Introduction

Campylobacter jejuni is the most common cause of campylobacteriosis, one of the most common gastrointestinal diseases in humans in the US [1]. It is a food-borne, Gram-negative, motile microaerophilic bacterium that causes disease by invading the epithelial cells of the host’s colon through subversion of microtubule structures in the epithelial cells [2,3]. Damage to these cells disrupts the normal absorptive function of the colon and causes disease [3,4]. Disease progression typically consists of abdominal pain, watery or bloody diarrhea, and fever which persists up to a week [4,5]. Most cases of campylobacteriosis are self-limiting, but medical costs and losses of productivity total more than 2.4 billion a year [6]. Additionally, approximately 1/1000 clinical cases may result in long term neurological defects, including Guillain-Barre syndrome [7,8]. This is particularly significant in immune-compromised populations, which comprise an estimated 3.6% of the total population in the US, who are more likely to become ill from Campylobacter and also more likely to experience long term sequelae [9,10]. Further, the elderly population, which is likely to represent more than 20% of the US population in the next few decades, are also at increased risk for food-borne illnesses; including Campylobacter [11–13]. Therefore Campylobacter infections may become both more prevalent and severe as the population ages, making understanding Campylobacter cell physiology and controlling this pathogen urgently relevant.

Despite its success as an endemic gastrointestinal pathogen, the keys to Campylobacter’s virulence remain enigmatic. Unlike other enteric bacterial pathogens, Campylobacter does not have the advantages of a Type III secretion system, toxins, or other conventional pathogenicity factors [14]. Campylobacter is not known to proliferate outside of the host, but does adapt quickly to environmental stressors including osmotic pressure, nutrient starvation and antibiotic exposure [15]. One of the factors necessary for stress-survival and successful colonization is the Twin Arginine Translocation (TAT) system [16]. The TAT system is an inner membrane translocase that transports proteins folded in the cytoplasm across the inner membrane. Although the TAT system is ubiquitous among bacteria and archaea, it does not have any animal homologues [17].
Using in silico screens, we have previously hypothesized that Campylobacter transports over a dozen proteins through TAT system [16]. Several proposed TAT substrates have been confirmed by laboratory methods, including HlyB, MrIB, and CJ0415, MrLA, TorA, CsoA, FdhA, the YedY homologue CJ0379c, and PhoX [19–20]. Some of these proteins are thought to have a role in cell physiology including stress survival, and we have previously found that the C. jejuni ΔattC mutant is unable to establish long term colonization in chickens [16]. The mechanism of this survival defect, however, has not been well-studied.

One of the proteins that has been identified as a TAT substrate is PhoA(CJ), the only alkaline phosphatase in Campylobacter species [21]. The PhoA(CJ) requires transport into the periplasm to become active and there it provides Campylobacter with inorganic phosphate (P1) through the hydrolysis of phosphate groups from more complex organophosphate molecules. Also, the regulation of phosphate sensing and alkaline phosphatase activity in Campylobacter species is thought to have evolved separately from those in the more commonly studied Escherichia coli and Bacillus subtilis, and lacks some of the features common to these bacteria, including auto-regulation [20]. The Campylobacter alkaline phosphatase is more similar to those found in Vibrio cholera, Pseudomonas, as well as marine and soil bacteria [22,23]. This family of alkaline phosphatases (PhoX) are typified by cytoplasmic folding, divergent sequences with little homology, and functional dependence of Ca2+ in place of Mg2+ or Zn2+ [23]. Since recent research has shown persuasively that Campylobacter alkaline phosphatase has evolutionary and functional similarities to the PhoX family of alkaline phosphatases, here we will refer to PhoA(CJ) as PhoX [21–25].

Since Pi in the environment is typically low, alkaline phosphatases are necessary for nutrient and ATP homeostasis [24]. Pi is necessary for the PPK1 mediated formation of poly P, which has an important role in both basic metabolism and stress response [25]. Poly P has been shown to insulate cells in an alkali environment and be necessary for long term growth and survival in Salmonella [25,26]. Additionally, cellular phosphate levels are often used by bacteria as an indicator of environmental resources; cellular responses to phosphate levels are closely connected to the control of starvation response as well as flagella growth, quorum sensing, and the production of virulence factors [24,27,28]. Many aspects of the Campylobacter phosphate response are mediated by the two component system phoR/S, a homologue to the E. coli phoR/B system [20]. In a microarray comparison of phosphate-starvation response, wild-type cells showed transcriptional modification of over 200 genes [29]. While some of these genes are likely affected by secondary responses, it is clear that the PhosR/S regulon is likely similarly important to cellular metabolism. Genetically similar phoX genes found in V. cholerae have been shown to be involved in environmental survival mechanisms including control of biofilm formation, adjustment of cyclic-di-GMP levels, aerobic and heat-stress tolerance, stringent response as well as flagella function [24,30–32]. Inorganic phosphate is known to also regulate additional genes including spoT, which subsequently affects the global regulator ppGpp [33]. Therefore, we hypothesize that some of the basic stress response defects seen in the Campylobacter TAT knockout mutant are caused by the inhibition of PhoX transport. We also hypothesize that these phenotypes may be linked with the subsequent interruption of poly P and poly P mediated stress responses. For both a way to understand the underlying physiology of an important pathogen as well as potential investment into understanding a mechanism that can be exploited for therapeutic use, we determined the role of phoX in C. jejuni pathophysiology.

### Materials and Methods

#### Ethics statement

Animal experiment was conducted according to the guidelines of Association for Assessment and Accreditation of Laboratory Animal Care International (AAALAC). The animal studies are approved by The Institutional Animal Care and Use Committee (IACUC), The Ohio State University, under the protocol number 2010A00000155. As we study complex host pathogen interactions that are applicable to human health, the use of laboratory animals is unavoidable and justifiable. Chickens were housed at the Food Animal Health Research Program Animal Care Facility. The facility is fully accredited by AAALAC and animals were supervised by our senior veterinarian Dr. J. Hanson. Infectious agents were administered using manual restraint for less than one minute to minimize distress. Chickens were euthanized by carbon dioxide inhalation, which is rapid and painless. This method is consistent with the recommendations of the panel on euthanasia of the American Veterinary Medical Association and by The Ohio State University Institutional Laboratory Animal Care and Use Committee.

#### Bacterial strains, plasmids, and culture conditions

Bacterial strains and plasmids used in this study are listed in Table S1. C. jejuni strain 81–176 (WT), a highly invasive strain originally isolated from an outbreak associated with raw milk [34], was used to generate the phoX deletion mutant. C. jejuni strains were routinely grown on Mueller-Hinton broth (MH; Oxoid) microaerobically [85% N2 (v/v), 10% CO2 (v/v) and 5% O2 (v/v)] in a DG250 Microaerophilic Workstation (Microbiology International) at 42°C. MH agar plates were supplemented with Campylobacter selective supplement (SR117E, Oxoid) when isolating C. jejuni from chicken feces and organs. For growth curve and stress survival assays, C. jejuni was grown microaerobically in MH broth with appropriate antibiotics at 42°C with shaking at 200 rpm. E. coli DH5α was used for plasmid propagation and cloning purposes and was routinely cultured on Luria-Bertani (LB) medium at 37°C overnight. Growth media was supplemented with appropriate antibiotics, chloramphenicol (20 µg ml−1 for E. coli, 10 µg ml−1 for Campylobacter), kanamycin (30 µg ml−1) and zeocin (50 µg ml−1), where necessary.

#### Construction of the phoX deletion mutant

Cloning and other molecular biology techniques were performed according to Sambrook & Russell [35]. Oligonucleotides were designed using Vector NTI® software (Invitrogen, Carlsbad, CA) and commercially synthesized by Integrated DNA Technologies (Skokie, IL). All the oligonucleotides used in the present study are listed in Table S2. Masterpure® DNA purification kit and FastLink DNA ligation kit were purchased from Epicentre (Madison, WI). Restriction enzymes were purchased from Promega. QIAquick® PCR purification kit and QIAprep® spin mini prep kit for plasmid isolation were purchased from Qiagen (Valencia, CA). Zero background cloning vector pZero1-O1 and E. coli DH5α competent cells were purchased from Invitrogen.

Deletion of the phoX gene (CjJ_0181) was achieved by double crossover homologous recombination using a suicide vector containing approximately 1 kb of homologous sequences on either side of the phoX gene as described previously [16]. Briefly, the phoX along with 1 kb flanking region on either side of the target gene was amplified by PCR using PhoX-F and PhoX-R primers from C. jejuni 81–176 genomic DNA. The amplified PCR product was ligated into pZErO-1 to generate plasmid pZero1-phoX. Inverse PCR was performed on pZerol-phoX using PhoX INV-F and...
PhoX INV-R primers to delete majority of the phoX coding sequence. Kanamycin cassette from pUC4K was then cloned into inverse PCR product, the resulting suicide vector designated, pZero1-phoX, was electroporated into C. jejuni 81–176 as described [16,36]. Recombinants were selected on MH agar plates containing kanamycin, kanamycin resistant colonies were streak purified and one such mutant designated AphoX was used for further studies. The deletion of the phoX gene was confirmed by PCR.

**Complementation of the AphoX mutant**

The complementation of AphoX mutant was accomplished by the insertion of a wild type phoX using pRRC integration vector [37]. The coding region of phoX along with its ribosome binding site was amplified by PCR using primers PhoX COMP-F and PhoX COMP-R. Restriction site (BamHI) was included in each primer to facilitate cloning. Following digestion with BamHI, product was ligated into the pRRC vector. The resulting construct was electroporated into C. jejuni ΔphoX::cin, and putative complemented clones were recovered on plates containing kanamycin and chloramphenicol. Insertion of the phoX gene in the mRNA spacer region was confirmed by PCR and further the constitutive expression of phoX was confirmed by phosphatase assay. One of the confirmed complemented clone designated phoX was used for further analysis.

**Alkaline phosphatase assay**

Alkaline phosphatase activity was determined as previously described [20]. Briefly, the ΔphoX, ΔphpK1, ΔtatC, ΔphoXc, and WT 81176 strains were grown overnight on MH plates with appropriate antibiotic selection. The cultures were gently scraped, washed in minimal essential medium (MEM), resuspended in MEM and incubated at 42°C microaerobically with shaking for 2 hours. Cultures were then centrifuged for 10 minutes at 7000 × g and supernatant was removed. Cells were gently washed with 50 mM MOPS buffer (pH 7.4) (Sigma), OD600 was measured. The cells were pelleted, supernatant was removed, and cells were resuspended in PNPP buffer with 2 mM PNPP (Sigma) and 50 mM MOPS buffer (pH 7.4) (Sigma), OD600 was measured. The coding region of phoX was amplified along with its ribosome binding sequence. Kanamycin cassette from pUC4K was then cloned into PhoX INV-R primers to delete majority of the phoX coding sequence. Kanamycin cassette from pUC4K was then cloned into pRRC vector. The resulting construct was electroporated into C. jejuni ΔphoX::cin, and putative complemented clones were recovered on plates containing kanamycin and chloramphenicol. Insertion of the phoX gene in the mRNA spacer region was confirmed by PCR and further the constitutive expression of phoX was confirmed by phosphatase assay. One of the confirmed complemented clone designated phoX was used for further analysis.

**Isolation/Detection of poly P**

Poly P was extracted using glassmilk and quantified using toluidine blue O as described earlier [39]. Poly P was quantified from mid-log, and stationary phase cultures by measuring the absorbance ratio at 530 to 630 nm spectrophotometrically using appropriate concentrations of phosphorous standard (Sigma Aldrich). The experiment was performed a total of three times under rich media (MH) and minimal media (MEM) conditions.

**Nutrient downshift assay**

The role of phoX in C. jejuni survival under nutrient downshift was assessed using MEM with glutamine (Gibco 11095) or without glutamine (Sigma, M2279) and in the presence or absence of 1 mM Pi (Inorganic Ventures) as described previously [40]. In experiments where Pi was added to the media, the final concentration was 1 mM. This amount is similar to a previous study by Wosten et al [20] where concentrations above 0.4 mM show greater than 90% inhibition of alkaline phosphatase activity. However alkaline phosphate is induced in MEM which has low Pi concentration. Therefore, here we used MEM and 1 mM Pi. Additionally 1 mM Pi concentration had no effect on C. jejuni growth [20]. Briefly, mid-log-phase cultures of the WT, ΔphoX, and ΔphoXc, strains were pelleted, washed twice and resuspended in MEM with or without glutamine or Pi, and the OD600 was adjusted to 0.05. The suspensions were then incubated microaerobically at 42°C with shaking at 200 rpm. At different time points, 100 μl of cultures were serially diluted (10-fold) in respective MEM media and plated onto MH agar in triplicate. The plates were incubated microaerobically, and the number of CFU ml⁻¹ was calculated. The experiment was performed three times.

**Osmotic stress response assay**

Survival of C. jejuni phoX deletion mutant in the presence of osmotic stress was tested as described previously [39,40]. To assess the osmotic stress survival in liquid culture, bacterial strains were grown to mid-log phase, adjusted to an OD600 of 0.05 in MH broth with and without 0.25 M NaCl and incubated microaerobically at 42°C for 48 hours with shaking at 200 rpm. A 100 μl of the culture at different time points was serially diluted (10-fold) and plated on MH agar plates. The plates were incubated microaerobically and CFU were determined. To assess the strains ability to tolerate osmotic stress on a solid medium, the WT, ΔphoX, and ΔphoXc strains were grown to mid-log phase, serially diluted (10-fold), and 10 μl of diluted culture was spotted onto MH agar plates containing either 0.17 M NaCl or 0.17 M NaCl and 1 mM Pi. Plates were incubated microaerobically at 42°C for 2 days, the growth of C. jejuni was visualized and photographed. The experiment was repeated three times.

**Oxidative Stress response assay**

To determine oxidative stress response, WT, ΔphoX, and ΔphoXc strains were grown overnight on MH agar with the appropriate antibiotics at 42°C under microaerobic conditions. Cells were harvested the next day and 100 μl of bacterial culture containing 5×10⁶ CFU ml⁻¹ (OD600 of 0.5) were spread on MH plates with appropriate antibiotics in the presence or absence of 1 mM Pi. A 5 mm well was cut into the middle of each plate and filled with 30 μl of 20 mM paraquat or 0.3% H₂O₂. Plates were incubated for 24 and 48 hours under microaerobic conditions and the zone of inhibition was measured and photographed. The experiment was performed a total of three times.

**Motility and biofilm assays**

The C. jejuni wild type 81–176, ΔphoX, ΔphoXc, Δppk1, and Δppk1c strains were tested for motility in semisolid MH medium plates containing 0.4% agar in the presence or absence of 1 mM Pi. Cultures were grown on MH agar plate under microaerobic conditions at 42°C for 48 hours. Culture densities were adjusted to OD₆₀₀ of 0.05 and 2 μl of each culture was stabbed onto the surface of the motility plate. Plates were incubated at 42°C under microaerophilic conditions. Motility phenotypes were assessed after 24, and 48 hours, following inoculation. The motility assay was performed three times.

Static biofilm formation was assessed in borosilicate tubes as described previously [40,41]. Briefly, overnight grown cultures of C. jejuni strains were diluted in MH broth with or without Pi to an OD₆₀₀ of 0.05. Two milliliters of diluted culture was incubated at 42°C microaerobically for 3 days without shaking. Biofilms were visualized by staining with 250 μl of 1% (w/v) crystal violet for 15 min, rinsed 3 times with double distilled water. After third rinse, vials were photographed then quantified by measuring the absorbance at 570 nm after dissolving in 2 ml DMSO for 24 hours. The biofilm assay was performed three times.
Antimicrobial susceptibility testing

Susceptibility to azithromycin, ciprofloxacin, erythromycin, tetracycline, fleroxacin, nalidixic acid, telithromycin, clindamycin, and gentamicin was determined by using Sensititre® susceptibility plates for Campylobacter (TREK Diagnostic Systems, West Sussex, UK). Briefly, one hundred microliters of log-phase grown cultures adjusted to an OD<sub>600</sub> of 0.05 in MH broth was added to each well in the Sensititre® susceptibility plate and the wells were covered using the perforated adhesive seal. The plates were incubated microaerobically at 42°C for 24 hours and the minimum inhibitory concentration (MIC) was recorded. Results were read following the manufacturer's instructions and interpreted according to MIC interpretive guidelines by Clinical Laboratory Standards Institute. In addition, the following antimicrobials were also tested individually as described previously [16]: polymyxin B, cholic acid, taurocholic acid, deoxycholic acid, and an antimicrobial peptide of chicken origin, fowlacidin-1. One hundred microliters of the cultures grown above were added to serially diluted (2-fold) antimicrobials in a 96 well microtiter plate, mixed and the plates were incubated microaerobically at 42°C for 24 hours. The MIC was determined as the lowest concentration showing complete inhibition of visible growth. The susceptibility testing was repeated 3 times and the mean MIC (µg ml<sup>-1</sup>) was calculated.

Quantitative RT-PCR

Quantitative RT-PCR (qRT-PCR) was performed, targeting a key gene involved in stringent response, spoT; post transcriptional global regulator, ccrA; poly P associated enzymes, ppk1 and ppk2; phosphate metabolism, phoR, pstS, pstC; oxidative stress, pq0379, cgg1374, abpC, sodB, and the multdrug resistance efflux pump gene, cmeC [20,42,43,44]. Total RNA was extracted from log-phase grown bacterial cultures using RNeasy Mini Kit (Qiagen). The RNA concentration and purity was determined using NanoDrop ND-1000 spectrophotometer (Wilmington, DE). cDNA synthesis was carried out using SuperScript® III First-Strand Synthesis SuperMix (Invitrogen). Gene specific primers were designed to amplify the above mentioned genes along with atpA or 16SrRNA (internal controls) using Beacon Designer 7.0 (Palo Alto, CA). Primers were obtained commercially from IDT-DNA and are described in Table S2. qPCR was performed using SensiMixPlus® SYBR RT-PCR Kit (Quantrace, Norwood, MA) in a Realplex<sup>2</sup> Mastercycler (Eppendorf, Westbury, NY). The relative levels of expression of genes were normalized with either atpA or 16SrRNA amplified from the corresponding sample. The difference in expression of the genes was calculated using the comparative threshold cycle (CT) method to yield fold-difference in transcript levels [45]. The qRT-PCR was performed in duplicates and assay was performed a total of four times.

INT407 invasion and intracellular survival assay

Invasion and intracellular survival assays were performed as described previously [39,46]. Each well of a 24 well tissue culture plate was seeded with 1.4×10<sup>5</sup> INT407 cells in MEM with 10% fetal bovine serum (FBS) and incubated for 18 hours at 37°C with 5% CO<sub>2</sub>. C. jejuni strains were grown to mid-log phase in MH broth microaerobically, the cells were pelleted at 5,000 g for 10 min, washed twice with MEM containing 1% FBS, resuspended in MEM to an OD<sub>600</sub> of 0.02 (1.5×10<sup>7</sup> cells) and used for infection. INT407 cells were infected with a multiplicity of infection 1:100 for invasion and intracellular survival assays. For infection, 1 ml of bacterial cell suspension was pipetted on to INT407 cells, centrifuged at 1000 g for 3 min and incubated for 3 hours. For determining invasion, after 3 hours of incubation with bacteria, cells were treated with gentamicin (150 µg ml<sup>-1</sup>) and incubated for additional 2 h. After 2 hours of incubation, the infected cells were rinsed with MEM three times, lysed with 0.1% (v/v) Triton-X 100, serially diluted in MEM and plated on MH agar in duplicate to determine CFU. To assess intracellular survival, following 2 hours of gentamicin treatment the infected cells were washed with MEM three times and covered with MEM containing gentamicin (10 µg ml<sup>-1</sup>) and incubated for 24 hours. After 24 hours of incubation, the infected cells were washed with MEM, lysed with 0.1% (v/v) Triton-X 100, serially diluted in MEM and plated on MH agar in duplicate to determine CFU. The invasion and intracellular assays were performed three times.

Chicken colonization study

Chicken colonization studies were performed as described previously [40]. Briefly, day-old broiler chicks (n=6 for each strain) were infected with Campylobacter cells at a MOI of 100 and allowed to recover for 24 hours, while being kept in isolation cages. The infected chicks were then killed by cervical dislocation and the entire gastrointestinal tract was removed. The content from the first and second segment of the small intestine was collected and homogenized in 0.1% (v/v) Triton-X 100, serially diluted in MEM and plated on MH agar in duplicate to determine CFU. The backgrounder was performed three times.

Figure 1. The phoX and tatC deletion mutants are defective in alkaline phosphatase activity. A PNP based phosphatase assay was carried out in triplicate to determine alkaline phosphatase activity. The ΔphoX and the ΔtatC mutants show similar inhibition of alkaline phosphatase activity where as the ppk1 mutant has alkaline phosphatase activity similar to wild type. Each data point is the mean ± standard deviation of 3 independent experiments. * P<0.05. ** P<0.01. doi:10.1371/journal.pone.0026336.g001

Figure 2. The phoX mutant is defective in poly P accumulation. Poly P was extracted from stationary phase grown wild type and ΔphoX mutant using glassmilk. The amount of poly P in the cell was determined by toluidine blue O method. Each data point is the mean ± standard error of 3 independent experiments. * P<0.05. ** P<0.01. doi:10.1371/journal.pone.0026336.g002
group) from a local hatching facility (Food Animal Health Research Program, OARDC, Wooster, OH) were inoculated orally with $10^3$ and $10^5$ CFU of the C. jejuni WT and ΔphoX strains in 200 μl of PBS (pH 7.4). After 7 days post-inoculation, the chicks were euthanized. The ceca and feces were collected aseptically, weighed, homogenized, serially diluted in PBS (pH 7.4) and plated on MH agar containing Campylobacter selective supplement. Plates were incubated at 42°C microaerobically and CFU per gram of tissues were determined.

Statistical analysis
Statistical significance of data generated in this study was determined using one-way analysis of variance (ANOVA) followed by Tukey’s HSD (Honestly Significant Difference) test or Student’s t-test (paired 2-tailed). $P<0.05$ (α level) was considered statistically significant.

Results
The ppk1 is not a primary effector of alkaline phosphatase activity
We found that ΔphoX mutant was significantly defective ($P<0.01$) in alkaline phosphatase activity (27.0 mU) compared to wild type (123.0 mU) (Fig. 1). Preliminary studies suggested that alkaline phosphatase activity could be measured with least background activity when cultures were incubated in MEM followed by washing with MOPS buffer before the assay (Fig. S1). Further, complementation of the ΔphoX mutant restored the alkaline phosphatase activity similar to wild-type levels (Fig. 1). Similarly, the ΔtatC mutant also showed similar alkaline phosphatase activity (33.5 mU) compared to the ΔphoX mutant (Fig. 1). This confirms previous results that PhoX is the only alkaline phosphatase in C. jejuni and it is solely transported through

Figure 3. PhoX is necessary for survival under nutrient stress conditions. (a) Survival of phoX deletion, wild type, and the phoX complemented strains under nutrient downshift was assessed by growing bacterial strains in minimal essential medium and determining the colony forming units at different time points. Addition of either 2 mM glutamine (b) or 1 mM Pi (c) or both (d) to nutrient downshift media improves phoX deletion mutant survival compared to wild type cells. Each data point represents the mean ± standard error of 3 independent experiments. * $P<0.05$.

doi:10.1371/journal.pone.0026336.g003
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PhoX in C. jejuni Environmental Resilience

Figure 4. Motility and biofilm formation in the ΔphoX mutant. (a) Motility of the ΔphoX mutant is similar to the motility of the wild type strain and addition of Pi did not affect the motility. Values indicate average motility zone diameter ± standard error from 3 replicate experiments. Only representative images are shown. (b) The ΔphoX mutant shows enhanced biofilm formation. The biofilm was visualized by staining with 1% crystal violet for 15 min. (c) Quantification of biofilm using dimethyl sulfoxide (DMSO). The amount of biofilm formed was dissolved in 2 ml DMSO for 24 hours and quantified by measuring absorbance at 570 nm. (d) Inorganic phosphate (1 mM) rescues the increased -biofilm phenotype in the ΔphoX mutants, but wild type biofilm is not changed by the addition of inorganic phosphate. The effect of 1 mM Pi on each strain was calculated as fold change in biofilm = (absorbance 570\text{nm} with addition of 1 mM Pi)/(absorbance 570\text{nm} without Pi) Each bar represents the mean ± SD of 3 independent experiments. * P<0.05. ** P<0.01.

doi:10.1371/journal.pone.0026336.g004

the TAT system [21]. The ΔattC mutant, though not statistically significant, showed slightly increased alkaline phosphatase activity compared to the phoX mutant.

It is well known that poly P levels are influenced by a wide range of stress responses, including low-nutrient stress [39]. Further, ΔspoT deletion mutants have shown both diminished accumulation of poly P, decreased transcription of ppk1 as well as increased transcription of pssS and pstC, which are regulated as part of the PhoR/S phosphate uptake regulon [39]. Therefore, we investigated whether there was a direct relationship between alkaline phosphatase activity and poly P accumulation. The alkaline phosphatase level in the ppk1 deletion mutant was 92.7 mU, slightly but significantly decreased compared to the wild-type (P<0.05) (Fig. 1). This suggests that ppk1 expression does not severely affect alkaline phosphatase expression; they are potentially part of a separate, although intersecting function, of the phosphate utilization pathway.

Poly P accumulation is reduced in both the tatC and phoX deletion mutants

The phoX deletion mutant showed a statistically significant decrease (P<0.05) in poly P accumulation during stationary phase in the minimal media compared to the wild-type. The poly P levels of ΔphoX mutant were similar to the poly P levels of the tatC deletion mutant (Fig. 2). Both ΔphoX and ΔattC had an average accumulated poly P level of approximately 21 nM poly P mg\(^{-1}\) of total protein and 25 nM poly P mg\(^{-1}\) of total protein respectively, compared to the wild-type cells (37 nM poly P mg\(^{-1}\) of total protein) (Fig. 2). However, ppk1 deletion mutant showed a larger accumulation defect with only 14 nM poly P mg\(^{-1}\) total protein (P<0.01) (Fig. 2). Difference in poly P accumulation between ppk1 and phoX mutants is likely due to ability of the phoX mutant to ameliorate the poly P defect with Pi obtained by other sources such as through phosphonate catabolism. C. jejuni has been shown to catabolize phosphonate [47,48]. However, ppk1 mutant can not synthesize poly P even in the presence of other Pi sources including phosphonates. This result suggests that interruption of phoX may be sufficient to cause the poly P defects seen in the tatC mutant as both ΔattC and ΔphoX mutants showed similar poly P levels. Although, the ΔattC mutant was defective in growth in rich media, the ΔphoX mutant grew similar to wild-type (data not shown, [21]). Similarly the poly P accumulation was significantly decreased in the phoX deletion mutant in the rich media during stationary phase (Fig S2). There was no difference in the accumulation of poly P between mutant and the wild-type in the log phase (data not shown). Although alkaline phosphatase activity is necessary for ppk1 mediated accumulation of poly P, the ppk1 mutant had only a small defect in alkaline phosphatase activity. Alkaline phosphatase activity may be pleiotropically mediated by the function of ppk1 or by other parts of the phosphate pathway under ppk1 regulation.

The phoX deletion results in sensitivity to nutrient but not osmotic and oxidative stress response

It is known that inorganic polyphosphate (poly P) is important to stress responses (nutrient, osmotic, osmotic and oxidative), and specifically the C. jejuni ppk1 and ppk2 mutants are sensitive to nutrient and osmotic stresses [23,40,46]. Since the ΔphoX mutant was defective in poly P accumulation; we investigated whether phoX deletion mutant is sensitive to various stresses. We found that the phoX deletion mutant had a significant defect (P<0.05) in nutrient stress survival after 24 hours of incubation in minimal media without glutamine (Fig 3a). This survival defect was restored in the complemented strain (Fig 3a). Similarly the phoX deletion mutant shows a statistically significant reduction in survival after 48 hours. The survival defect was even greater (approximately 3 logs), compared to wild type cells, at 60 hours (Fig 3a).

Inorganic phosphate is required for deadenylation of glutamine synthetase, which is required for the synthesis of glutamine [49]. Since the phoX mutant is defective in Pi generation, we hypothesized that supplementation of glutamine might correct the nutrient downshift defect. The survival of the ΔphoX mutant was similar to WT in minimal media containing 2 mM glutamine.

Table 1. Antibiotic susceptibility of the ΔphoX mutant.

| Antibiotic     | ΔphoX MIC ± SE (fold change) | WT | ΔphoX |
|---------------|-----------------------------|----|-------|
| Zithromycin   | 0.05±0.02 (1.5)             | 0.03 | 0.03±0.02 |
| Ciprofloxacin | 0.19±0.08 (3.0)             | 0.06 | 0.06 |
| Erythromycin  | 0.25±0.14 (1.4)             | 0.19±0.09 | 0.12±0.09 |
| Gentamicin    | 1.50±0.58 (1.5)             | 1.0 | 1.0 |
| Tetracycline  | 0.31±0.19 (5.2)             | 0.06 | 0.06±0.04 |
| Florfenicol   | 1.50±0.58 (3.0)             | 0.50 | 1     |
| Nalidixic Acid| 16.0±9.2 (4.0)              | 4   | 4     |
| Telithromycin | 1.0±0.58 (2.0)              | 0.50 | 0.50 |
| Clindamycin   | 0.38±0.19 (2.0)             | 0.19±0.09 | 0.19 |
| Polymixin B   | 6.3 (0.5)                   | 12.5 | 12.5 |
| Cholic Acid   | 10,000 (2.0)                | 5,000 | 7,500±3.5 |
| Taurocholic Acid | 100 (1.2)              | 83.3±28.9 | 100 |
| Deoxycholic Acid | 25,000 (1.0)          | 25,000 | 25,000 |
| Fowlidicin-1  | 16 (1.0)                    | 16   | 16    |

* fold change is the quotient of ΔphoX/wild type resistance for a given antibiotic.

*All calculations are the average of three independent repetitions of the assay. Where standard error measurements are absent, measurement for all tests was same and standard error is zero. MIC; Minimal Inhibitory Concentration; SE, Standard Error.

doi:10.1371/journal.pone.0026336.t001
even after 60 hours of incubation (Fig. 3b). Further, addition of 1 mM Pi to the media also corrected the survival defect of the ΔphoX mutant even the absence of glutamine (Fig. 3c). Since Pi is required for deamidation of glutamine synthetase, which is required for synthesis of glutamine, the nutrient survival defect is likely due to insufficient Pi in the ΔphoX mutant.

The ΔphoX mutant had an osmotic stress and oxidative stress response similar to the wild-type strain both in the presence or absence of Pi (Fig. S3a–b). However, the qRT-PCR results indicated a downregulation of katA suggesting that mechanisms other than katA may play a role in oxidative stress response of C. jejuni [30]. Though the ΔphoX mutant showed consistently increased resistance to osmotic stress, it was not significant.

The phoX deletion does not affect motility but has enhanced biofilm

Poly Pi/inorganic phosphate stores have been linked to changes in both motility and biofilm [31,40]. Additionally, regulation of Pi has been implicated in the ability of V. cholera to transit between marine and gastrointestinal lifestyles [30]. Therefore, we investigated whether phoX deletion would induce changes in motility and biofilm formation. The phoX deletion mutant did not show any motility defect on a semisolid agar. The motility of the phoX mutant was comparable to the wild-type strain (Fig. 4a). Further, addition of Pi, though increased the motility, there was no significant difference between the WT and phoX mutant. Similarly there was no significant difference in the motility of the Δppk1 mutant compared to WT (Fig. 4a). On the other hand, the phoX deletion resulted in significantly enhanced biofilm formation (P<0.01) compared to the wild-type C. jejuni and the complementation with the wild type copy of the phoX restored biofilm to wild type levels (Fig. 4b–c). This result is surprising, since phosphate starvation in V. cholera and Pseudomonas is thought to reduce biofilm formation [30,51].

Since phoX is closely associated with Pi levels, it is suggested that phosphate levels are an environmental indicator for biofilm regulation [30]. Therefore, we further tested whether biofilm formation was affected by Pi alone. We found that the ΔphoX mutant’s increase of biofilm can be rescued with the addition of 1 mM Pi (P<0.01), strongly suggesting that the increase in biofilm is affected by the decreased availability of phosphate in the phoX deletion mutant (Fig. 4d). However, the wild-type strain did not show any changes in biofilm formation after the addition of 1 mM Pi (Fig. 4d), nor did the addition of 1 mM Pi affected the biofilm formation in the complemented strain (Fig. 4d). Consistent with our earlier finding, the Δppk1 mutant also has an enhanced biofilm phenotype; however, addition of 1 mM Pi further increased the biofilm formation (P<0.05) in the Δppk1 mutant (Fig. 4c–d) [40].

The phoX deletion mutant has increased resistance to antimicrobials

Both the Δppk1 and ΔtatC deletion mutants have been shown to have increased sensitivity to antibiotics [16,40]. Since the phoX deletion mutant has a defect in poly P accumulation, we further investigated whether this would also result in susceptibility to antimicrobials. In contrast to Δppk1 and ΔtatC deletion mutants, we found that the ΔphoX mutant has increased resistance to some common antimicrobials, including tetracycline, nalidixic acid and, to a lesser degree, to ciprofloxacin (Table 1). The phoX deletion mutant had 3, 4, 3-fold greater resistance to tetracycline, nalidixic acid, florfenicol and ciprofloxacin than wild-type (Table 1). The complementation with the wild type copy of the phoX restored susceptibility to wild-type levels (Table 1). Since biofilms have been implicated as a contributor to increased resistance to antibiotics, it is possible that the increased biofilm activity in the ΔphoX mutant is contributing to this phenotype. In previous studies, tetracycline, and nalidixic acid have been known to kill biofilm-forming cells less efficiently than non-biofilm forming cells [52]. This may suggest that the increased biofilm phenotype seen in ΔphoX may be a mechanism of innate increased antibiotic resistance.

### Table 2. qRT-PCR analysis of change in expression of selected genes in wild type, ΔtatC, and ΔphoX strains.

| Gene/ORF | Fold change in ΔphoX compared to WT | Fold change in ΔtatC compared to WT |
|----------|------------------------------------|------------------------------------|
| ΔahpC (CJJ_0298) | No change | −9.3 |
| ΔahpC (CJJ_0356) | No change | −16.2 |
| CJJ_0379 | No change | −18.9 |
| ppk1 | No change | −4.4 |
| CJJ_1374 | −10.4 | −15.7 |
| csrA | 2.6 | No change |
| katA | −8.5 | −19.2 |
| proP | 4.3 | −27.5 |
| cmcC | −5.8 | No change |
| pstC | No change | Not tested |
| pstD | −2.4 | Not tested |
| ppk1 | −2.1 | −7.1 |
| spoT | −2.9 | 5.5 |
| sodA | No change | −29.3 |

*The difference in gene expression was determined by the threshold cycle(CT) method, and the assay was repeated three times with two replicates each time for each sample. Data represent the mean relative fold change in expression.

WT, Wild Type; ORF, Open Reading Frame.

doi:10.1371/journal.pone.0026336.t002
The phoX deletion results in transcriptional changes in key genes involved in phosphate uptake and stress responses

We used qRT-PCR to investigate how changes in alkaline phosphatase and inorganic phosphate starvation, caused by the phoX deletion, affected the transcription of genes that are commonly associated with environmental stress response or are believed to be part of the phosphate regulon. We found that ppk1 was down regulated 2.1-fold in the ΔphoX mutant compared with a 7-fold down regulation of ppk1 in the ΔtatC mutant (Table 2). However, no changes in the transcription of ppk2 were observed in the phoX deletion mutant although the tatC mutant showed 4-fold down-regulation (Table 2).

ProP has been known to confer osmotic protection [53]. The expression of proP was increased (4.3-fold) in the phoX deletion mutant, compared to a >25-fold down regulation in the tatC deletion mutant (Table 2). However, the ΔphoX mutant has a slightly increased, though not significant, resistance to osmotic stress (Fig. S3b-c) [16]. While phoX deletion increases proP expression, there may be other TAT-substrates which have additive and pleiotropic effects that result in the overall down-regulation of proP in the tatC deletion mutant. Also, we found that CJJ_1374, a VacJ homolog, known to be upregulated during oxidative stress, was down-regulated in both ΔphoX and ΔtatC mutants (Table 2) [43]. While CJJ_1374 was 1.5-fold more down-regulated in the tatC mutant than the ΔphoX mutant, the tatC mutant has an increased sensitivity to oxidative stress and the ΔphoX mutant does not (Table 2) [16]. Similarly, we saw an 8.5-fold down regulation of katA in the ΔphoX mutant, compared to a 19.2-fold down-regulation in the tatC mutant. Additionally, cmcC was down-regulated (5.8-fold) in the ΔphoX mutant. However, the phoX mutant was resistant to certain antimicrobials. It is possible that the antimicrobial resistance is mediated by changes other than CmeABC efflux pump, such as increased biofilm formation (Fig. 4; [54]).

We measured whether pstS and pstC, which are concurrently regulated by the PhosR/S regulon, would be transcriptionally affected by phoX deletion. We found a 2.4-fold down regulation of pstS and no change in the pstC gene (Table 2). This result was

![Graph](image)

**Figure 5. Invasion and intracellular survival of the ΔphoX mutant in INT407 cells.** The ΔphoX mutant displays decrease in invasion (a) and intracellular survival (b) in INT407 human intestinal epithelial cells. Similarly, the ΔtatC mutant is also significantly deficient in invasion as well as intracellular survival in the INT 407 cells. The data represents the average of 3 experiments with 2 replicates in each experiment. Detection limit is represented by a dotted line. * P≤0.05.

doi:10.1371/journal.pone.0026336.g005
unexpected; we predicted that *pstS* would be upregulated during the putative phosphate depletion resulting from the deletion of *phoX*. This result may indicate that an indirect effect of *phoX* deletion, such as nutrient stress contributing to *pstS* regulation [55,56]. We also observed a 2.9-fold down-regulation of *spoT*, a primary effecter of ppGpp degradation. In addition, a post-transcriptional global regulator, *csrA* [24] was upregulated by 2.6-fold.

Survival of the Δ*phoX* mutant is slightly diminished in INT407 cells

To investigate if *phoX* is involved in virulence-associated phenotypes, we examined whether the Δ*phoX* mutant could invade and survive within INT407 human intestinal epithelial cells. The *phoX* deletion mutant was defective in invasion compared to wild-type cells by 4.4 fold (Fig. 5a). The Δ*phoX* mutant’s intracellular survival was also similarly reduced by 4.0 fold compared to wild-type cells in INT407 cells while the complementation restored the defect to WT levels (Fig. 5a–b). However, the Δ*tatC* mutant had more than 7000-fold invasion defect in INT407 cells and no intracellular bacteria were recovered 24 hours post infection suggesting a severe intracellular survival defect (Fig. 5a–b). This suggests that *phoX* activity is not essential for successful invasion and intracellular survival; however, it should be noted that invasion and intracellular assays were performed using rich media (MEM supplemented with fetal bovine serum) where Pi was abundant and readily available even to the Δ*phoX* mutant. Since Δ*phoX* mutant, though defective in generation of Pi from complex organophosphate ester, is not defective in uptake of free Pi because it still carries intact Pi uptake *pst* genes [20,24].

The Δ*phoX* mutant shows diminished ability to colonize day-old chicks

To assess the contribution of *phoX* to *C. jejuni* host colonization, we tested the ability of the Δ*phoX* mutant to colonize day-old chicks. We found that there was 0.8 log difference (P<0.05) in the number of wild-type (average CFU 2.34×10^6 g^-1) and *phoX* deletion mutant (average CFU 3.17×10^5 g^-1) bacteria recovered from the ceca 7 days after chicks were orally infected with 10^3 CFU of each strain (Fig. 6). Similarly, there was approximately 1 log difference (P<0.05) in the number of C. *jejuni* wild-type (average CFU 3.41×10^8 g^-1) and *phoX* deletion mutant (average CFU 5.29×10^7 g^-1) recovered from chick ceca when the inoculation dose was 10^5 CFU/chick (Fig. 6). Chicks inoculated with the *phoX* deletion mutant also had fewer bacteria in the feces compared to wild type. On an average there were approximately 1.1 and 1.2 log fewer bacteria in feces (P<0.05) in chicks inoculated with 10^3 and 10^5 CFU of the mutant (average CFU 1.38×10^5; 3.3×10^4 g^-1), respectively, compared to chicks inoculated with similar amounts of the wild-type strain (average CFU 2.50×10^6; 4.71×10^5 g^-1) (Fig. 6). These results suggest that there is a small but consistent defect in colonization caused by *phoX* deletion. It is possible that the effects of *phoX* deletion are ameliorated by using inorganic phosphate produced by alkaline phosphatases native to the chicks’ gastrointestinal tract [52].

Discussion

In this study we explored how one of the substrates of the TAT system, alkaline phosphatase, contributes to *C. jejuni* pathophysiology. Also we have expanded a model for poly P mediated responses in *C. jejuni* (Fig. 7) [39,40,46]. From our results, we can conclude the following about the phosphate metabolism of *C. jejuni*;

alkaline phosphatase (*phoX*) is necessary for extracellular Pi acquisition and the effects of low inorganic phosphate are a significant part of stress metabolism through poly P and biofilm formation. In particular, *phoX* is important for survival in stationary phase in nutritionally limited conditions where it affects poly P metabolism. Further, without the ability to synthesize inorganic phosphate from extracellular sources via alkaline phosphatase, poly P accumulation is reduced. This suggests that extracellular sources of Pi are necessary for efficient poly P synthesis; *C. jejuni* is partially reliant on the environment to accumulate this important stress response and metabolic mediator. Since little is known about poly P regulation, understanding the origin of the inorganic phosphate that make up the poly P macromolecule is novel.

Both the deletion of *tatC* and *ppk1* have been shown to have pleiotropic effects on bacteria including *C. jejuni* [16,39,40,57].
Deletion of \( \text{tatC} \) can result in diverse phenotypes which can be a direct effect due to defect in translocation of necessary proteins or can be an indirect effect for which the mechanisms are not yet understood. Similarly, \( \text{PPK1} \) has been linked with regulation of several genes in \( \text{P. aeruginosa}, \text{P. gingivalis} \) and \( \text{E. coli} \) \cite{25,58}.

Specifically, microarray analysis of \( \text{ppk1} \) mutants in \( \text{P. aeruginosa} \) and \( \text{E. coli} \) revealed up-regulation of over 250 genes and down-regulation of more than 450 genes. Thus, poly P and its associated enzymes may be identified as "global regulators". Our finding shows that how the deletion of a single, specific variable (\( \text{phoX} \)) contributes to \( \text{C. jejuni} \) resilience to environmental stresses and provides an insight into understanding the complex mechanisms behind poly P and TAT system mediated stress responses.

The \( \Delta \text{phoX} \) mutant has a 3 log nutrient stress defect that is rescued by the addition of 2 mM glutamine (Fig. 3), suggesting that the \( \Delta \text{phoX} \) nutrient stress phenotype is caused by a defect in glutamine metabolism. Interestingly the nutrient downshift defect was also rescued by the addition of 1 mM Pi even in the absence of glutamine. Since it is known that inorganic phosphate is crucial for the deadenylation of glutamine synthetase \cite{49}, it is likely that the nutrient stress defect is associated with defective inorganic phosphate in the \( \Delta \text{phoX} \) mutant. Further, it is reasonable to assume that defective translocation of alkaline phosphatase may be responsible for the nutrient stress phenotype seen in the \( \Delta \text{tatC} \) mutant since both \( \Delta \text{phoX} \) and \( \Delta \text{tatC} \) mutants appear to have similar nutrient downshift defects (Fig. 3) \cite{16}.

The \( \Delta \text{phoX} \) has an increased biofilm phenotype that can be rescued with the addition of 1 mM Pi (Fig. 4d). This suggests that reduced Pi concentration, and consequently reduced poly P, may be a factor that allows \( \text{C. jejuni} \) to sense an environment hostile to growth and initialize defensive measures. This could promote \( \text{C. jejuni} \) survival on surfaces—which are typically low phosphate environments—and improve the pathogen’s ability to endure until it can reach a suitable host. The biofilm results may explain the \( \text{phoX} \) deletion mutant’s decrease in sensitivity to tetracycline (Table 1), a drug that is known to have reduced penetration of biofilms \cite{52}.

The rescue of the biofilm phenotype by addition of 1 mM Pi (Fig. 4d) suggests that the biofilm phenotype is not a direct result of \( \text{phoX} \) deletion, but may be result of cellular response to the Pi depleted conditions that the \( \text{phoX} \) deletion mutant created. In contrast, in the \( \Delta \text{ppk1} \) mutant biofilm formation was further increased with the addition of 1 mM Pi. Since, the \( \text{ppk1} \) mutant is defective in the synthesis of poly P, unlike the \( \text{phoX} \) mutant, the different outcome in the \( \text{ppk1} \) mutant in response to Pi is likely due to pleiotropic effects that the \( \text{ppk1} \) deletion has on the \( \text{C. jejuni} \) \cite{39,40}. This response suggests that although low levels of inorganic phosphate may be sufficient to increase biofilm, poly P

![Figure 7. Hypothetical model illustrating interaction of different poly P-associated proteins in \( \text{C. jejuni} \). Low phosphate conditions activate PhoS phosphorylation of PhoR. Alkaline phosphatase (PhoX) is transcriptionally regulated by two component system PhosRS and is translocated by the TAT system to the periplasm where it is active. PhoX hydrolyzes organophosphate esters to release inorganic phosphate (Pi), which is a preferred source of phosphate for bacteria. Pi uptake into the cytoplasm is mediated by phosphate uptake proteins (PstSCAB) which are also directly regulated by PhoR. Pi is used directly by glutamine synthetase for glutamine metabolism. Also Pi appears to influence biofilm formation. ATP generated from Pi will be utilized for poly P synthesis by PPK1. PPK2 utilizes poly P to generate GTP, while PPX hydrolyzes poly P back to Pi. SpoT synthesizes pppGpp using GTP and ATP. pppGpp will be converted to ppGpp, a molecule that mediates stringent response in \( \text{C. jejuni} \) and other bacteria. SpoT also mediates hydrolysis of ppGpp to GDP and inorganic phosphate. PPK1 may interact with SpoT to regulate its functions. Dashed lines and arrows indicate possible direct or indirect interaction of proteins. Question mark indicates potential pathways predicted based on the experimental evidence from other bacteria.](https://doi.org/10.1371/journal.pone.0026336.g007)
and other factors modify this response. Since the Δppk1 mutant showed increased biofilm in the presence of Pi, we hypothesize that there is a phosphate-related biofilm response parallel to the poly P mediated response and perhaps tied to other environmental stressors.

Our qRT-PCR data shows that a slight down-regulation of ppk1 transcription in the ΔphoX mutant in the stationary phase. This result agrees with the poly P accumulation defect that we observed. Additionally, we saw that the ΔstatC mutant had a greater down-regulation of ppk1 than ΔphoX, this may suggest that an additional TAT substrate contributes to poly P synthesis. Although we found that phoX deletion results in transcriptional down-regulation of oxidative stress genes (datA, CjL_0379, vacJ homologue CjL_1374, abpC) these transcriptional changes did not result in reduced oxidative stress survival phenotype (Fig. S3). These genes are universally more down-regulated in the ΔstatC mutant where the mutant has a significantly increased sensitivity to oxidative stress [16]. However, when grown in minimal media, phoX deletion in general caused fewer transcriptional changes (data not shown); there was a 2.4-fold upregulation of ppk1 and 3.5-fold decrease in ppk2 transcription compared to wild-type. This may suggest that ppk1 and ppk2 regulation is pleiotropically affected by nutritional stressors.

We also conclude that phoX is unlikely to be more than a peripheral mechanism for the survival defect in INT407 cultured cells as well as in vivo chicken colonization. This agrees with previous research on phoB deletion mutant in chickens; the host’s gastrointestinal surfaces have their own alkaline phosphatase enzymes, allowing the ΔphoX mutant to compensate for its alkaline phosphatase deficiency by uptake of host derived free phosphate [59,60]. Less inorganic phosphate availability and other factors such as differences in temperature and oxygen levels in feces compared to ceca could explain an increased survival defect of ΔphoX mutant in feces compared to ceca.

In C. jejuni we found that deletion of phoX resulted in increased biofilm formation. However, in V. cholera, low intracellular phosphate results in decreased biofilm formation that is regulated by PhoB [59]. This is surprising because biofilm inhibition in V. cholera is thought to be part of a cellular transition to an aquatic lifestyle. Similar to V. cholera, C. jejuni contamination of water is a known source of infection [5,59]. C. jejuni’s increase of biofilm in response to deletion of phoX and presumably diminished intracellular phosphate is more similar to plant and soil dwelling pathogens such as A. tumefaciens [61]; this may suggest that the phosphate stressor response that C. jejuni has evolved is more tuned towards survival on an exposed surface rather than a marine environment [62]. Although V. cholera and Pseudomonas have evolved similar phoX genes, there may be crucial phenotype differences that are likely unique to C. jejuni and have an impact on its cell physiology and survival [51,59]. For instance, phosphate starvation in Campylobacter increases biofilm rather than decreasing it, and phosphate starvation did not increase oxidative stress sensitivity, or cause a change in motility [20,63].

In summary, these findings reinforce an important, central theme to alkaline phosphatase and phosphate utilization in C. jejuni. It is nearly universal that the ability to use inorganic phosphate from extracellular environments is critical for bacterial physiology. The molecular response to this information however is highly dependent on the bacterial species and its environment. Therefore we hope that our study will advance understanding of the phosphate utilization in Campylobacter a little further and perhaps suggest additional possible combinations of cellular responses to those modeling other phoX containing bacteria.

Supporting Information

Figure S1 C. jejuni alkaline phosphatase activity in different culture media. Background phosphatase activity in the ΔphoX mutant grown in different Campylobacter culture media. The background alkaline phosphatase activity in the ΔphoX mutant was least in minimal essential medium. Additional washing with MOPS buffer reduced variation and improved alkaline phosphatase activity in the wild type. Each data point is the mean ± standard deviation of 3 experiments. * P≤0.05. (TIF)

Figure S2 Poly P Accumulation in ΔphoX mutant grown in rich media. The ΔphoX mutant is defective in poly P accumulation. Using glassmilk, Poly P was extracted from stationary phase wild type, ΔphoX, ΔstatC and Δppk1 strains grown in MH media. The amount of poly P in the cell was determined by toluidine blue O method. Each data point is the mean ± standard deviation of 3 independent experiments. * P≤0.05. (TIF)

Figure S3 Oxidative and osmotic stress tolerance of the ΔphoX mutant. (a) The ΔphoX mutant has a similar zone of inhibition to the wild type strain when exposed to 20 mM paraquat or 0.3% H2O2 for 24 hours under microaerobic conditions. Addition of 1 mM Pi did not affect the oxidative stress response. Values indicate average zone of inhibition diameter ± standard deviation from three replicate experiments. Only representative images are show. (b-c) wild type, ΔphoX, and ΔphoX strains were grown to mid-log phase, oxidative stress tolerance was determined either on solid media (MH agar) containing 0.17 M NaCl (b) or in liquid media (MH broth) containing 0.25 M NaCl (c). These experiments were performed three times. (TIF)

Table S1 Bacterial strains and plasmids used in this study. (DOC)

Table S2 Oligonucleotide primers used in this study. (DOCX)

Acknowledgments

We thank Dr. Juliette Hanson for assistance with chicken colonization studies. We thank Dr. Jun Lin for providing chicken antimicrobial peptide, fowliecidin-1.

Author Contributions

Conceived and designed the experiments: GR MD DG. Performed the experiments: DG MD ZL. Analyzed the data: DG MD GR ZL. Wrote the paper: MD GR.

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