutralized by adding an equal volume of 40 mM Tris-HCl (pH 5.0). The genomic DNA plates were stored at 4°C.

TILLING. Targeting induced local lesions in genomes (TILLING) was used to identify specific mutants in the library and was performed as described previously (13). Specific primers (see Table S6 in the supplemental material) were used to amplify targeted PZ genes from template genomic DNA plates, which were extracted from the mutant libraries. These PCR products were annealed slowly (80°C; 7 s; 60 cycles, decreasing by 0.3°C every cycle) and then were digested with CEL1 endonuclease. Digestions were performed for 10 min at 45°C using an empirically determined volume of CEL1 extract (range, 0.1 to 1 μ l) diluted in 1 \times CEL1 buffer (0.01 M MgSO₄, 0.01 M HEPES, pH 7.5, 0.02 M KCl, 0.005% Triton X-100, 2 μ g/ml bovine serum albumin). The digestions were stopped by adding 10 μ l EDTA (0.15 M, pH 8.0). The digested products were separated on 2% agarose Tris-borate-EDTA (TBE) gels, stained with Midori Green (Bulldog Bio), and visualized using a ChemiDoc XRS+ device (Bio-Rad Laboratories, Inc., Hercules, CA).

Cytotoxicity assays. McCoy cells seeded on coverslips in 24-well plates were infected with *C. trachomatis*, *C. muridarum*, or cytotoxin mutants at an MOI of 10 or 250 in the presence of 0.1 μ g/ml rifampin. The cells were fixed in 3.7% formaldehyde at 3 h p.i. and then permeabilized with 1% Triton X-100. Following staining with Alexa Fluor 488-phalloidin for visualization of actin filaments, the coverslips were mounted on slides using ProLong Gold antifade reagent with DAPI (4',6-diamidino-2-phenylindole) (Life Technologies). The cells were imaged using a Nikon Eclipse Ni-E fluorescence microscope (Nikon Instruments, Melville, NY).

Lactate dehydrogenase (LDH) assays were performed according to the manufacturer's instructions (OPS Diagnostics). McCoy cells were infected with various strains at a range of MOI (10, 25, 100, and 500) in the presence of rifampin. To determine maximal LDH release, cells in control wells were lysed with 10% Triton X-100 10 min prior to supernatant collection. The supernatants were removed from the wells 3 h p.i. after brief centrifugation of the plates to remove debris. Twenty-five microliters of the supernatants was mixed with 75 μ l of the dye-buffer solution. The A₄₉₀ was measured after 15 min incubation at 37°C.

Animal experiments. C57BL/6J mice were obtained from Jackson Laboratories (Bar Harbor, ME). All animal experiments were approved by the Institutional Animal Care and Use Committees of the University of Arkansas for Medical Sciences and Indiana University. Female mice between 6 and 10 weeks old were treated with 2.5 mg medroxyprogesterone (Depo-Provera) diluted in 100 μ l PBS 10 and 3 days prior to infection. Prior to infectious challenge, the vaginal vault was swabbed to remove excess mucus. Mice were inoculated vaginally with *C. muridarum* or PZ mutants at a concentration of 5×10^4 IFU in 5 μ l SPG. The infections were monitored by enumeration of IFU from cervical swabs collected at various time points. At 73 days postinfection, the mice were euthanized and examined visually for hydrosalpinx.

Sequencing library construction and whole-genome sequencing. Gradient-purified EB were treated with DNase I (NEB) at 37°C for 1 h, and then 5 mM EDTA was added and the mixtures were incubated at 75°C for 10 min to halt digestion. Total DNA was extracted using the DNeasy blood and tissue kit (Qiagen), and DNA concentrations were determined using the Quant-It high-sensitivity double-stranded DNA (dsDNA) assay (Life Technologies Corp.). The dsDNA fragments were generated by treating 1 μ g of genomic DNA with NEBNext dsDNA Fragmentase (NEB). DNA sequencing libraries were prepared using the TruSeq Nano DNA sample preparation kit (Illumina Inc.) according to the manufacturer's protocols. Samples were multiplexed using TruSeq Single Index sequencing primers. Equimolar concentrations of the indexed libraries were combined into a single pool and submitted for Illumina HiSeq sequencing at the Tufts University Genomics Core Facility. Paired-end 100-bp sequencing was performed on the Illumina HiSeq 2500 platform.

Genome assembly and sequence analysis. For single nucleotide polymorphisms (SNP), nucleotide insertion and deletion (indel) analysis, raw sequence data were mapped to the corresponding reference genome (GenBank accession number AE002160.2) using Bowtie2 (56). The resulting SAM files were converted to a BAM file format and sorted. SNPs/indels were called using a Samtools mpileup function. The remaining ambiguous sequences from the data set and mutation calls with low-quality scores were resolved by PCR and Sanger sequencing.

Statistics. Data were analyzed using Prism 6.0 software (GraphPad). For comparisons of multiple groups with more than one variable, two-way analysis of variance (ANOVA) with Bonferroni posttest was used. Two-way ANOVA with Bonferroni posttest was also used to analyze the differences in chlamydial burdens between groups of infected mice. Multiple comparisons for data with a single variable were analyzed by one-way ANOVA with Dunnett's posttest. Differences in the development of hydrosalpinx between *C. muridarum*-infected mice and mutant-infected mice were analyzed by Fisher's exact test. Differences were considered statistically significant when the *P* value was <0.05.

RESULTS

The *C. muridarum* plasticity zone is transcriptionally active.

The *C. muridarum* PZ is defined as the 50-kb locus between *tc0431* (MACPF) and *tc0448* (*dsbB*) (Fig. 1). The *C. muridarum* PZ is poorly characterized, and the only evidence for gene activity in the region comes from a prior transcriptional study of the cytotoxin ORFs (*tc0437* to *tc0439*) (19). To determine if and when other PZ ORFs were expressed, we examined the entire locus for transcriptional activity. RT-PCR analysis of RNA isolated from *C. muridarum*-infected McCoy cells at 24 h p.i. indicated that all the PZ genes were transcribed (Fig. 2A). Next, we determined the transcriptional and temporal expression profiles of a subset of PZ ORFs (*tc0431* and *tc0436* to *tc0443*) using qRT-PCR analysis at various time points p.i. The developmentally regulated genes *euo*, *ompA*, and *omcB* were analyzed as representative chlamydial early, mid-late, and late genes, respectively (19). Increased expression of *euo* was detected by 6 h p.i., whereas *ompA* transcripts first increased between 6 and 12 h p.i. and *de novo omcB* expression was

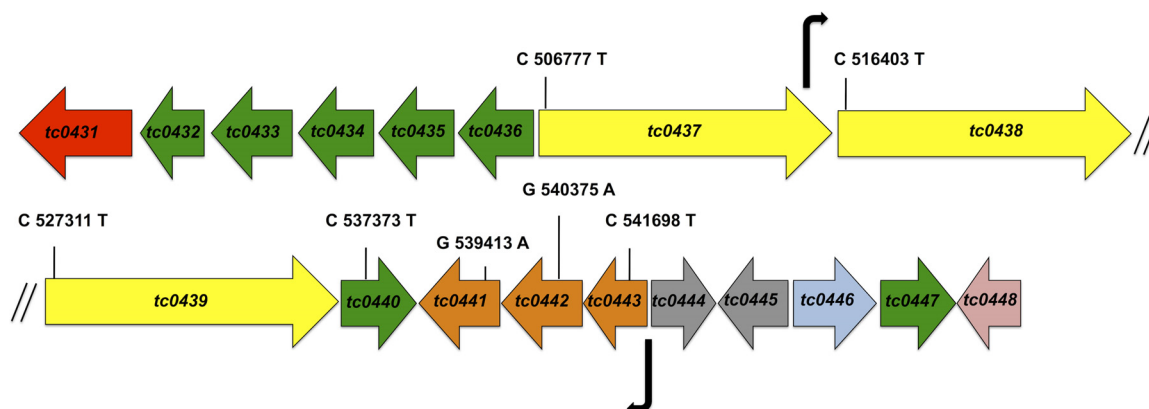


FIG 1 Map of the *C. muridarum* PZ and diagram of PZ ORFs. The ORFs are color coded according to function: red, MACPF; green, PLD; yellow, cytotoxin; orange, purine nucleotide synthesis; gray, conserved hypothetical; blue, peptide ABC transporter; pink, *dsbB*. The locations of promoters identified by β -galactosidase assays in this study are shown by the bent arrows. The genomic locations and nucleotide changes of the nonsense mutations discussed in this study are indicated in the corresponding ORFs. The diagram is not drawn to scale.

detected only at 12 h p.i. and did not peak until 18 h p.i. (Fig. 2B). The expression trend for all of the PZ genes most closely resembled that of *euo*, and the absolute peak numbers of PZ ORF transcripts were 1 to 3 log units lower than those of *ompA* and *omcB*. Taken together, these results showed that expression of most PZ ORFs initiates in the early to middle developmental cycle and that they are expressed at low levels relative to known highly expressed chlamydial genes.

We next sought to identify the promoters of PZ genes. The abilities of a series of overlapping cloned fragments of the *C. muridarum* PZ to drive expression of a promoterless *lacZ* gene were assayed in *E. coli* to identify promoters. Only two of the constructs exhibited strong promoter activity. One promoter locus (nucleotide positions 542868 to 542368 in GenBank AE002160.2) was located immediately upstream of *guaB* (see Table S5 in the supplemental material) and was in the correct position and orientation to drive expression of the putative operon *guaBA-add* (4). RT-PCR analysis of *guaB*, *guaA*, and *add* using primer pairs that amplified across intergenic regions of adjacent genes confirmed that these genes were cotranscribed (see Fig. S1 in the supplemental material). A specific primer pair spanning *add* and *tc0440* also gave rise to an overlapping amplification product from extracted RNA. However, since these two genes are predicted to be located on different strands, the RT-PCR product is likely a result of enzyme run-through. Two putative cytotoxin genes (*tc0437* and *tc0438*) were found to be in a single transcript (see Fig. S1 in the supplemental material), but the presence of an active internal promoter at the 3' end of *tc0437* (nucleotide positions 515330 to 515896) suggested that *tc0438* can also be expressed as a separate transcript (see Table S5 in the supplemental material). No promoter likely to drive expression of the operons *tc0436-tc0433* and *tc0432-tc0431* or the cytotoxin ORFs *tc0437* and *tc0439* was identified (see Fig. S1 and Table S5 in the supplemental material). Failure to identify other promoters for PZ genes whose expression was confirmed by RT-PCR could indicate that some PZ promoters were not functional in *E. coli* or were located outside the regions interrogated in the assay.

Members of three PZ gene families are dispensable for *C. muridarum* survival and proliferation *in vitro*. Since little is known about the potential functions of most PZ ORFs, we next

tested if they were dispensable using a reverse genetic approach. An EMS-mutagenized *C. muridarum* library was screened by TILLING for isolates that had nonsense mutations in any of eight *C. muridarum*-specific PZ ORFs: *tc0437*, *tc0438*, *tc0439*, *tc0440*, *guaA* (*tc0441*), *guaB* (*tc0442*), *add* (*tc0443*), and MACPF (*tc0431*) (13). The TILLING library was constructed with a high mutation load to facilitate faster screening. To estimate the number of EMS-induced point mutations in a given gene that would have to be isolated in order to recover at least one nonsense mutation, we assumed a Poisson distribution described as follows: $P(x \geq 1) = 1 - P(x = 0) = 1 - e^{-a}$, where $a = N \times p$ (N is the number of mutations in gene X that need to be isolated, and p is the probability of obtaining a nonsense mutation in gene X). The majority of mutations induced by EMS are GC \rightarrow AT transitions. A total of 5 GC transitions in 4 amino acid codons (CAA, CGA, CAG, and TGG) can yield a stop codon. We use *tc0442* as an example to illustrate how we arrived at an approximation of screen size (N). The probability p can be calculated as the number of possible nonsense GC transitions in the analyzed region of *tc0442* divided by the number of GC base pairs in the same region of *tc0442* ($p = 19/585 = 0.032$). For 95% screen saturation, screen size N was determined as follows: $P(x \geq 1) = 0.95 = 1 - e^{-0.032N}$, or $n = 94$. In other words, if *tc0442* is not essential, there is a 95% probability that at least one of 94 isolated mutants is a nonsense mutant. The screen sizes for other PZ genes were calculated by the same method (see Table S7 in the supplemental material).

One or more nonsense mutants in *guaBA-add*, the *C. muridarum*-specific PZ PLD *tc0440*, and the cytotoxins (*tc0437* to *tc0439*) were isolated at the anticipated frequencies, indicating that these ORFs are not essential for chlamydial survival *in vitro*. Despite attaining 95% saturation in our screen for nonsense mutants of *tc0431* (MACPF), all of the mutants that we isolated contained either missense or silent mutations (see Table S7 in the supplemental material). While it is possible that a larger screen of *C. muridarum* EMS mutants could have revealed a nonsense mutant, the relatively high frequency at which other PZ nonsense mutants were obtained argues against this interpretation. *Chlamydia abortus* does not encode MACPF (31, 32), and the *Chlamydia caviae* and human *Chlamydia pneumoniae* MACPF genes

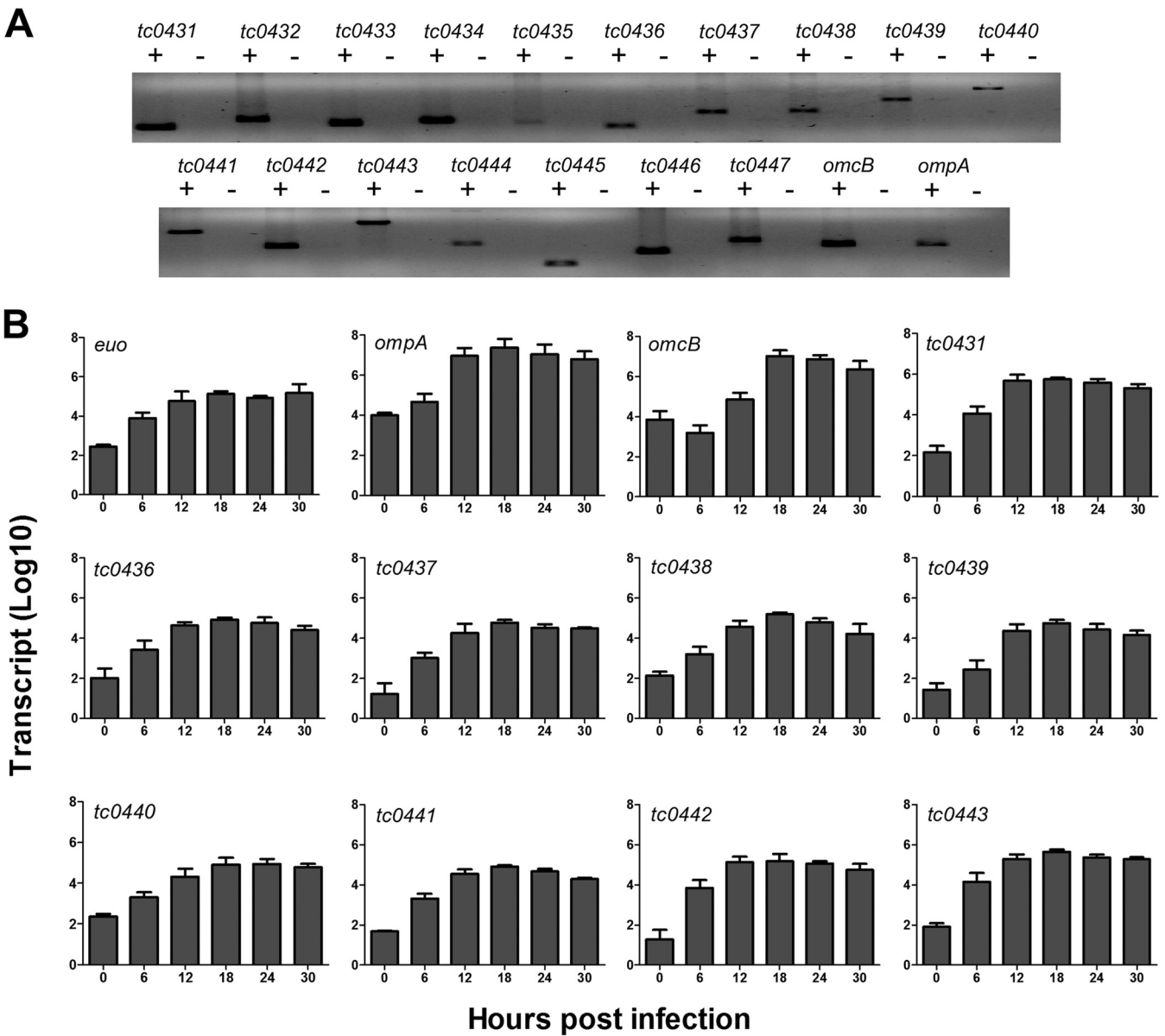


FIG 2 Transcription of *C. muridarum* PZ ORFs initiates in the early to middle phase of the developmental cycle. Total RNA was isolated from *C. muridarum*-infected McCoy cells at 0, 6, 12, 18, 24, and 30 h p.i. (A) RT-PCR at 24 h p.i. indicated that all genes in the PZ are transcribed. Whether amplification was performed with (+) or without (–) reverse transcriptase is indicated. (B) The kinetics of transcription of select PZ ORFs was characterized by qRT-PCR. Transcript levels were normalized to standard curves of dilutions of *C. muridarum* chromosomes. The data represent the mean transcript levels with standard deviations (SD) from three independent experiments. *euo*, *ompA*, and *ompB* represent early, mid-late, and late genes, respectively.

contain frameshifts (5, 32, 33), so our result could indicate that MACPF plays an essential, species-specific role in *C. muridarum*. **PZ mutants display mild *in vitro* growth defects.** The PZ nonsense mutants were initially characterized using temporal recoverable IFU assays. Since multiple independent mutants of *tc0440* and *guaA* were recovered by TILLING, we selected a representative mutant of each with a nonsense mutation located closest to the predicted 5' end of these ORFs for further analysis. The growth kinetics of most of the PZ mutants paralleled those of *C. muridarum*, with the exception of *add* and *guaB*, which attained maximal EB production at 18 h p.i. as opposed to 30 h p.i. None of the mutants exhibited dramatic growth defects, although the IFU

production of most of the mutants was significantly reduced compared to that of *C. muridarum* (Fig. 3). This modest reduction in IFU yield could be caused either by mutations in the PZ or by background mutations in the strains. Confocal microscopy failed to reveal obvious differences between the morphology of the mutant and *C. muridarum* inclusions (data not shown). These results implied that the presence of nonsense mutations in PZ ORFs did not hinder the ability of *C. muridarum* to form inclusions, complete development, or produce infectious EB. **The cytotoxicity of cytotoxin nonsense mutants is reduced.** Since none of the PZ mutants displayed pronounced growth or morphological defects, we focused on examining phenotypes that

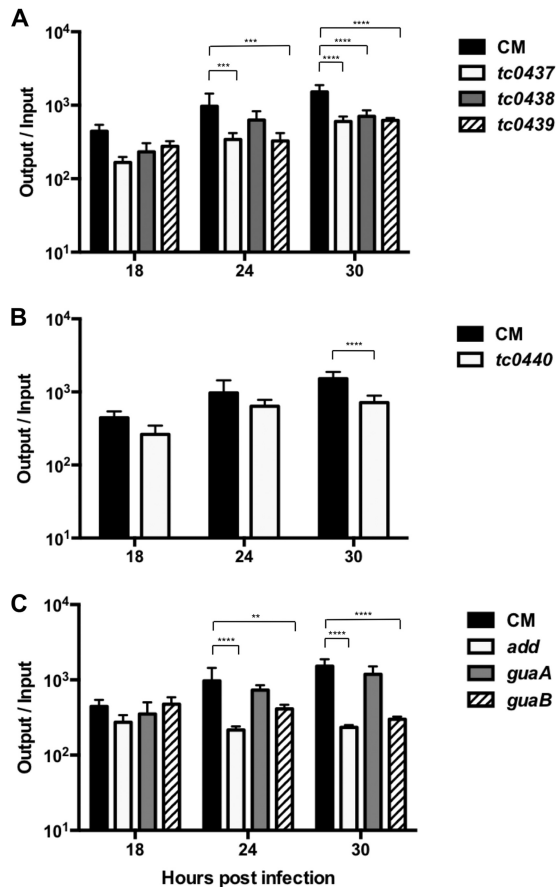


FIG 3 *C. muridarum* PZ mutants exhibit mild growth defects. rIFU analysis was performed for *C. muridarum* (CM) and PZ mutants at various time intervals following infection of McCoy cells. The data shown are from experiments performed in parallel. The data represent the average (\pm SD) ratios of input to output IFU from three independent experiments performed in triplicate. ****, $P < 0.0001$; ***, $P < 0.001$; **, $P < 0.01$ by two-way ANOVA with Bonferroni *post hoc* test.

have been indirectly attributed to the chlamydial cytotoxins in previous studies. The *C. muridarum* LCT-like cytotoxins are purported to mediate multiplication-independent cytopathic effects of *C. muridarum* EB on host cells at high MOI (19). This phenomenon, referred to as “immediate cytotoxicity” (34), resembles the cytoskeletal collapse observed in cultured cells treated with clostridial TcdB (19).

To determine if the *C. muridarum* cytotoxin gene (*tc0437*, *tc0438*, and *tc0439*) nonsense mutants were less cytotoxic, HeLa cells were infected with EB at an MOI of 250 in the presence of rifampin to block chlamydial multiplication. The cells were fixed and stained with phalloidin at 3 h p.i. and examined by fluorescence microscopy. HeLa cells infected with *C. trachomatis* serovar L2 strain 434/Bu (*C. trachomatis*), a cytotoxin-negative strain, did not differ in morphology from uninfected cells. In contrast, cell rounding and alterations in actin filament morphology were observed in infections with *C. muridarum* and the cytotoxin mutants (Fig. 4A). This implied that the single cytotoxin mutants retained some cytotoxic activity.

Microscopy could not distinguish small variations in cytotoxicity, so we measured LDH release at 3 h p.i. from cells infected with *C. muridarum*, *C. trachomatis*, and the mutants at various MOI. As expected, *C. trachomatis* elicited minimal LDH release even at an MOI of 500. In contrast, *C. muridarum* caused cells to release significantly more LDH at high MOI. Interestingly, the *tc0437* and *tc0439* mutants elicited lower LDH release than *C. muridarum* at the highest MOI tested (Fig. 4B). This could indicate that *tc0437* and *tc0439* wild-type alleles encode active toxins or, alternatively, that background mutations in the mutant strains affected EB cytotoxic activity. Undiminished cytotoxicity of the *tc0438* mutant implies that TC0438 may not be functional or has an unrelated function.

IFN- γ resistance is unaltered in *C. muridarum* PZ mutants.

While high-multiplicity infections were important in associating a toxin-like activity with chlamydial EB, these doses may not be relevant in natural chlamydial disease. Roles for the chlamydial cytotoxins in host cell attachment and entry, vesicle trafficking,

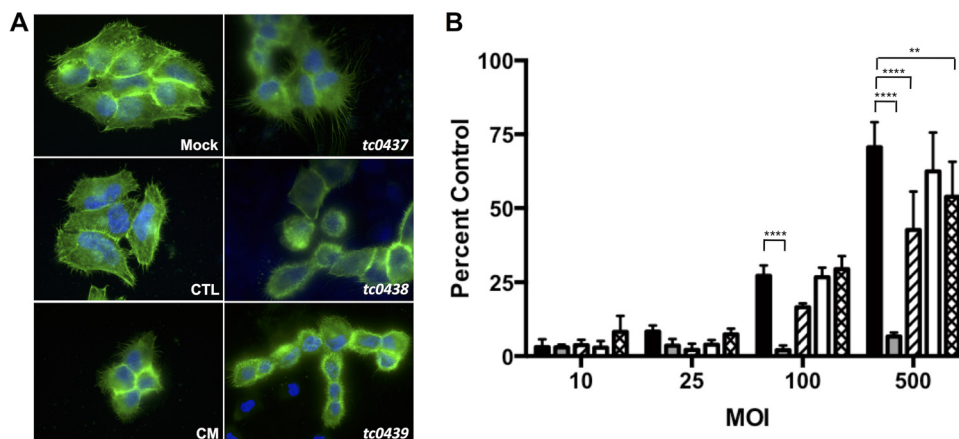


FIG 4 *tc0437* and *tc0439* mutants have reduced cytotoxicity. (A) HeLa cells were infected at an MOI of 250 in the presence of rifampin and were fixed 3 h p.i. Cell morphology and actin structure were visualized by staining with Alexa Fluor 488-phalloidin (green) and DAPI (blue). Overlays of fluorescence micrographs of cells that were mock infected or infected with *C. muridarum* (CM), *C. trachomatis* (CTL), *tc0437*, *tc0438*, or *tc0439* are depicted. The images are representative of the results of three independent experiments. (B) LDH in supernatants of HeLa cells infected at various MOI in the presence of rifampin. Shown are *C. muridarum* (black), *C. trachomatis* (gray), *tc0437* (hatched), *tc0438* (white), and *tc0439* (crosshatched). Triton X-100-treated cells were used as positive controls for LDH release. The data show the mean percentages (\pm SD) of the positive control from three independent experiments performed in triplicate. ****, $P < 0.0001$; **, $P < 0.01$ by one-way ANOVA with Dunnett's *post hoc* test.

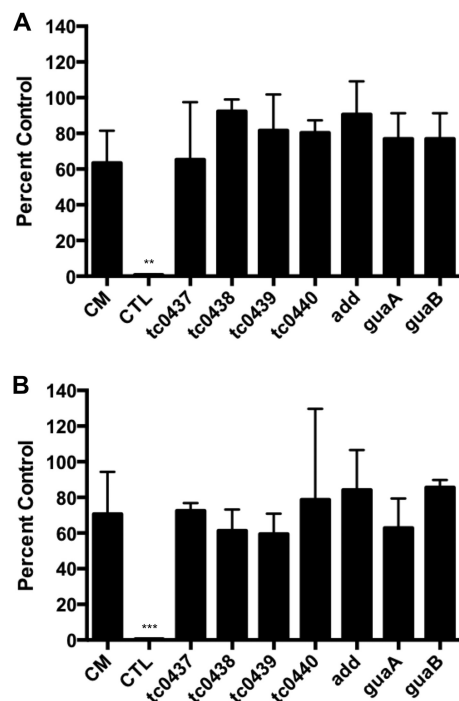


FIG 5 PZ mutants are resistant to IFN- γ . The sensitivity to IFN- γ treatment of *C. muridarum* (CM), *C. trachomatis* (CTL), and various PZ mutants was assessed by IFU assay (A) and rIFU assay (B). The data are represented as the average (\pm SD) percentages of untreated (no IFN- γ) infected controls from three independent experiments performed in triplicate. ***, $P < 0.001$; **, $P < 0.01$ by one-way ANOVA with Dunnett's *post hoc* test.

and modification of host cell GTPases have been reported (35–37). The *C. muridarum* cytotoxins have also been proposed to mediate evasion of the inhibitory effects of IFN- γ responses in murine cells (12). To test the latter hypothesis, McCoy cells were treated with IFN- γ , followed by infection with *C. trachomatis*, *C. muridarum*, or cytotoxin mutants. At 24 h, inclusions were either directly counted (Fig. 5A) or harvested for enumeration of viable EB contained by the inclusions using IFU assays (Fig. 5B). *C. trachomatis*, but not *C. muridarum*, was dramatically inhibited by IFN- γ . IFN- γ treatment did not significantly affect the survival or infectivity of the cytotoxin mutants compared to *C. muridarum*. Parallel experiments with other PZ nonsense mutants indicated that they were also similarly resistant to IFN- γ (Fig. 5). Thus, disruption of individual cytotoxin or other PZ ORFs did not confer IFN- γ sensitivity to *C. muridarum*.

PZ mutants retain virulence in the mouse genital tract infection model. Intravaginal inoculations of mice with human *C. trachomatis* isolates result in mild infections of short duration with minimal upper genital tract infection and pathology (38). In comparison, *C. muridarum* is more virulent in this model, and the postinfection sequelae mimic human disease (39). To determine if the *C. muridarum* PZ genes were niche-specific virulence factors in mice, PZ mutants were analyzed in the murine genital tract infection model. The chlamydial burden in mice infected with the tc0439 cytotoxin mutant was significantly lower at 3 days postinfection than in animals infected with *C. muridarum*. Infection trajectories in mice inoculated with the tc0437 and tc0438 mutants also trended lower during the first week of infection. However, chlamydial recoveries of all three cytotoxin mutants appeared

identical to *C. muridarum* infection at later time points (Fig. 6A). We also did not observe any significant deviation in bacterial loads of mice infected with the tc0440 mutant compared to mice infected with *C. muridarum* (Fig. 6B). Interestingly, the *guaA* nonsense mutant behaved identically to *C. muridarum*, whereas *guaB* and *add* mutants displayed reduced pathogen burdens at early data points (Fig. 6C and D). The *guaBA-add* operon putatively encodes products that function in the same purine-biosynthetic pathway. The absence of bacterial-shedding phenotypes in the *guaA* mutant suggested either that the phenotypes of the *guaB* and *add* mutants were due to background mutations or that *guaBA-add* acts noncanonically in *C. muridarum*. Since none of the PZ mutants were completely attenuated *in vivo*, we conclude that these genes are dispensable for infection of the murine genital tract.

Infected mice were also scored at 73 days postinfection for the presence of hydrosalpinx, an immunopathological consequence of chlamydia genital infection. Eighty percent (24/30) of the mice infected with *C. muridarum* developed hydrosalpinx compared to 63% (10/16) of *guaA*, 63% (5/8) of *guaB*, 13% (1/8) of *add*, 75% (9/12) of *pld*, 38% (3/8) of tc0437, 50% (4/8) of tc0438, and 38% (3/8) of tc0439 mutant-infected mice. The development of hydrosalpinx was significantly lower in *add*, tc0437, and tc0439 mutant-infected mice ($P < 0.001$, $P = 0.03$, and $P = 0.03$, respectively). It should be noted, however, that the differences observed in hydrosalpinx cannot necessarily be attributed to specific PZ gene mutations due to other background mutations found in the mutants.

PZ mutants contain multiple background mutations. To identify a plausible explanation for the differing phenotypes of the purine pathway mutants, we sequenced the genomes of the parent strain and PZ mutants to an average coverage of 2,075 \times on the Illumina HiSeq platform (Table 1; see Tables S8 to S14 in the supplemental material). Comparison to published *C. muridarum* genomes indicated that our parent most closely resembled a *C. muridarum* Weiss isolate that was recently sequenced by K. H. Ramsey et al. (Table 1). On average, the PZ mutants contained 36.4 mutations, 66.67% of which were nonsynonymous substitutions. Most (98.43%) of the mutations were GC \rightarrow AT transitions, consistent with EMS-induced changes. Several synonymous and nonsynonymous mutations were shared by some PZ mutant strains, which was a consequence of our serial mutagenesis strategy. They included mutations in *recA*, *alaS*, *tc0237*, *tc0283*, *tc0471*, *tc0490*, *rpoC*, *aroA*, *upp*, and *tc0877* that were common to the *add*, tc0437, and tc0438 mutants. The *add* and tc0437 mutant strains contained identical mutations in *tc0035*, *tc0069*, *tc0438*, *tc0575*, *tc0917*, and *pfkA-2*. The tc0438 mutant shared mutations in *murB*, *tc0330*, and *tc0575* with the *add* mutant and a *pmpB-pmpC-2* mutation with the tc0437 mutant. Finally, mutations in *tc0054*, *tc0191*, *pmpD*, *tc0250*, *tc0290*, *tc0312*, *tc0600*, and an intergenic region (GenBank genome position 126188) were present in the tc0439 and *guaB* mutants. Interestingly, multiple mutant strains had nonsense mutations in *tc0412*, an ortholog of *C. trachomatis* *ct135* that has been linked to virulence of *C. trachomatis* in the murine genital tract model (40). None of the tc0412 mutants except *add* were significantly attenuated in the murine GT, and an identical stop mutation was present in the tc0437 mutant strain, suggesting that attenuation of the *add* mutant was not linked to this tc0412 allele. No common mutations were present in the *guaB* and *add* mutants, which is consistent with the hypothesis that

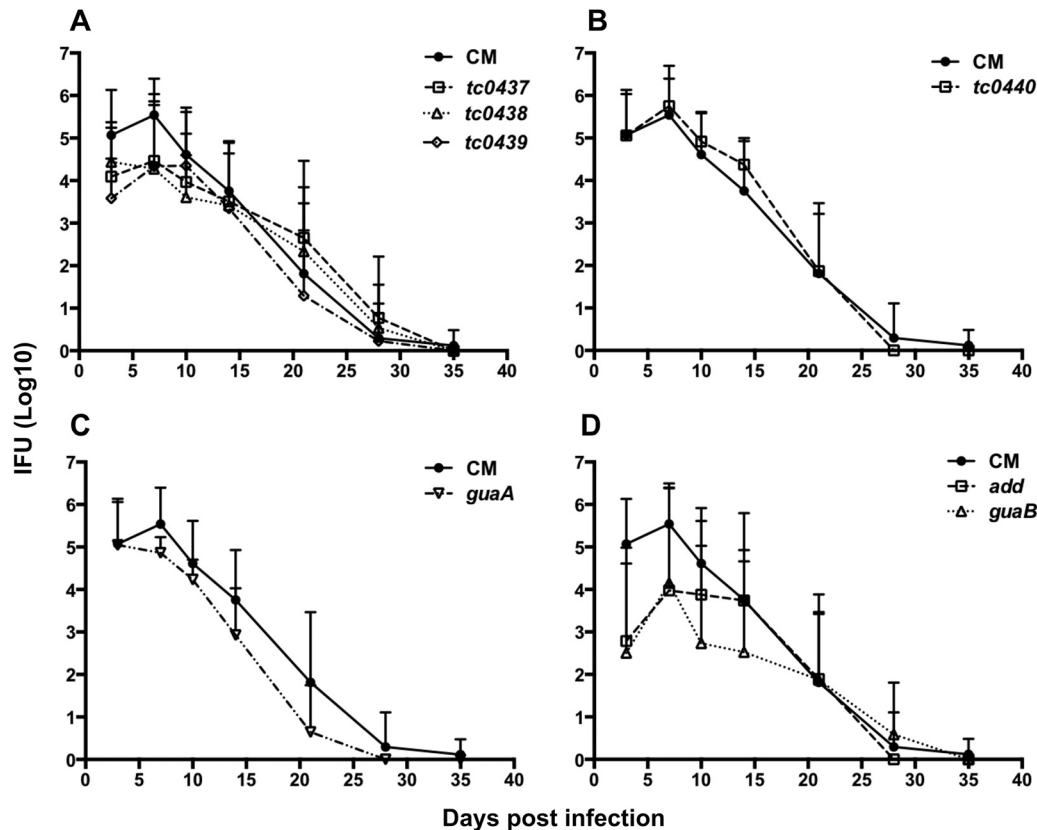


FIG 6 Mouse genital tract infections with *C. muridarum* PZ mutants. Groups of mice were challenged intravaginally with 50,000 IFU of *C. muridarum* (CM) or various PZ mutants. The infection curve for *C. muridarum*-infected mice ($n = 30$) is reproduced in each panel for comparison. (A) Lower genital tract shedding of IFU by mice infected with *C. muridarum* differed significantly from that by mice infected with the *tc0439* mutant ($n = 8$) at day 3 postinfection ($P < 0.01$), but not that by mice infected with *tc0437* ($n = 8$) or *tc0438* ($n = 8$). (B and C) IFU shedding from mice infected with CM did not vary significantly from that from mice infected with the *tc0440* ($n = 12$) (B) or *guaA* ($n = 16$) mutant (C). (D) Shedding by mice infected with the *add* ($n = 8$) mutant was significantly less than shedding by mice infected with *C. muridarum* at 3 days ($P < 0.0001$) and 7 days ($P < 0.01$) postinfection. IFU shedding from mice infected with the *add* ($n = 8$) mutant was also significantly reduced at 3 days ($P < 0.0001$), 7 days ($P < 0.05$), and 10 days ($P < 0.001$) postinfection. The data are presented as mean numbers of IFU (\pm SD) for the mice in each group. Statistical differences were analyzed by two-way ANOVA with Bonferroni *post hoc* test.

guaB and *add* might function in a noncanonical pathway independently of *guaA*.

DISCUSSION

C. muridarum and *C. trachomatis* are genetically closely related, and genital infections of their respective hosts, mice and humans, have many similarities (4, 41). While T-cell immunity is critical for the resolution of chlamydial infection in both mice and humans, other aspects of immunity, most notably IFN- γ responses, appear to differ and are incompletely characterized (38). The results of our study suggest that although the PZ is the site of maximal divergence between the genomes of these pathogens, *C. muridarum* IFN- γ resistance determinants may not localize to the region.

We show that most PZ ORFs are transcribed in the early to middle phase of the developmental cycle and are expressed at relatively low levels. Although we identified two promoter regions that support our RT-PCR findings and that could provide plausible explanations for how some PZ ORFs are transcribed, strong promoters that could drive expression of one of the cytotoxin ORFs (*tc0437*) and an adjacent operon of PZ PLD (*tc0436* to *tc0432*) that is relatively conserved in *Chlamydia* spp. were not

identified. The putative start sites of *tc0436* and *tc0437* are located less than 1,000 bp apart in opposite strands of the *C. muridarum* genome, so perhaps our chosen cloning site disrupted two promoters in the same narrow region on opposing strands, thus compromising their detection. Another possibility is that *E. coli* sigma factors cannot recognize the chlamydial promoters in this region. For example, the auxiliary *C. trachomatis* transcription factor GrgA, which is not conserved in *E. coli*, directs expression from some chlamydial σ^{66} -dependent promoters (42). Chlamydial σ^{28} also cannot complement an *E. coli* σ^{28} mutant, indicating that even conserved *E. coli* and chlamydial sigma factors are not completely interchangeable (43). We attempted to identify additional promoters using transcriptome sequencing (RNA-seq), but low expression of PZ ORFs relative to other, highly expressed genes yielded insufficient coverage for mapping transcriptional start sites in this region. As costs of deep RNA-seq decrease further and methods for enrichment of chlamydial transcripts from host cell transcripts improve, it might make sense to revisit this approach in the future. Better understanding of transcriptional organization could inform subsequent attempts to inactivate PZ operons.

Two of three cytotoxin mutants displayed reduced cytotoxicity *in vitro*, which suggests that these alleles are functional. We hope

TABLE 1 Comparison of the genome of the wild-type *C. muridarum* isolate used in this study to those of other *C. muridarum* reference strains

Gene ID	Description	Position	Base		
			<i>C. muridarum</i> used in study	<i>C. muridarum</i> Nigg ^a	<i>C. muridarum</i> Weiss ^b
tc0007	Exodeoxyribonuclease V, beta chain, putative	10505	A		
		10558	A,C		
		10564	A		
tc0027	Conserved hypothetical protein	32915	G		
tc0052	Major outer membrane protein, porin; OmpA	58882	T		
		58904	T		
		59065	C	C	T
tc0107	Lipoprotein	126406	G	G	
		126416			A
		126383	G		
None	Intergenic	126436	A		
		126475		A	
tc0124	Transcription repair-coupling factor; TrcF	151212	T	T	
tc_r05	rRNA-23S rRNA	158800	G		
tc0138	Phospho-N-acetylmuramoylpentapeptide transferase; MraY	169451	TTTT	T	T
tc0168	Ribosomal protein L34; RpmH	200671	C	G	C
tc0301	Methionyl tRNA synthetase; MetG	358414	T		
tc0338	ABC transporter, periplasmic substrate binding protein	401302	G		
tc0341	ABC transporter, permease protein	403623		T	
		403624	C	A	A
		403626			C
		403652	C,G	T	C
		403714	G		
tc0342	ABC transporter, permease protein	404473	T		
		404884	A	A	
		405819	G		
tc0343	1-Deoxy-D-xylulose 5-phosphate reductoisomerase; Dxr	419119	C		
tc0359	Hypothetical protein	468932	T	G	T
tc0408	Hypothetical protein	473118	A	A	
tc0412	Hypothetical protein	515204	A	C	C
tc0437	Adherence factor	600435	A	C	C
tc0493	Phenylacrylic acid decarboxylase; UbiX	712587	C		
tc0708	Hypothetical protein	846475	G	G	
		866121	T	G	T
tc0727	Outer membrane protein; OmcB	967487	A	A	
tc0832	DNA polymerase III alpha subunit; DnaE	1022529	A		
tc0879	Hypothetical protein	1022551	A		
		1035913	A		
tc0893	groEL-2				

^a *C. muridarum* Nigg isolate accession number: [AE002160.2](#).
^b *C. muridarum* Weiss isolate accession number: [ACOW00000000](#).

to verify this in future experiments using allelic complementation or by utilizing isogenic cytotoxin mutant strains. The biological role of chlamydial cytotoxins is still largely a matter of speculation. Polymorphisms in the cytotoxins have been associated with differences in the ocular and genital tropisms of *C. trachomatis* isolates (20). On the basis of their homology to LCTs and YopT, we have previously proposed that the cytotoxins may inactivate interferon-regulated GTPases that inhibit *C. trachomatis*, but not *C. muridarum*, in IFN-γ-treated mouse cells (44). Finally, mutants of *E. coli efa1*, another chlamydial cytotoxin gene homolog, exhibit reduced adherence to epithelial cells and fail to colonize colonic tissue effectively in mice (45). The observed trend toward a lower chlamydial burden during the first week of infection in mice inoculated with the chlamydial cytotoxin mutants could indicate similar roles for *C. muridarum* cytotoxins in adhesion and colonization, but this hypothesis requires experimental validation. An

additional future direction would be to evaluate the IFN-γ resistance and virulence of double and triple cytotoxin mutants. One of the most surprising findings of our study was that the *guaB* (IMP dehydrogenase) and *add* (adenosine deaminase) mutants were moderately attenuated in the GT while a *guaA* (GMP synthase) mutant was not. Genome sequencing confirmed that the phenotypes of the *guaB* and *add* mutants were not caused by a common background mutation. Thus, if unrelated background mutations were not responsible for the very similar GT phenotypes, our results suggest *C. muridarum* could encode an additional cryptic GMP synthase enzyme or that *C. muridarum* purine salvage acts noncanonically. The *guaBA-add* cluster has been retained by several chlamydial species, including those that cause respiratory illness, such as *Chlamydia psittaci*, *Chlamydia felis*, and human isolates of *C. pneumoniae*, but it is unknown if they are functional (33, 46, 47). *guaA* and *guaB* are known virulence deter-

minants in other pathogenic organisms, like *Borrelia burgdorferi* and *Francisella tularensis*, during intraperitoneal and subcutaneous infection of mice, indicating that one or more purine nucleoside monophosphates (NMP) are limiting in certain mammalian niches (48, 49). Thus, an explanation of our results could be that NMP levels in the murine GT are sufficient to support chlamydial growth, whereas other tissues, like the lung, are starved for purine nucleotides. We hope to differentiate these possibilities by constructing additional *guaA*, *guaB*, and *add* nonsense mutants and comparing their virulence in murine GT and lung infection models.

Although we were unable to identify a role for individual PZ genes in IFN- γ resistance and murine genital infection, functional redundancy in the cytotoxin and PLD superfamily of proteins could have masked the effects of the loss of a single family member. However, our data suggest that cytotoxin functions are not wholly redundant, and true functional redundancy of these alleles is not consistent with the relatively low amino acid identity to one another (40 to 42%). While it would be possible to disrupt all members of the cytotoxin family and the *guaBA-add* operon sequentially, variable alleles, including *tc0412*, are a practical impediment to constructing isogenic strains by iterative TILLING (40, 50). Another possible explanation for the lack of phenotype of select PZ mutants is the emergence of suppressor mutations. Although this possibility is difficult to exclude, the fact that all of the PZ mutants were isolated from pools of mutants at the expected frequencies indicates that it is unlikely to have been a significant issue. We are currently investigating the possibility of disrupting the MACPF-PZ PLD (*tc0432-tc0431*; *tc0436* to *tc0433*), cytotoxin (*tc0437-tc0438*), and *guaBA-add* operons using polar transposon insertions by the Targetron method developed by Johnson and Fisher (51).

In summary, our results imply that all the examined PZ ORFs are nonessential for virulence in the mouse GT. However, we cannot rule out their importance in other niches. *C. muridarum* was originally isolated from the lungs of symptom-free laboratory mice. Disease was observed only upon inoculation of mice with serially passaged lung homogenates, indicating that very low levels of *C. muridarum* are typically present in pulmonary tissue (52). Later studies demonstrated that *C. muridarum* is transmitted through the oral-fecal route, which suggests that the gut is a reservoir and that small numbers of *C. muridarum* organisms enter the lungs during feeding (53). Finally, it is notable that *guaBA-add* and cytotoxin homologs of other intracellular pathogens modulate infections of the lung and gastrointestinal tract, respectively (45, 48, 54, 55). Thus, examination of *C. muridarum* PZ mutants in lung or gastrointestinal tract models of mouse infections is warranted. A positive implication of our study is that successful modeling of human disease in the murine GT does not appear to require much of the *C. muridarum* PZ. Future investigations of polymorphisms outside the PZ will be important in the identification of *C. muridarum* alleles that confer IFN- γ resistance and dictate virulence in the murine GT model.

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