Gene Expression Profiling of Transcription Factors of *Helicobacter pylori* under Different Environmental Conditions

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*Helicobacter pylori* is a Gram-negative bacterium that colonizes the human gastric mucosa and causes peptic ulcers and gastric carcinoma. *H. pylori* strain 26695 has a small genome (1.67 Mb), which codes for few known transcriptional regulators that control bacterial metabolism and virulence. We analyzed by qRT-PCR the expression of 16 transcriptional regulators in *H. pylori* 26695, including the three sigma factors under different environmental conditions. When bacteria were exposed to acidic pH, urea, nickel, or iron, the sigma factors were differentially expressed with a particularly strong induction of *fliA*. The regulatory genes *hrcA*, *hup*, and *crdR* were highly induced in the presence of urea, nickel, and iron. In terms of biofilm formation, *fliA*, *flgR*, *hp1021*, *fur*, *nikR*, and *crdR* were induced in sessile bacteria. Transcriptional expression levels of *rpoD*, *flgR*, *hspR*, *hp1043*, and *cheY* were increased in contact with AGS epithelial cells. Kanamycin, chloramphenicol, and tetracycline increased or decreased expression of regulatory genes, showing that these antibiotics affect the transcription of *H. pylori*. Our data indicate that environmental cues which may be present in the human stomach modulate *H. pylori* transcription.

**Keywords:** *H. pylori*, transcription factors, environmental cues

### INTRODUCTION

*Helicobacter pylori* is a Gram-negative bacterium, a member of the Epsilon proteobacteria that colonizes the human gastric mucosa and is responsible for causing peptic ulcers and gastric carcinoma (Marshall and Warren, 1984; Parsonnet et al., 1991; Uemura et al., 2001). *H. pylori* survives in the hostile environment found in the stomach, which is partially attributed to the expression of virulence factors, such as secretion systems, cytotoxins, flagella, and adhesins. Unlike other Gram-negative bacteria such as *Escherichia coli* or *Salmonella enterica*, the *H. pylori* genome encodes only few known transcriptional regulators, which control expression of genes involved in bacterial metabolism and pathogenicity. This limited repertoire is likely due to its life style highly adapted to one particular niche, the human gastric mucosa. *H. pylori* strain 26695 has a small
The genome of 1.67 Mb (Tomb et al., 1997), and possesses three genes that code for sigma factors: rpoD (σ\(^{54}\)), rpoN (σ\(^{54}\)), and fliA (σ\(^{28}\)). σ\(^{54}\) is a homolog of Gram-negative vegetative sigma factors responsible for the transcription of housekeeping genes (Tomb et al., 1997; Beier et al., 1998), whereas σ\(^{54}\) and σ\(^{28}\) are two alternative sigma factors dedicated mostly to control expression of flagella components (Fujinaga et al., 2001; Josenhans et al., 2002; Niehus et al., 2004). The response regulator FlgR is also involved in regulation of flagella synthesis (Spohn and Scarlato, 1999b), whereas bacterial chemotaxis is controlled by the CheY protein (Foynes et al., 2000; Terry et al., 2005). Master regulators of response to metals such as Fur, NikR, and CrdR activate (Foynes et al., 2000; Terry et al., 2005). Little is known about the effects of environmental cues on the expression of H. pylori regulatory genes, including some poorly investigated transcriptional regulators.

In this work, we determined the expression profile of the transcriptional repertoire of H. pylori strain 26695 under several environmental conditions relevant for adaptation to its particular ecological niche of the human stomach, such as acidic pH, urea, nickel, and iron. In addition, we analyzed the expression of regulatory genes in biofilm formation and in the presence of AGS gastric epithelial cells. Finally, we studied the effect of the antibiotics kanamycin, chloramphenicol, and tetracycline on the transcription of regulatory genes. Our study describes the transcriptional expression of H. pylori regulatory genes in response to different environmental conditions.

### MATERIALS AND METHODS

**In silico Identification of H. pylori Transcription Factors**

Selection of H. pylori transcription factors was performed as previously reported in the literature (see Table 1) (Danielli...
et al., 2010)]. Sequence data and loci annotations from 260 *H. pylori* genomes were retrieved from the NCBI database by a series of custom Perl scripts. In addition, the genomes of 57 *Helicobacter* non-*pylori* strains were included in the comparative analysis (Supplementary Table S1). Each putative transcriptional factor was queried using PSI-BLAST (Altschul et al., 1997) under the following parameters: matrix = BLOSUM62, word size = 3, PSI-BLAST threshold = 0.005, expect threshold = 10, and without filtering low complexity regions. Hits were carefully examined and selected according to their functional annotation.

### Bacterial Strains and Culture Conditions

*H. pylori* 26695 was grown for 3 days on blood agar plates containing 10% defibrinated sheep blood, at 37°C under microaerophilic conditions. A bacterial suspension was prepared in *Brucella* broth (BB), and adjusted to an optical density of 0.1 at 600 nm (2 × 10⁶ CFU/ml). *H. pylori* was then grown at 37°C for 24 h (logarithmic growth phase) or 48 h (stationary growth phase) in BB supplemented with 10% decomplemented fetal bovine serum (BB + FBS) under the following conditions: adjusted to pH 5.5, or containing either urea [5 mM CO(NH₂)₂], nickel [250 mM NiCl₂], or iron [150 mM (NH₄)₂Fe(SO₄)₂·6H₂O] as previously described (Contreras et al., 2003; Wen et al., 2003; Vannini et al., 2014; Cardenas-Mondragon et al., 2016). Fold-changes in transcription were determined by calculating the relative expression of transcription regulator genes under different environmental conditions as compared to expression in BB + FBS. Experiments were performed in triplicate on three different days and the results shown are the mean of the data produced.

### RNA Isolation and Quantitative RT-PCR

Total RNA was extracted from bacteria grown under different culture conditions using the hot phenol method (Jahn et al., 2008) with some modifications. Briefly, after the lysate was obtained, an equal volume of phenol-saturated water was added, mixed and incubated at 65°C for 5 min. The samples were chilled on ice and centrifuged at 19,000 g for 10 min at 4°C. The aqueous layer was transferred to an 1.5 ml Eppendorf tube, RNA was precipitated with cold ethanol and incubated at −20°C overnight. The RNA was pelleted by centrifugation at 19,000 g for 5 min. The samples were air dried for 15 min in the Centrifugal Vacuum Concentrator 5301 (Eppendorf). The pellets were resuspended in 100 μL of DEPC-treated water. Purification of RNA was performed using the TURBO DNA-free kit (Ambion, Inc.). Quality of RNA was assessed using a NanoDrop (ND-1000; Thermo Scientific) and a bleach 2% agarose gel as previously described (Aranda et al., 2012). qRT-PCR was performed as previously reported (Ares et al., 2016). Specific primers were designed with the Primer3Plus software and are listed in Table 2. The absence of contaminating DNA was controlled by lack of amplification products after 35 qPCR cycles using RNA as template. Control reactions with no template (water) and with no reverse transcriptase were run in all experiments. 16S rRNA (HPrrnA16S) was used as a reference gene for normalization and the relative gene expression was calculated using the 2^−ΔΔCt method (Livak and Schmittgen, 2001). Expression of 16S rRNA remained unaffected in all conditions tested (Supplementary Figure S1).

### Biofilm Formation

*H. pylori* was grown on blood agar medium supplemented with 10% defibrinated sheep blood at 37°C under microaerophilic conditions. Biofilm formation on abiotic surface (polystyrene) was analyzed using 6-well polystyrene plates, inoculated with 3 ml of a bacterial suspension (in BB + FBS, at a final concentration of OD₆₀₀nm = 0.1) in each well. The plates

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1 ftp://ftp.ncbi.nlm.nih.gov/genomes/genbank/bacteria/

2 http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi/

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### Table 2: Primers for qPCR used in this study.

| Primers | Sequence | Target gene |
|---------|----------|-------------|
| rpoD-F  | TAT GGC TCA AGT GCC AGA AG | rpoD |
| rpoD-R  | TGT TGG GGG CTA GAT CAA AG | |
| rpoN-F  | CAG CGG GTT GAA TAA TGA GG | rpoN |
| rpoN-R  | ACG CTT CGG CAC TTT TTC | |
| filA-F  | TCG TCT AAA AGA GCG CTT GC | filA |
| filA-R  | CTT CGC ATA CCC CCA AAA AG | |
| hrcA-F  | TTT CTT CGC CAC TGG ATT AC | hrcA |
| hrcA-R  | GAA AGA AGC AGC TAT GGA GC | |
| ansR-F  | GAG CGA GTT TCT GCT CCA AC | ansR |
| ansR-R  | GCC CGT CTA AAT TAG GCA AAG | |
| hp0222-F | CTA GQA CGC AAA CCA AAA GC | hp0222 |
| hp0222-R | CCC ACG CCT TCT TCT TCT | |
| hp0564-F | GTC GCT GTA GAT GAG CTG AAA C | hp0564 |
| hp0564-R | GCC GGT TQA CAA AAG ATT TG | |
| flgR-F | CAG GCC TTA AAA GTC GCA AG | flgR |
| flgR-R | CTC TAT AAA AGG GTG CTG TG | |
| hup-F | GTG QAG TTG ATC GGT TTT G | hup |
| hup-R | TTA GGC ACC GTG TTT TCT | |
| hp1021-F | GGT GCG CAA GAT CCT CTA GA | hp1021 |
| hp1021-R | GCG GGT TQA GTA TTA TCT TG | |
| hspF-F | CGG QGC TGG ATT ATT TCT TG | hspF |
| hspF-R | TGT TTG TCG AGA GCG TCG T | |
| fur-F | GAA GAA GTG GTG AGC GTT TTG | fur |
| fur-R | CTT TCT GTG QGA TAG AGC | |
| hsrA-F | GGA AGA AGT CCA TGC GTG TG | hsrA |
| hsrA-R | CAA AGC AGC CTC AAT CTT TG | |
| cheY-F | TGG AAG CTT GTG AGA AAC TG | cheY |
| cheY-R | CAG AGC GCA CCT TTT TAA CG | |
| nikR-F | CAT CGC CTT GTG TCT TG | nikR |
| nikR-R | CAT GTC GCG CAC TAA TCT TG | |
| crdR-F | CTT AGG CTT GCC TAA AAT GC | crdR |
| crdR-R | CAA AGC CCC CAA AAA CAC | |
were incubated during 3 days at 37°C under microaerophilic conditions as previously reported (Cardenas-Mondragon et al., 2016). Supernatant (planktonic) and adhered (sessile) bacteria were recovered for RNA extraction. Fold-change in gene transcription was determined by calculating the relative expression of transcription regulator genes within biofilms (sessile bacteria) as compared to planktonic bacteria. Quantifications were performed in triplicate on three different days and the results shown are the mean of the three experiments.

Infection of AGS Cells
AGS gastric epithelial cells were grown to about 75% confluence in RPMI-1640 medium containing 10% FBS, and washed thrice with PBS before adding fresh RPMI media with 10% FBS. *H. pylori* 26695 was grown in BB + FBS for 24 h, suspended in RPMI, and added to the AGS cell culture at a multiplicity of infection (MOI) of 100 (bacteria/cell). Infected cells were incubated at 37°C under microaerophilic conditions for 0 or 6 h, and bacteria were recovered. At the end of the incubation period, the *H. pylori*-infected AGS cells were washed thrice with PBS and lysed with 0.1% Triton X-100 for 10 min. Large debris and nuclei were removed by centrifugation for 5 min at 200 × g and adhered bacteria were pelleted at 20,000 g for 10 min. RNA was extracted from adhered bacteria to determine gene expression. Fold-change in gene transcription was determined by calculating the relative expression of the transcription factors genes with respect to bacteria at time 0 of infection. Fold-change in gene transcription of *H. pylori* grown in RPMI-1640 + FBS (for 0 or 6 h) was calculated as control of expression. Assays were performed in triplicate on three different days and the results shown are the mean of the three experiments.

Transcription in the Presence of Antibiotics
*H. pylori* was grown in BB + FBS at 37°C for 48 h (stationary phase), with gentle shaking under microaerophilic conditions. The antibiotics kanamycin (Km, 50 µg/mL), chloramphenicol (Cm, 30 µg/mL) or tetracycline (Tc, 10 µg/mL) were added and the cultures were incubated for 1 h as previously described (Christensen-Dalsgaard et al., 2010; Cardenas-Mondragon et al., 2016). Antibiotics were used at the minimal inhibitory concentrations that have been reported for *E. coli* and *S. enterica* (Christensen-Dalsgaard et al., 2010; Maisonneuve et al., 2011; Silva-Herzog et al., 2015; Li et al., 2016). Fold-change in gene transcription was determined by calculating the relative expression of the transcription regulators genes in the presence of each antibiotic as compared to bacteria growing without antibiotics for 1 h. Experiments were performed in triplicate on three different days and the results shown are the mean of the three experiments.

Heatmap Construction
To show the fold-changes in gene expression, we selected the “heatmap.2” function of the R software, using the “gplots” package. The rows (culturing conditions) were hierarchically clustered (“hclust” function, “ward.D” method) according to the absolute fold-changes in gene expression.

In order to illustrate the presence/absence of transcription factors in all *Helicobacter* genomes, an amino acid sequences content matrix (“heatmap” function) was built using the R software1 (v3.2.4). 260 *H. pylori* and 57 *H. pylori* genomes were retrieved from the NCBI database4 by a series of custom Perl scripts. These paired loci were hierarchically clustered (“hclust” function, “ward.D” method) according to their loci-content using a sidelong dendrogram.

Statistical Analysis
For statistical differences, one-way ANOVA followed by the Tukey’s comparison test was performed using Prism5.0 (GraphPad Software Inc., San Diego, CA, USA). $p \leq 0.05$ was considered statistically significant.

RESULTS
Environmental Cues that Trigger the Expression of Transcription Factor Genes
*H. pylori* adaptation to the gastric mucosa conditions is mediated by a limited number of regulatory genes. An analysis of the reports of 26695 *H. pylori* strain identified 16 genes that code for transcriptional regulators, including three sigma factors (Table 1). We performed qRT-PCR on RNA extracted from bacteria grown during both exponential (24 h) and stationary phase (48 h) and the expression of all regulatory genes was calculated during both growth phases. Expression of most genes was higher during stationary phase than in exponential phase, except for *rpoD* and *hp0564* (Figure 1A). Therefore, we determined the expression of all transcription regulators during stationary phase in media with acid pH or in the presence of urea, nickel, or iron. None of these environmental variations promoted or inhibited growth of *H. pylori* (Figure 1B). Interestingly, the conditions tested resulted mostly in increased expression of transcription regulators (Figures 1C–F, 4). Regarding sigma factors, *fliA* expression increased with all treatments, with the highest induction levels observed in response to nickel. The same was true for *rpoD* with exception of treatment with iron, which resulted in down regulation of the gene (Figure 1F); whereas *rpoN* expression significantly increased only after exposure to urea or nickel (Figures 1D,E). Concerning the other transcriptional regulators, acidic pH resulted in down regulation of *arsR*, *hp0564*, and *flgR* and a moderate increase of *hup*, *cheY*, or *cdrR*, whereas the remaining genes were unaffected (Figure 1C). Exposure of bacteria to urea and nickel ions resulted in more pronounced transcriptional changes (Figures 1D,E). However, whereas expression of most transcription factors increased considerably, *hp0564* and *fur* showed only subtle

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1http://www.R-project.org/
2ftp://ftp.ncbi.nlm.nih.gov/genomes/genbank/bacteria/
changes in response to urea and nickel. hp0564, fur, and rpoD were the only genes tested to be down regulated in response to iron, whereas expression of the other transcription regulators increased (hrcA, hup, hp1021, hsrA, cheY, nikR, and crdR), or did not change (arsR, hp0222, flgR, and hspR) (Figure 1F).
The Effect of Biofilm Formation and Interaction with Gastric Epithelial Cells on Expression of H. pylori Transcription Factors

During infection, H. pylori closely interacts with the cells of the gastric mucosa, which may result in bacterial biofilm formation in later stages of infection (Carron et al., 2006). To study the effect of bacterial interaction with cells of the gastric mucosa or the growth in biofilms on the expression of transcription regulators, bacteria were grown either stationary on polystyrene surfaces or brought into contact with AGS cells, and their transcription profiles were analyzed. As control for the interaction with AGS cells, H. pylori was grown in RPMI-1640 medium for the same amount of time, which did not result in any changes in gene transcription. Expression of rpoN did not change during growth in biofilm or upon attachment to AGS cells, whereas rpoD expression increased under both conditions (Figure 2A). However, the most striking effect among the three sigma factors was a dramatic increase of fliA expression in response to biofilm formation (Figure 2). Only few of the other regulatory genes remained unaffected by the interaction with abiotic surfaces (hrcA), or epithelial cells (hp1021, fur, nikR, and crdR). All other transcriptional regulators were upregulated upon contact with AGS cells, and to a greater extent during biofilm formation (Figures 2, 4).

Antibiotic Exposure Decreases Expression of Most H. pylori Transcription Regulators

Our group recently reported that antibiotics affect the expression of virulence factors in H. pylori (Cardenas-Mondragon et al., 2016). Whereas the environmental conditions tested here mostly upregulated expression of the transcription regulators analyzed, exposure to different antibiotics resulted predominantly in gene repression (Figures 3, 4). This was likely not due to compromised cell growth, since the antibiotic concentrations used here did not affect the viability of the bacteria (Supplementary Figure S2). Among the three sigma factors, rpoD expression was not affected by exposure to kanamycin or tetracycline and increased in response to chloramphenicol (Figure 3), whereas expression of rpoN and fliA was downregulated or not affected after exposure to all three antibiotics tested (Figure 3). Expression levels of the other transcription regulators were mostly repressed in response to antibiotic treatment, particularly upon exposure to kanamycin or chloramphenicol (Figures 3A, B). Only hrcA and hup mRNA levels were slightly increased in the presence of kanamycin, whereas those of hp1021 were not affected (Figure 3A). While negatively regulating expression of most transcription factors, chloramphenicol led to a mild increase of hup and hp1021 levels, and did not affect expression of fur (Figure 3B). In contrast, tetracycline had stimulating effects on the expression of several transcription factors, including hp0166, hp0222, hp0564, hup, and hp1021 (Figure 3C). Transcription of flgR, nikR, and crdR decreased upon tetracycline treatment, whereas transcription levels of hrcA, hspR, and cheY were not affected (Figure 3C).

Transcriptional Regulator Genes Are Highly Conserved in H. pylori Strains

We performed a Blast search in genomes deposited in GenBank using the amino acid sequences of the 16 transcriptional regulators identified in H. pylori 26695. The transcription factors were highly prevalent and well conserved among different H. pylori isolates (Figure 5). We studied the occurrence of these genes in other Helicobacter species, and found that their presence changed from one species to another. Two species that are phylogenetically closely related to H. pylori, H. acinonychis

http://www.ncbi.nlm.nih.gov/genbank/
Expression of Transcriptional Regulators of *H. pylori*

**FIGURE 3** | Effect of antibiotics on expression of transcription factors. Expression levels of transcription factors after treatment of bacteria with different antibiotics (A) Kanamycin, (B) Chloramphenicol, and (C) Tetracycline for 1 h were determined by qRT-PCR and compared to those in untreated bacteria. Data are expressed as fold-change expression and represent the means and standard deviations of at least three independent experiments. ns, not significant; statistically significant ***p < 0.001, **p < 0.01, *p < 0.05.

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*H. pylori* is a highly specialized bacterium that is exclusively found in the human gastric mucosa. In this work, we describe the expression of *H. pylori* transcriptional regulatory genes under different environmental conditions. Most transcription factors were highly expressed when *H. pylori* reached the early stationary phase. At this growth phase, *H. pylori* is exposed to specific stress signals such as pH changes, starvation, reactive oxygen species that would activate its transcriptional repertoire, suggesting that the stationary phase may mimic the conditions found by the bacteria in the host. Acid pH, the presence of urea, nickel, or iron are environmental cues required for optimal adaptation of *H. pylori* to its natural niche. Whereas all of the above conditions boosted *fliA* expression, *rpoD* showed only a mild increase in transcription after bacterial exposure to acid pH, urea, and nickel, and a decrease in response to iron. The expression of *rpoN* significantly increased upon treatment of bacteria with urea or nickel; RpoN was initially described as regulator of flagellar genes, but recent studies show that it also controls several bacterial regulatory processes involved in energy metabolism, biosynthesis, protein fate, oxidative stress, and virulence (Sun et al., 2013). *fliA* was strongly expressed even under environmental conditions that are not common inducers of flagella synthesis, but are known to affect other bacterial components or pathways. In fact, FliA regulates expression of outer membrane proteins, lipopolysaccharide synthesis, DNA restriction, and CagA (Josenhans et al., 2002; Baidya et al., 2015), a protein involved in virulence and associated with the development of gastric carcinoma (Ohnishi et al., 2008). Our data support the notion that FliA regulates different bacterial functions other than the flagellum.

*H. pylori* is exposed to changes in pH while passing from the stomach lumen through the mucus layer to interact with epithelial cells, and this pH gradient is used by the bacteria for spatial orientation (Schreiber et al., 2004). Accordingly, changes in pH strongly affect expression of transcriptional regulators that control genes involved in colonization and persistence in the human host. Our data show that an acidic pH repressed *arsR*, *hp0564*, and *flgR*, while stimulating *hup*, *cheY*, and *crdR*. The *hup* gene codes for the HU nucleoid protein, which has a regulatory role in the response to acid stress in *H. pylori*. Thus, *hup* mutants are less viable than wild type bacteria at pH 5.5 and during stomach colonization due to down regulation of both urease (*ureA*) and arginine decarboxylase (*speA*) in the absence of HU nucleoid protein (Wang et al., 2012; Almarza et al., 2015). CheY and CrdR response regulators are also crucial for a successful colonization of the animal stomach (Foynes...
et al., 2000; Panthel et al., 2003; McGee et al., 2005; Terry et al., 2005). CheY expression is essential for the chemotactic motility required to reach and colonize the gastric epithelia, and is likely to be triggered in the acidic milieu of the stomach lumen. In contrast, CrdR has not been shown to be involved in the regulation of gene expression in response to acidic pH (Pflock et al., 2007b), although copper can be present in the acidic environment of the human stomach, and regulation of its uptake is important for keeping the balance between supplying copper as respiration co-factor, and avoiding copper-induced toxicity (Haley and Gaddy, 2015). Interestingly, the master regulator of the acid response arsR was repressed in acidic pH, which confirms reports that ArsR may also act as transcriptional auto-repressor under an acidic pH (Dietz et al., 2002).

Urea, nickel, and iron are crucial for H. pylori pathogenesis and they control regulatory networks responding to their presence. We found that expression of most of the transcription regulators tested increased when bacteria were exposed to urea, nickel, or iron. Nickel serves as essential co-factor for the urease enzyme, which enables H. pylori survival at acidic pH (Khan et al., 2009). The up regulation of nikR expression that we observed contrasted with the auto-negative regulation reported for the nikR promoter (Delany et al., 2002; Contreras et al., 2003). The conditions of growth (stationary phase) and nickel concentrations (250 μM) that we tested resulted in increased nikR expression. However, nikR expression showed slight variations in response to low (1 μM) and high (100 μM) concentrations of nickel (Davis et al., 2006), suggesting that nickel may modulate nikR transcription in a concentration-dependent manner. Interestingly, rpoD and fur were down regulated in the presence of iron. While iron-mediated fur repression can be explained by the negative auto-regulation of this transcription factor upon iron-binding (Delany et al., 2002), the decrease in rpoD levels is hard to explain. Whilst the evaluation of each environmental condition provides relevant information about H. pylori physiology, the combination of these stimuli could better mimic the in vivo response of the bacteria in the infection context.

One of the strategies employed by H. pylori to persist and colonize the stomach is biofilm formation. Analysis of bacteria grown in biofilm revealed an interesting expression pattern of the three sigma factors: whereas rpoN was not affected, expression of rpoD and fliA increased during biofilm formation. FliA has been found to control the lpxC gene, which is involved in the early steps of lipid A synthesis in H. pylori (Josenhans et al., 2002). The marked increase of fliA expression that we found in sessile, aggregated bacteria is in agreement with reports about the effect of lipid A architecture on biofilm formation (Gaddy et al., 2015). In addition, with the exception of hrcA, expression of all transcription factor genes studied increased during biofilm formation, and the relative increase of several of them was the highest increase observed across all the conditions tested. This remarkably activated state of the regulatory transcriptome highlights the importance of forming sessile microbial communities in H. pylori ecology.

Similar to the response during biofilm formation, the presence of gastric epithelial cells significantly increased expression levels of several transcription factors, except for fur, nikR, and crdR. Expression of these three master regulators of virulence remained unaffected in our AGS cell model, which correlates with the previously reported lack of activation or repression of these regulators after the interaction with gastric epithelial cells (Kim et al., 2004).

It has been hypothesized that the reduced number of transcriptional regulators in the H. pylori genome has been compensated by gain of functions in the remaining transcription factors, as compared to their functions by homologs found in other species (Ernst et al., 2005a). For instance, the H. pylori Fur protein was not only found to be involved in iron homeostasis, but it also participated in several other additional pathways.
including those of oxidative stress resistance (Harris et al., 2002) and acid regulation (Bijlsma et al., 2002; van Vliet et al., 2004), and has been found essential for colonization of the gastric mucosa (Bury-Mone et al., 2004). Moreover, unlike Fur homologs in other species, H. pylori Fur has been found to mediate gene regulation even in its iron-free (apo) form (Bereswill et al., 2000; Delany et al., 2001; Carpenter et al., 2013). Interestingly, whereas most conditions tested here showed only moderate effects on Fur expression, biofilm formation resulted in a marked up regulation of the gene, suggesting functions beyond regulation of iron metabolism.

The presence of antibiotics can alter the expression of genes related to the bacterial stress and virulence on a transcriptional level. Interestingly, most regulatory genes were repressed in response to antibiotic treatment. rpoN, fliA, fliR, and crdR genes presented a negative regulation profile in the presence of kanamycin, chloramphenicol, and tetracycline. In contrast, expression levels of rpoD and hup were highly stimulated under chloramphenicol or tetracycline treatment. These last antibiotics inhibit bacterial translation, differentially affecting the 50S and 30S ribosomal subunits, respectively. The molecular mechanisms responsible for the regulation in expression of transcriptional regulators in the presence of antibiotics have been poorly studied. About this, 16S rRNA expression was affected in the presence of kanamycin and chloramphenicol, showing that this gene was not completely stable and that antibiotic treatment may have affected the expression of this reference gene under these conditions. For a better analysis in presence of these antibiotics, it is necessary the selection and validation of other reference genes for qRT-PCR normalization as was recently reported (Martins et al., 2017).

During analysis of Helicobacter sequences we found that the transcriptional regulators were highly identical among H. pylori species. Interestingly, H. acynonichis and H. cetorum grouped together with H. pylori, corroborating the close phylogenetic relation between these species. The transcriptional regulators HP0222 and HP0564 appear to be conserved in H. pylori and its closely related species, while they were absent in most of the remaining Helicobacter species. Since H. pylori, H. acynonichis, and H. cetorum are all found within mammalian stomachs, these two regulators may confer an adaptive advantage in this particular ecological niche. In line with these findings, expression levels of both, hp0564 and hp0222 increased in contact with AGS gastric epithelial cells, corroborating a report by Kim et al. (2004) on hp0222. However, we did not observe enhanced hp0222 expression under acidic pH, contrasting with the report by Ang et al. (2001).

Recently, we reported the transcriptional profiling of type II toxin-antitoxin genes under different environmental conditions (Cardenas-Mondragon et al., 2016). The type II antitoxins function as transcriptional repressors of their own expressions (Yamaguchi and Inouye, 2011) and also regulate the expression of other genes related with cellular functions such as biofilm formation, persistence, and the general stress response (Wang and Wood, 2011; Hu et al., 2012). Our findings here expand the transcriptional repertoire of H. pylori to respond to the different stresses found in the stomach niche.
In summary, our data show that the repertoire of transcriptional regulators of *H. pylori* possesses a functional plasticity needed to respond to different environmental cues and to integrate them for the survival and persistence of this bacterium in the stomach niche.

**AUTHOR CONTRIBUTIONS**

Conceived and designed the experiments: MDC. Performed the experiments: MDC, KvB, MA, LP, JM-C, HV-S, and CJ-G. Analyzed the data: MDC, KvB, and MA. Wrote the paper: MDC, KvB, and JT.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2017.00615/full#supplementary-material

**FIGURE S1** | Expression of reference gene (HPrrnA16S) under different environmental conditions. Panels show the expression of reference gene during stationary phase in BB + FBS with changes in pH and concentrations of urea, nickel, and iron (A), or in presence of antibiotics (B) or in contact on abiotic and biotic surfaces (C). (-) Indicates the BB + FBS plain (neutral pH with no addition of components). Quantification of expression is showed as copies of HPrrnA16S/µg RNA.

**FIGURE S2** | Effect of antibiotics on *H. pylori* growth. Determination of colony forming units (CFU) of *H. pylori* 26695 grown during 1 h in presence of antibiotics (Km, Kanamycin; Chloramphenicol, Cm; Tetracycline, Tc).
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