The *Legionella pneumophila* GIG operon responds to gold and copper in planktonic and biofilm cultures

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Abstract

*Legionella pneumophila* contaminates man-made water systems and creates numerous exposure risks for Legionnaires’ Disease. Because copper/silver ionization is commonly used to control *L. pneumophila*, its mechanisms of metal response and detoxification are of significant interest. Here we describe an *L. pneumophila* operon with significant similarity to the GIG operon of *Cupriavidus metallidurans*. The *Legionella* GIG operon is present in a subset of strains and has been acquired as part of the ICE-βox 65-kB integrative conjugative element. We assessed GIG promoter activity following exposure of *L. pneumophila* to multiple concentrations of HAuCl₄, CuSO₄ and AgNO₃. At 37˚C, control stationary phase cultures exhibited GIG promoter activity. This activity increased significantly in response to 20 and 50uM HAuCl₄ and CuSO₄ but not in response to AgNO₃. Conversely, at 26˚C, cultures exhibited decreased promoter response to copper. GIG promoter activity was also induced by HAuCl₄ or CuSO₄ during early biofilm establishment at both temperatures. When an *L. pneumophila* GIG promoter construct was transformed into *E. coli* DH5α, cultures showed baseline expression levels that did not increase following metal addition. Analysis of *L. pneumophila* transcriptional regulatory mutants suggested that GIG up-regulation in the presence of metal ions may be influenced by the stationary phase sigma factor, RpoS.

Introduction

*Legionella pneumophila*, the etiological agent of Legionnaires’ Disease (LD), is the leading cause of bacterial waterborne disease outbreaks in the United States [1]. This Gram-negative bacterium is ubiquitous in both natural and man-made aquatic environments, where it replicates as an intracellular parasite of free-living protozoa [2]. Most cases of LD can be traced back to human-made aquatic systems with above-ambient water temperatures: cooling towers, hot water heaters, fountains, and air conditioning units have all served as sources of outbreaks [2].
Legionella persist as part of the biofilm community in human-made aquatic systems, and these systems are routinely treated to inhibit microbial growth. Chlorine and chloramine are the most common disinfectants used in the US. Other treatments include chlorine dioxide, UV radiation, hyperchlorination, bromine, and copper/silver ionization. While such treatments work well against fecal coliforms and other bacteria that enter the system from outside sources, they are less effective at eliminating pathogens like Legionella that exist within resistant biofilms of the system itself. Copper/silver ionization is often used to control Legionella in large recirculating water systems, particularly industrial plumbing, but its effectiveness is controversial [3–6]. Some studies indicate that high levels of copper inhibit Legionella growth and survival, but others have demonstrated increased persistence of Legionella in biofilms formed on copper [7]. Legionella also demonstrates increased resistance to copper at lower temperatures [8–9].

Small amounts of copper are required for mitochondrial electron transport and other enzymatic reactions, but high intracellular copper levels are considered toxic to most prokaryotes [10–12]. General mechanisms of bacterial metal resistance include export across the plasma membrane, sequestration by binding proteins, and reduction to a less toxic state. While Legionella is sensitive to elevated concentrations of numerous metals [13], few of its metal resistance mechanisms have been described. A Legionella copper-translocating PIB-type ATPase (CopA) was shown to confer copper resistance when expressed in a copper-sensitive E. coli strain [14], and the helABC locus was reported to encode three proteins involved in heavy metal resistance and cytopathogenicity [15]. Additional mechanisms by which Legionella sense and respond to metal ions in their environment remain to be characterized.

Here we describe a Legionella pneumophila operon (lpq2105-2108) with significant homology to the “gold induced genes” (GIG) operon of Cupriavidus metallidurans [16]. The operon appears to have been acquired by a subset of L. pneumophila strains as part of the horizontally-transferred ICE-βox integrative conjugative element [17]. In planktonic cultures, promoter expression occurs at 37˚C in response to gold and copper, but at 26˚C it occurs only in response to growth phase. Under biofilm conditions, promoter response to metal ions is seen at both temperatures.

**Material and methods**

**Identification of the L. pneumophila GIG operon and homologous operons**

The L. pneumophila GIG operon was originally identified through a BlastP search of proteins from the L. pneumophila subsp. pneumophila str. Philadelphia 1 genome (NCBI NC_002942.5), using the four C. metallidurans GIG proteins as queries (Table 1). Additional homologous operons in L. pneumophila were subsequently identified through BlastP searches of five sequenced L. pneumophila subsp. pneumophila strains, using the L. pneumophila GIG proteins as queries. Hits with a query cover of at least 80% and an E-value of less than 1.0E−15 were retained in this analysis. Homologous operons were also identified in the genomes of the Francisella tularensis (NC_006570.2), Burkholderia pseudomallei (NC_012695.1), Polaramonas sp. JS666 (NC_007948.1) and Pseudomonas fluorescens (NC_007492.2) using similar methods. Predicted protein sequences from each operon were concatenated and aligned with MAFFT v.7 [18]. A maximum likelihood tree was constructed in PhyML using the LG amino acid substitution model and the SPR method of topology estimation [19]. Reliability of the tree was assessed with 500 bootstrap replicates, and branches reproduced in fewer than 50% of the replicates were collapsed.
Strains and media

In all experiments, wild type *L. pneumophila* strain Lp02 was cultured at 37˚C or 26˚C for three days on buffered charcoal yeast extract agar with 100μg/ml thymidine (BCYE-T). *L. pneumophila* Lp02 is a derivative of *L. pneumophila* Philadelphia 1 and contains the complete ICE-βox region. Lp02ΔletA and ΔrpoS mutants were grown on BCYE with 20μg/ml kanamycin. All *Legionella pneumophila* strains were provided by Michelle Swanson (University of Michigan). *Escherichia coli* DH5α was grown at 37˚C or 26˚C for 24 hours on Tryptic Soy Agar (TSA). Broth cultures consisted of ACES (N-(2-Acetamido)-2-aminoethane sulfonic acid)-buffered yeast extract (AYE) for *Legionella* strains (with antibiotics and thymidine as necessary) and Tryptic Soy Broth (TSB) for *E. coli* strains.

Reverse Transcriptase-PCR. Total RNA was isolated from stationary phase cultures using the Promega SV RNA Isolation kit. RNA was DNase treated to ensure removal of contaminating DNA prior to RT-PCR analysis. Three sets of primers spanning the four genes of the operon (2105–2106; 2106–2107; 2107–2108) were used to confirm the single transcript containing the four genes. The Verso 1-step Reverse Transcriptase PCR kit was used following manufacturer’s instructions. Reactions were carried out in 25μl final volume using each primer set and 15ng of RNA per reaction. Amplified fragments were analysed on a 1.0% agarose gel.

Construction of pGIG reporter gene vector

The flaA promoter of pflaA (a GFP reporter gene vector provided by M. Swanson, [20]), was replaced with 180nt of the upstream lpg2105-2108 predicted promoter region using standard cloning methods. The 180nt upstream of lpg2105-2108 represents the intergenic region between lpg2108 and lpg2109. Bacterial promoter regions have been shown to be enriched in the intergenic regions of the genome and depleted from coding regions [21]. The entire intergenic region was cloned in order to capture as many potential promoter binding sites as possible for subsequent experiments. The resulting plasmid, pGIGgfp, was transformed into *L. pneumophila* Lp02, *E. coli* DH5α, *L. pneumophila* Lp02 ΔletA, and *L. pneumophila* Lp02 ΔrpoS.

Reporter gene activity in planktonic cultures

The effect of metal ions on *L. pneumophila* growth kinetics and pGIGgfp reporter gene activity were measured by incubating planktonic cultures for 72 hours at 150rpm in the presence or absence of gold, copper or silver ions. Cultures were incubated at 37˚C with 20μM or 50μM of...
gold chloride (HAuCl$_4$), 20µM or 50µM copper sulfate (CuSO$_4$), or 50µM or 150µM silver nitrate (AgNO$_3$), or no additional ions (control). Cultures were incubated at 26˚C with 5µM and 10µM HAuCl$_4$, 200µM and 275µM CuSO$_4$, or no additional ions (control). Every three hours, absorbance (OD$_{600}$) was measured using a Genesys 6 spectrophotometer, and GFP fluorescence (485 nm excitation/528 nm emission) was measured using a Biotek Synergy H1 plate reader.

The effect of metal ions on *E. coli* growth kinetics and pGIGgfp reporter gene activity was measured by incubating planktonic cultures for 24 hours at 150rpm in the presence or absence of gold or copper. Cultures were incubated at 37˚C with 20µM or 50µM HAuCl$_4$, 20µM or 50µM CuSO$_4$, or no additional ions. Absorbance and GFP fluorescence were measured hourly as described above.

All experiments were performed in triplicate. Data were normalized by dividing the GFP relative light units (RLU) by the OD$_{600}$. The effects of metal ion, ion concentration, growth phase, and their interactions on the magnitude of GFP fluorescence were each analyzed separately.

**Reporter gene activity in biofilms**

*L. pneumophila* Lp02 pGIGgfp biofilms were established as previously described [22–23]. Briefly, bacteria were inoculated into glass petri dishes containing glass slides in 10% AYE solution and incubated for 24 hours, then transferred to 100% AYE for the remainder of the incubation. Cultures were incubated at 26˚C or 37˚C with 20µM HAuCl$_4$, 20µM CuSO$_4$, or no additional ions. Biofilms were grown for 120 hours and assessed for GIG activity at 24, 48, 72, 96, and 120h. At each time point, biofilms were washed twice with sterile ultrapure water (UPW) to remove non-attached bacteria. Slides were aseptically removed, briefly air dried, and fixed in paraformaldehyde for 10 minutes. Slides were rinsed with UPW and dried; then coverslips were mounted using a 50/50 v/v solution of glycerol:phosphate buffered saline (1X PBS). All experiments were performed in triplicate.

**Image analysis**

Biofilms were imaged using a Leica SPE spectral confocal microscope (63X, oil immersion objective, NA = 1.30; Leica Microsystems, Buffalo Grove, IL) in the Clemson Light Imaging Facility. Three DIC (differential interference contrast) images and three corresponding GFP images were obtained for each slide. For assessment of promoter activity, ImageJ software and DIC images were used to generate regions of interest (ROIs) corresponding to individual bacteria within the biofilm. The ROIs were outlined and numbered, and their areas were measured. A binary image “mask” of the ROIs was used as an overlay on the corresponding GFP image. The signal intensity within each ROI on the GFP image was measured and used as an indication of GFP expression in each bacterium (ROI). All biofilm samples produced some level of green fluorescence. To account for this primary fluorescence, the intensity of control biofilms was determined and gated out (subtracted) from the intensity of treated biofilms to determine GFP expression due solely to the addition of gold or copper. The 37˚C control fluorescence intensity value was applied to all samples. Treatment effects were assessed using Z-scores with a 99% confidence interval.

To measure the biofilm biomass that demonstrated GIG activity, confocal images from each time point were analyzed using COMSTAT software [24]. Bio-volume was estimated from calculations of biofilm biomass, which were based on the number of bacteria-containing pixels in all images of a stack, multiplied by voxel size and divided by the stack’s substratum
area. A one way analysis of variance (ANOVA) was used to compare bio-volumes among biofilms and time points. A significance level of \( p < 0.05 \) was used for all tests.

**Results**

**Description of the *L. pneumophila* GIG operon**

The *L. pneumophila* GIG operon spans 2421 bases from 2352473 to 2354894 in the *L. pneumophila* subsp. *pneumophila* str. Philadelphia 1 genome (NC_002942.5). It is located within the ICE-\( \beta \)ox, a 65-kB integrative conjugative element found in approximately 18% of surveyed *Legionella* strains and associated with increased virulence and oxidative stress tolerance [17, 25]. The operon contains four open reading frames encoding proteins of unknown function (\( lpg2105 \)-\( lpg2108 \)).

The first *L. pneumophila* GIG gene, \( lpg2105 \), encodes a 165-aa predicted inner membrane protein with a DoxX domain (PF07681). DoxX proteins exhibit some similarity to the thiosulphate:quinone oxidoreductase small subunit, DoxD, but their precise function is unknown [26]. The second GIG gene, \( lpg2106 \), encodes a 259-aa protein with an N-terminal DUF2063 domain that is predicted to function in DNA binding and transcriptional regulation [27]. The third GIG gene, \( lpg2107 \), encodes a 284-aa protein assigned to the DUF692 family of uncharacterized bacterial proteins. Other members of this family are key enzymes in the biosynthesis of methanobactins, secreted copper-binding and copper-reducing peptides produced by a variety of bacteria [28]. The final gene, \( lpg2108 \), encodes a small 97-aa protein assigned to the DUF2282 family of putative integral membrane and signal peptide proteins.

**Characterization of homologous operons**

The *L. pneumophila* str. Philadelphia 1 genome also contains three additional regions with significant sequence similarity to the GIG operon, designated here as homologous operons H2, H3 and H4 (Table 2). The H2 operon (\( lpg0665 \)-\( lpg0669 \)) is structurally similar to the GIG operon and contains homologs of all four GIG genes, arranged in an identical order and strand orientation. The H3 operon (\( lpg2253 \)-\( lpg2255 \)) lacks an \( lpg2105 \) homolog and therefore contains only three of the four GIG genes, again arranged in identical order and strand orientation. The H4 operon (\( lpg0671 \)-\( lpg0676 \)) contains homologs of all four GIG genes, but in a different order and on different strands. The first H4 gene (\( lpg0671 \)) shares a DoxX domain with \( lpg2105 \) but encodes a much larger NADH dehydrogenase transmembrane protein. Two additional genes are also present: an acetoacetate decarboxylase gene (\( lpg0672 \)) and an adenylate cyclase gene (\( lpg0674 \)). Rather than an operon, H4 may be better described as a cluster of genes that includes the four GIG homologs. They are unlikely to be transcribed as a unit, given their differing strand orientations.

We surveyed the genomes of four additional *L. pneumophila* strains for the presence of the GIG, H2, H3 and H4 operons (Table 2). While all strains contained copies of H2, H3 and H4, the GIG operon was present only in *L. pneumophila* Philadelphia 1. This is consistent with its acquisition as part of the ICE-\( \beta \)ox. A maximum likelihood phylogenetic tree indicated that the *L. pneumophila* GIG operon was most similar to the *C. metallidurans* GIG operon, as well as to number of homologous operons from multiple environmental and/or pathogenic bacteria (Fig 1).

**\( lpg2105-2108 \) operon**

The four genes—\( lpg2105-2108 \)—are predicted to be transcribed as a single unit based on genomic analysis. Using reverse transcriptase PCR, we confirmed that the genes are co-transcribed.
Using three sets of primers spanning the end of one gene and the beginning of the next, RT-PCR reactions for gene combinations 2105–2106, 2106–2107 and 2107–2108 were positive (Fig 2).

### GIG promoter activity in planktonic cultures

Expression of the *C. metallidurans* GIG operon is influenced by the presence of gold and copper ions in the growth medium [16]. To test whether the *L. pneumophila* GIG operon was similarly affected, we created a GFP reporter construct whose expression was driven by the 180nt upstream promoter region of the *L. pneumophila* GIG operon. We measured GFP fluorescence to monitor expression of the pGIgGfp reporter in *L. pneumophila* Lp02 exposed to multiple concentrations of gold, copper, and silver ions. At both 37˚ and 26˚C, we observed an extended lag phase in a concentration dependent manner when HAuCl4 or CuSO4 was added to the cultures. Cultures grown in the presence of 50 μM of gold or copper never reached the same maximum OD\textsubscript{600} as controls.

To account for the individual growth rates of each culture, mid-exponential (ME) and late stationary (LS) growth phases were identified in individual cultures, and the promoter activity was compared at each growth phase. Late stationary phase was defined as the time point prior to decline of OD\textsubscript{600} that preceded death of the culture. GFP was normalized to OD\textsubscript{600} at each time point to accurately compare promoter activity across cultures.

At 37˚C, control cultures of *L. pneumophila* Lp02 pGIgGfp showed similar levels of lpg2105-2108 promoter activity across all growth phases. The addition of 50μM AgNO\textsubscript{3} did not affect

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**Table 2. Homologous operons in *L. pneumophila* genomes to lpg2105-2108 GIG operon.**

| Strain  | GIG   | H2      | H3      | H4      |
|---------|-------|---------|---------|---------|
|         | Start position and strand | Start position and strand | Start position and strand | Start position and strand |
| Philadelphia | lpg2105 | 2352970 (-) | lpg0665 | 716610 (-) | lpg0671 | 719987 (+) |
| NC_002942.5 | lpg2106 | 2353739 (-) | lpg0666 | 717386 (-) | lpg2253 | 2557140 (-) | lpg0675 | 725484 (-) |
|          | lpg2107 | 2364586 (-) | lpg0667 | 718248 (-) | lpg2254 | 2558033 (-) | lpg0676 | 726280 (-) |
|          | lpg2108 | 2354894 (-) | lpg0669 | 718508 (-) | lpg2255 | 2558333 (-) | lpg0673 | 723047 (+) |
| Paris    | LPP_RS03625 | 801134 (-) | LPP_RS03650 | 804510 (+) |
| NC_006368.2 | LPP_RS03630 | 801910 (-) | LPP_RS03670 | 809982 (-) |
|          | LPP_RS03635 | 802772 (-) | LPP_RS03675 | 810802 (-) |
|          | LPP_RS03640 | 803032 (-) | LPP_RS03660 | 807569 (+) |
| Corby    | LPC_RS03790 | 827443 (-) | LPC_RS03815 | 830819 (+) |
| NC_009494.1 | LPC_RS03795 | 828003 (-) | LPC_RS03835 | 836292 (-) |
|          | LPC_RS03800 | 829080 (-) | LPC_RS03840 | 837112 (-) |
|          | LPC_RS03805 | 829340 (-) | LPC_RS03825 | 833879 (+) |
| Lens     | LPL_RS03540 | 790101 (-) | LPL_RS03565 | 793477 (+) |
| NC_006369.1 | LPL_RS03545 | 790877 (-) | LPL_RS03585 | 798950 (+) |
|          | LPL_RS03550 | 791739 (-) | LPL_RS03590 | 799770 (-) |
|          | LPL_RS03555 | 791999 (-) | LPL_RS03575 | 796537 (+) |
| Alcoy    | LPA_RS03750 | 813482 (-) | LPA_RS03775 | 816859 (+) |
| NC_014125.1 | LPA_RS03755 | 814258 (-) | LPA_RS03795 | 822332 (-) |
|          | LPA_RS03760 | 815120 (-) | LPA_RS03800 | 823152 (-) |
|          | LPA_RS03765 | 815380 (-) | LPA_RS03785 | 819919 (+) |

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culture growth or induce a promoter response. 150μM AgNO₃ inhibited culture growth but did not induce a promoter response (Fig 3). The promoter response to HAuCl₄ and CuSO₄ differed between growth phases. At the mid-exponential growth phase, only CuSO₄ treatment induced moderate promoter expression. At the late stationary growth phase, both HAuCl₄ and CuSO₄ induced higher levels of promoter expression (Fig 3).

At 26°C, L. pneumophila Lp02 growth was inhibited by the 20μM HAuCl₄ used at 37°C, and HAuCl₄ concentrations were therefore reduced to 5μM and 10μM. Legionella demonstrates increased resistance to copper at lower temperatures [9], and 26°C cultures showed no inhibition to the 20μM and 50μM CuSO₄ used at 37°C. CuSO₄ concentrations were therefore increased to 200μM and 275μM at 26°C to assess the promoter response under similar growth kinetics.

The effect of growth phase on promoter expression was more pronounced at 26°C than at 37°C. Control cultures showed a significant increase in promoter activity at LS phase, a result which was not observed at 37°C and that suggested temperature regulation of the operon. 10μM HAuCl₄ caused a modest increase in promoter activity at ME but reduced the level of promoter activity at LS. The addition of copper caused a significant decrease in promoter activity at LS (Fig 4).
**Fig 2. Reverse Transcriptase PCR using gene spanning primers.** M-Marker; 1: positive control 2105F/2106R primers; 2: positive control 2106F/2107R primers; 3: positive control 2107F/2108R primers; 4: no template control; 5: no RT control; 6: RNA 2105F/2106R primers; 7: RNA 2106F/2107R primers; 8: RNA 2107F/2108R primers.

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**Fig 3. Effect of gold, copper and silver on pGIGgfp activity in *L. pneumophila* at 37°C.** GIG operon activity measured by GFP expression of *pGIGgfp* in control media, with 20μM HAuCl₄, 50μM HAuCl₄, 20μM CuSO₄ and 50μM CuSO₄ at mid-exponential (ME) and late stationary (LS) growth phases (a). GFP expression in response to 50μM and 150μM AgNO₃ (b). Data presented from three independent experiments ± SD (* = p < 0.05).

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The complex promoter response to temperature, growth phase and metal addition suggests that multiple levels of operon regulation interact in planktonic culture. To further investigate mechanisms of operon regulation, we examined promoter expression in *E. coli* DH5α transformed with pGIGgfp as well as in two *L. pneumophila* mutants in the major stationary phase regulation pathways. *E. coli* DH5α, which does not possess the operon, showed no change in growth kinetics in the presence of HAuCl₄ or CuSO₄. At neither 37˚C nor 26˚C did *E. coli* show promoter activity to gold or copper (Fig 5). These results suggest a unique regulator in *L. pneumophila* for metal induction of the GIG operon. Significant increase in promoter expression in response to HAuCl₄ and CuSO₄ was seen during stationary phase, and we therefore investigated known regulators of *L. pneumophila* stationary phase gene expression. Mutants for the stationary phase regulators LetA and RpoS were transformed with pGIGgfp to assess their potential involvement.

The *Legionella* two-component regulatory system, LetA/S, induces the expression of virulence and transmission traits at stationary phase [29–30]. Lp02 ΔletA pGIGgfp cultures showed delayed entry into exponential phase when exposed to HAuCl₄ or CuSO₄, similar to that seen in wild type Lp02 pGIGgfp cultures. Lp02ΔletA cultures with added HAuCl₄ or CuSO₄ exhibited greater promoter activity compared to Lp02 ΔletA cultures with no metal addition. The magnitude of metal-induced promoter up-regulation was greater at the LS phase. However, wild type and ΔletA cultures responded similarly to metal addition: there were no significant differences in promoter activity between Lp02 ΔletA cultures and wild type Lp02 cultures at the same metal concentration.
Lp02 ΔrpoS pGIGgfp cultures exhibited an extended lag phase in response to HAuCl$_4$ or CuSO$_4$, and 50 μM CuSO$_4$ cultures showed increased promoter expression at ME phase. The promoter response to metal addition was dampened in Lp02 ΔrpoS pGIGgfp stationary phase cultures: mutant cultures showed reduced promoter activity compared to wild type in response to both HAuCl$_4$ and CuSO$_4$ addition, suggesting that RpoS may play a role in promoter expression, particularly at stationary phase (Fig 6).

GIG promoter activity in biofilms

*L. pneumophila* primarily persists in the environment as a biofilm [31–33]. To assess operon activity under more natural growth conditions, Lp02 pGIGgfp biofilms were established and grown at two temperatures in the presence and absence of 20 μM HAuCl$_4$, 50 μM HAuCl$_4$, 20 μM CuSO$_4$, or 50 μM CuSO$_4$ at mid-exponential (ME) and late stationary (LS) growth phases. Data presented are from three independent experiments ± SD.

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![Fig 5. Effect of gold and copper on pGIGgfp activity in *E. coli* at 37°C. GIG operon activity measured by GFP expression of pGIGgfp in control media with addition of with 20 μM HAuCl$_4$, 50 μM HAuCl$_4$, 20 μM CuSO$_4$, or 50 μM CuSO$_4$ at mid-exponential (ME) and late stationary (LS) growth phases. Data presented are from three independent experiments ± SD.](https://doi.org/10.1371/journal.pone.0174245.g005)

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Similar to planktonic cultures, greater promoter expression was measured in 37°C biofilms than in 26°C (Fig 7b). Increased promoter expression was seen in early phase biofilms in response to gold and copper at 24h and in response to gold at 48h and 72h.

To determine if biomass were responsible for the increase in promoter activity, biofilm biomass was assessed from confocal images using COMSTAT. There were no differences in
biomass among control, plus gold, and plus copper samples at each temperature. There was a significant decrease in biofilm biomass in all treatments from 48h to 72h, likely due to the dispersal event typically seen at this stage in *Legionella* biofilm development (Fig 8).

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**Fig 6.** Effect of gold and copper on pGIGgfp activity *L. pneumophila* Lp02 ΔletA and ΔrpoS. GIG operon activity measured by GFP expression of pGIGgfp in control media with addition of 20μM HAuCl₄, 50μM HAuCl₄, 20μM CuSO₄, or 50μM CuSO₄ at mid-exponential (ME) and late stationary (LS) growth phases. Data presented are from three independent experiments ± SD. (*) = significantly different from WT with same metal addition in same panel; p < 0.05, † = significantly different from mutant with no metal addition in first panel; p<0.05

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**Fig 7.** Analysis of pGIGgfp expression during biofilm establishment and development at a) 26°C in *L. pneumophila* control biofilms and with the addition of 20μM HAuCl₄ or 20μM CuSO₄ and at b) 37°C in *L. pneumophila* control biofilms and with the addition of 20μM HAuCl₄ or 20μM CuSO₄. Percent GFP intensity above basal level (determined at 26°C) is shown. Data presented are from three independent experiments ± SE (p<0.05)

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Discussion

Based on sequenced strains, the GIG operon appears to be rare in the *Legionella* genus, existing in only 3 sequenced strains—Philadelphia 1, Lp02 and SVir, each of which possess the full operon. Multiple homologs of the operon exist in *L. pneumophila* at genes lpg0665-0669, lpg0671, 0673, 0675, 0676, and lpg2253-2255. Species within the *Legionella* genus appear to contain multiple paralogous copies of the operon, which group into four distinct clades (Fig 1, Table 1). The lpg2105-2108 operon in *L. pneumophila* Philadelphia 1 is distinct from the others.

The response of the *L. pneumophila* GIG operon to HAUCl$_4$ and CuSO$_4$ is similar to that reported for the GIG operon of *C. metallidurans* [16]. At 37˚C, planktonic cultures expressed GIG operon promoter activity that increased with the addition of HAUCl$_4$ or CuSO$_4$ at stationary phase. Conversely, promoter activity was repressed in response to HAUCl$_4$ or CuSO$_4$ at 26˚C (Fig 4). Although not well explored, temperature regulation of genes is not unknown in *Legionella*. Piao, et al., [34] showed that biofilms alter their morphology based on temperature and surface material. Biofilms formed at 37˚C and 42˚C resembled filamentous mycelial mats, while at 26˚C, short, rod shaped bacteria made up the majority of the biofilm. Genes involved in type IV pilus biogenesis and type II protein secretion also showed temperature regulation, with increased expression at 30˚C compared to 37˚C [35]. The increased HAUCl$_4$ sensitivity and opposite expression patterns between 26˚C and 37˚C in both biofilm and planktonic cultures suggest that temperature regulation may be involved with operon activity. *Legionellae* growing at low-temperatures have a more unsaturated membrane lipids, suggesting that
changes in membrane fluidics may also be involved in gold sensitivity if membrane proteins are involved in the response to gold [36]. The lpg2107 gene appears to possess transmembrane domains and may therefore be localized to the bacterial membrane.

Stationary phase changes in promoter activity in both Legionella and E. coli suggest that known stationary phase regulatory proteins may regulate the operon. L. pneumophila exhibits unique biphasic gene expression with tight control over exponential vs. stationary phase proteins. Determinants involved in metal or metalloid resistance that are up-regulated after treatment with gold complexes are often controlled by MerR- or ArsR-type regulators, which usually bind “soft” metals or metalloids. In addition to C. metallidurans, gold response systems exist in E. coli, controlled through CueR [37] and in S. typhimurium controlled by GolS [38]. C. metallidurans also possesses a CupR system that responds to both gold and copper [39]. Homologs of CueR and GolS are found in the L. pneumophila genome but have not been characterized.

Previous transcriptomic work showed expression of the Legionella GIG operon during amoebal infection, suggesting a potential role in virulence [40]. Many Legionella virulence genes are regulated by the rpoS, csrA, and letA genes [41–43]. LetA/S activates transmission phase genes, including but not limited to mip (macrophage infectivity protein), dot/icm T4SS, and flaA [29, 43, 44–45]. Deletion of letA significantly inhibits virulence [29]. However, we saw no changes in promoter activity in Lp02 ΔletA pGIGgfp compared to WT Lp02 pGIGgfp, suggesting that LetA does not directly regulate expression of the operon. A bioinformatic search for the conserved LetA promoter binding sequence upstream of the GIG operon was also negative, further supporting a lack of direct LetA involvement in operon regulation.

Lp02 ΔrpoS cultures showed no response to added HAuCl₄ or CuSO₄ when compared to the ΔrpoS control. Compared to the Lp02 WT however, promoter activity in ΔrpoS strains was significantly repressed (Fig 6). RpoS typically represses motility, infectivity, and cytotoxicity during exponential phase, and up-regulates them during stationary phase [46–47]. It is required for expression of virulence traits and growth within amoebae [41]. RpoS functions in response to stressors in the environment and interacts with other stationary phase regulators such as LetA/S, FliA, letE, and LqsR [47–48]. Our data suggest that RpoS may regulate the GIG operon similarly to the way it regulates other stationary phase virulence genes. In fact, regulation may require LetA to release exponential phase repression by RpoS and allow for stationary phase expression, as seen in the regulation of the sRNA molecules RsmYZ [44]. The data support the possibility of multiple regulators for this operon, one for basal expression, and a second for gold/copper response. Investigations into mutants of fliA and letE, which also coordinate differentiation from replicative phase to transmissive [29,49], as well as csrA, a global regulator responsible for the repression of transmission phase genes and activation of replicative phase genes [42], may lead to a better understanding of how this operon is controlled. The recent publication of a csrA homolog within the ICE-βox unit–csrT–is also a potential regulatory candidate [50].

Biofilms grown at 26˚C with 20μM gold or copper showed increased response to the metal ions compared to planktonic cultures at 26˚C, with no obvious toxicity. Biomass of biofilms exposed to copper and gold was equivalent to controls (Fig 8). The matrix produced by biofilms that protects bacteria from the effects of toxic metals is not found in planktonic cells. Analyses of operon expression through multiple stages of biofilm development revealed that while baseline expression is present in controls, HAuCl₄ or CuSO₄ treatments increase promoter activity, particularly at early developmental stages. Specific up-regulation of the promoter in response to gold occurred at 24h and 72h. Up-regulation early in biofilm development suggests that after the matrix is in place, there is less need for metal response system expression. The up-regulation in response to gold at 72h corresponded with the timing of initial dispersal events in this model (Fig 8). Biofilm dispersal is evident in the difference in
biomass between 48 and 72h, but a significant increase in promoter activity is seen at 72h. The up-regulation at this time point may be from cells newly exposed to the external environment.

The GIG operon is present in several other virulent pathogens, including a category B biological agent *Burkholderia pseudomallei*, the causative agent of melioidosis, and a Category A bioterrorism agent, *Francisella tularensis*, the causative agent of tularemia. The presence of this operon in *F. tularensis* is particularly interesting since its abundance of pseudogenes and disrupted metabolic pathways are indicative of genome reduction [51–52]. The idea of "use it or lose it" can be applied to this process: genes that are nonessential to the survival of the organism that are mutated or lost. That this operon has been retained suggests it may have an essential function in survival or environmental persistence that could be exploited as a target for antimicrobials.

Many studies have been conducted on the efficacy of water treatment methods on *Legionella*, but few have looked at possible genetic response systems to those treatments, specifically those involved with metal ion response. The controversy over the use of copper/silver ionization as an effective means to protect water systems from *Legionella* colonization may be better explained after the metal response systems in the bacteria are more thoroughly characterized. Lack of information on bacterial response to metals hinders development of effective disinfection strategies. Based on the persistence of *Legionella* in the presence of metal-based disinfection treatments, an increased understanding of biofilm ecology, and in particular the environmental ecology of *Legionella*, is necessary. The results of this study lay groundwork for continued investigation of *Legionella* responses to potentially toxic metal ion concentrations.

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