ArsRS-Dependent Regulation of \textit{homB} Contributes to \textit{Helicobacter pylori} Biofilm Formation

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One elusive area in the \textit{Helicobacter pylori} field is an understanding of why some infections result in gastric cancer, yet others persist asymptotically for the life-span of the individual. Even before the genomic era, the high level of intraspecies diversity of \textit{H. pylori} was well recognized and became an intriguing area of investigation with respect to disease progression. Of interest in this regard is the unique repertoire of over 60 outer membrane proteins (OMPs), several of which have been associated with disease outcome. Of these OMPs, the association between HomB and disease outcome varies based on the population being studied. While the molecular roles for some of the disease-associated OMPs have been evaluated, little is known about the role that HomB plays in the \textit{H. pylori} lifecycle. Thus, herein we investigated \textit{homB} expression, regulation, and contribution to biofilm formation. We found that in \textit{H. pylori} strain G27, \textit{homB} was expressed at a relatively low level until stationary phase. Furthermore, \textit{homB} expression was suppressed at low pH in an ArsRS-dependent manner; mutation of \textit{arsRS} resulted in increased \textit{homB} transcript at all tested time-points. ArsRS regulation of \textit{homB} appeared to be direct as purified ArsR was able to specifically bind to the \textit{homB} promoter. This regulation, combined with our previous finding that ArsRS mutations lead to enhanced biofilm formation, led us to test the hypothesis that \textit{homB} contributes to biofilm formation by \textit{H. pylori}. Indeed, subsequent biofilm analysis using a crystal-violet quantification assay and scanning electron microscopy (SEM) revealed that loss of \textit{homB} from hyper-biofilm forming strains resulted in reversion to a biofilm phenotype that mimicked wild-type. Furthermore, expression of \textit{homB} in trans from a promoter that negated ArsRS regulation led to enhanced biofilm formation even in strains in which the chromosomal copy of \textit{homB} had been deleted. Thus, \textit{homB} is necessary for hyper-biofilm formation of ArsRS mutant strains and aberrant regulation of this gene is sufficient to induce a hyper-biofilm phenotype. In summary, these data suggest that the ArsRS-dependent regulation of OMPs such as HomB may be one mechanism by which ArsRS dictates biofilm development in a pH responsive manner.

\textbf{Keywords:} \textit{homB}, ArsRS, \textit{H. pylori}, biofilms, OMPs
INTRODUCTION

*Helicobacter pylori* is an intriguing pathogen. Not only does it colonize the inhospitable gastric niche, but it does so with incredible efficiency. It is estimated that approximately 50% of the world’s population is colonized by *H. pylori* (Goh et al., 2011). In a majority of cases, infection with *H. pylori* remains in a relatively quiescent, asymptomatic state. For those individuals that develop *H. pylori*-associated diseases, symptomatic *H. pylori* infections can present as gastritis, duodenal or gastric ulcers, gastric adenocarcinoma, or mucosa associated lymphoid tissue (MALT) lymphoma (Bauer and Meyer, 2011). Given the broad range of clinical manifestations associated with *H. pylori* infection, one of the most tantalizing avenues of research is identification of the bacterial factors that dictate disease progression. Indeed, the extensive heterogeneity of the *H. pylori* genome makes this pathogen an attractive target in the hunt for factors that can predict disease outcome.

Greater than 20% of *H. pylori*’s gene content varies between strains (Dong et al., 2009). Many of these auxiliary genes encode for proteins of unknown function; however, genes of known function include those encoding for DNA modifying proteins, LPS modifying proteins, outer membrane proteins, and members of the Cag pathogenicity island (PAI) (Dong et al., 2009). In fact, one of the main predictors of severe disease outcome, particularly gastric cancer, is the presence of the cytotoxin, CagA, which is encoded on the PAI. Interestingly, while CagA-positive strains are associated with an increased risk for development of gastric cancer as compared to CagA-negative strains, the risk among CagA-positive strains is further delineated based on conserved polymorphisms in the CagA C-terminal EPIYA motif (Cover, 2016). Thus, as demonstrated by the previous example, *H. pylori*’s panmictic structure is based not only on the presence or absence of genes, but also on conserved polymorphisms within certain genes.

Another class of genes that has been of particular interest in terms of an association with disease outcomes are those that encode for outer membrane proteins (OMPs) (Cover, 2016). Gram-negative OMPs are unique because, unlike many integral membrane proteins that are comprised of alpha helices, OMPs are beta-barrel proteins (Koebnik et al., 2000; Schulz, 2002; Wimley, 2003). OMPs have been shown to function in a variety of roles: molecular transportation, antibiotic resistance, and virulence (Koebnik et al., 2000; Schulz, 2002; Wimley, 2003). For a pathogen that resides within a very specific niche, *H. pylori* encodes an extensive array of OMPs. In the majority of Gram-negative bacteria, two to three percent of the genome is typically dedicated to OMPs (Wimley, 2003). For a bacterium such as *Haemophilus influenza*, which has a genome size similar to *H. pylori*, this equates to approximately 40 proteins (Wimley, 2003). In comparison, up to 60 OMPs are predicted to be encoded by *H. pylori*’s relatively small genome (1.6 Mb); this is equal to approximately 4% of the genome (Alm and Trust, 1999; Alm et al., 2000). This number becomes even more impressive when one considers that *H. pylori* is only predicted to encode 17 regulatory proteins (Scarlatto et al., 2001).

Several *H. pylori* OMPs have been identified as adherence factors (reviewed in Oleastro and Menard, 2013). While adhesins may not directly damage the host, adherence is likely paramount to virulence for a persistent pathogen like *H. pylori*. This is supported by the fact that CagA is delivered to cells in a contact dependent manner. Without an adherent population, CagA would not reach the host epithelium (Jones et al., 2010). Indeed, in some strains the presence of hopQ was found to be required for CagA translocation (Belogolova et al., 2013). Additionally, the intimate attachment of *H. pylori* to the gastric epithelium is well documented (Schreiber et al., 2004) and believed to be crucial for maintenance of long term infection (Blaser and Kirschner, 1999). Finally, given that surface adherence is the initiating step in biofilm formation, OMPs likely contribute to biofilm formation as well (O’Toole et al., 2000). In fact, biofilm formation is thought to contribute to many chronic bacterial infections, and the presence of *H. pylori* biofilms in *vivo* has been observed (Carron et al., 2006; Coticchia et al., 2006; Cellini et al., 2008; Cammarota et al., 2010).

The high level of genetic diversity seen with the OMPs has led several groups to investigate the association between specific OMP(s) and clinical outcomes (reviewed in Oleastro and Menard, 2013). One OMP that shows an association with more severe disease outcomes is HomB. *homB* was identified in a screen for genes associated with formation of duodenal ulcers; the authors compared isolates from a duodenal ulcer patient and a gastritis patient (Oleastro et al., 2006). Further analysis showed that in individuals less than 40 years of age, *homB* was significantly associated with peptic ulcer disease (Oleastro et al., 2006, 2008, 2009b). Following this initial publication, several groups have studied the association between *homB* and disease severity in a variety of populations (Oleastro et al., 2008, 2009b; Jung et al., 2009; Talebi Bezmín Abadi et al., 2011; Kang et al., 2012).

Despite these initial findings, in comparison to other OMPs that are associated with disease outcomes, very little is known about this OMP or the larger Hom family at a molecular level. From the original genome publication, we know that HomB is a member of a small OMP family, the *Helicobacter* outer membrane protein or Hom family. There are four members that make up the hom family: HomA, HomB, HomC, and HomD. Based on the few molecular studies that have been conducted, HomB appears to be involved in adherence (Oleastro et al., 2008; Oleastro and Menard, 2013). Furthermore, serological evidence suggests that HomB is expressed *in vivo* (Oleastro et al., 2008). Additionally, two interesting observations that arose from the epidemiological studies were that (1) *homB* could occupy two different loci, locus A and locus B, and (2) strains can carry a single or double copy of *homB* (Oleastro et al., 2009a, 2010; Kang et al., 2012). To date, neither phase variation nor recombination has been demonstrated for *homB*; however, differences in distribution of *homB* location and copy number have been reported (Oleastro et al., 2009a; Servetas et al., 2018). This is particularly intriguing given that the *homB* genotype varies between Western and East Asian isolates and because the association between *homB* and disease severity is dependent on the population being studied (Oleastro et al., 2008, 2009b; Jung...
et al., 2009; Hussein, 2011; Talebi Bezman Abadi et al., 2011; Kang et al., 2012).

Given the lack of molecular detail concerning HomB, we compared the promoter regions of locus A and locus B, assessed homB expression, and examined the role of HomB in H. pylori biofilm formation. Our data indicate that both homB loci have nearly identical promoter regions, which we reason may maintain the presence of transcriptional binding sequences. Interestingly, subsequent evaluation of homB expression revealed that homB was suppressed in response to low pH and that this regulation occurs via the ArsRS two component system. ArsRS is an acid responsive two component system that is comprised of a sensor kinase, ArsS, and a response regulator, ArsR. ArsS autophosphorylates in response to a decrease in pH, and then transfers the phosphate to ArsR, which then acts to repress or activate expression of downstream genes. Previous work from our lab identified an enhanced biofilm phenotype in strains in which ArsRS was mutated (Servetas et al., 2016). Herein, we demonstrate that homB contributes to ArsRS-induced biofilm formation by H. pylori.

MATERIALS AND METHODS

Bacterial Strains and Growth Conditions

Strains and plasmids used in this study are listed in Table 1. Unless noted otherwise, H. pylori strains were grown as previously described (Carpenter et al., 2015). Briefly, H. pylori stock cultures were maintained at −80°C in H. pylori freezing media [brain heart infusion broth (BD Biosciences) containing 10% fetal bovine serum (FBS) and 20% glycerol (EMD chemicals, Inc)]. Freezer stocks were plated on horse blood agar (HBA) plates comprised of 4% Columbia agar (Neogen Corporation), 2 mg/mL β-cyclodextran (Sigma), and an antibiotic/antifungal cocktail [10 µg/mL vancomycin (Amresco), 5 µg/mL cefsulodin (Sigma), 2.5 U/ml polymyxin B (Sigma), 5 µg/mL trimethoprim (Sigma), and 8 µg/mL amphotericin B (Amresco)]. Following growth on HBA plates, H. pylori strains were grown in liquid culture. Liquid media consisted of brucella broth (Neogen Corporation) supplemented with 10% FBS (Gibco) and 20% glycerol (EMD chemicals, Inc); in addition, liquid cultures were grown shaking at 100 rpm.

Sequencing of homB Promoter Regions

Twenty-one homB positive strains from a collection of South Korean H. pylori clinical isolates (K42, K248, K23, K16, K82, K34, K183, K25, K209, K36, K44, K131, K3, K197, K10, K165, K26, K259, K104, K107, K197) were chosen for primer analysis (Table 1; Kang et al., 2012). homB locus A and locus B were amplified from these 21 clinical isolates and from DSM48 using the AF/AR and BF/BR primer pairs, respectively (Table 2). Sanger sequencing of the promoter regions was carried out with the forward primers (Af and Bf) of each locus; these

| TABLE 1 | Plasmids and strains used in this study. |
| Description | References |
| --- | --- |
| **PLASMIDS** | |
| pPSM 3 | pKSF-II | Copass et al., 1997 |
| pPSM 215 | pTM117 promoterless vector | Carpenter et al., 2007 |
| pPSM 463 | pTM117::G27::HP0073 promoter region | Carpenter et al., 2015 |
| pPSM 920 | pBlueScript:: ArsS::kan | Wen et al., 2006 |
| pPSM 1530 | pGEM-T Easy::HPG27_667::kansacB | This study |
| pPSM 1670 | pGEM-T Easy::G27Δhom | This study |
| pPSM 1532 | pTM117::pUreA-homB | This study |
| pPSM 1538 | pTM117::pFlaA-homB | This study |
| **STRAINS** | |
| DSM 1 | WT G27 | Baltrus et al., 2009 |
| DSM 215 | G27 pTM117, KanR | Carpenter et al., 2007 |
| DSM 983 | G27ΔarsS markerless | Carpenter et al., 2015 |
| DSM 1442 | G27ΔarsS markerless | This study |
| DSM 1443 | G27ΔΔarsSΔhom | This study |
| DSM 1446 | G27 ArsRD52N, unphosphorylatable, KanR | Servetas et al., 2016 |
| DSM 1530 | E.coli top10 pGEM HP27_667::kansacB | This study |
| DSM 1531 | G27 HP27_667::kansacB | This study |
| DSM 1532 | Top10 pUreA-homB | This study |
| DSM 1533 | G27 pUreA-homB | This study |
| DSM 1538 | Top10 pFlaA-homB | This study |
| DSM 1539 | G27 pFlaA-homB | This study |
| DSM 1540 | G27Δhom pFlaA-homB | This study |
| DSM 1543 | H. pylori G27 ArsR-D52N Δhom | This study |
| DSM 1547 | H. pylori G27Δhom pTM117 | This study |
| DSM 1546 | G27Δhom pUreA-homB | This study |

| STRAINS FOR LOCUS A AND LOCUS B PROMOTER ANALYSIS | |
| --- | --- |
| DSM 48 | H. pylori strain 7.13 | Franco et al., 2005 |
| K42 | Korean clinical isolate | Kang et al., 2012 |
| K248 | Korean clinical isolate | Kang et al., 2012 |
| K23 | Korean clinical isolate | Kang et al., 2012 |
| K16 | Korean clinical isolate | Kang et al., 2012 |
| K82 | Korean clinical isolate | Kang et al., 2012 |
| K34 | Korean clinical isolate | Kang et al., 2012 |
| K3 | Korean clinical isolate | Kang et al., 2012 |
| K197 | Korean clinical isolate | Kang et al., 2012 |
| K10 | Korean clinical isolate | Kang et al., 2012 |
| K166 | Korean clinical isolate | Kang et al., 2012 |
| K26 | Korean clinical isolate | Kang et al., 2012 |
| K259 | Korean clinical isolate | Kang et al., 2012 |
| K104 | Korean clinical isolate | Kang et al., 2012 |
| K107 | Korean clinical isolate | Kang et al., 2012 |
| K197 | Korean clinical isolate | Kang et al., 2012 |
promoter sequences were deposited in the NCBI GenBank database with the accession numbers MF572264 to MF572284. The transcriptional start site was identified using the predictive data from Sharma et al. (2010). The homB promoter regions from a handful of H. pylori genomes that are available in Pubmed (SJM180, J99, HUP-B14, Gambia94/24, G27, P12, HPAG1) were also used to supplement the data from the clinical isolates. In total, 30 sequences from 29 strains (both locus A and locus B from strain DSM48 were included in the analysis) that extend 180 bp upstream from the homB start codon were compiled and aligned using the Geneious software 9.1.6 (Biomatters Ltd.) (Supplementary Figure 1). A Weblogo was created based on nucleotide conservation (Figure 1). In addition, the homB promoter region from G27 was also run through the Basic Local Alignment Search Tool (BLAST) to get a more global view of the sequence conservation among homB promoters.

Construction of G27 ΔhomB
All isogenic mutant constructs and strains were created using the H. pylori G27 parental strain (Baltrus et al., 2009). The homB gene in locus A (HPG27_667) of G27 was replaced with kan-sacB. The kan-sacB cassette was amplified with primers K4A and PFA3 from pDSM3 (Copass et al., 1997); primer sequences used in this study can be found in Table 2. The kan-sacB cassette was then fused to the up-stream region of G27 Locus A (amplified with LocA_UpF/LocA_UpR) and the downstream region of G27 Locus A (amplified with LocA_DnF/LocA_DnR) by SOE PCR (Table 2). The full construct was then amplified

| TABLE 2 | Primers used in this study. |
|---|---|
| **Δhom CONSTRUCTION PRIMERS**<sup>A</sup> | **Sequence (5′ − 3′)** | **References** |
| K4A | GTGCCAGCAGCGGCGCCACATTTTGAAGTGA | Copass et al., 1997 |
| PFA3 | GTCTGACCGCCCTTAAGGCGATTTCTCATGC | Copass et al., 1997 |
| LocA_UpF | GTAATCGCTGGTTATTTGAAG | This study |
| LocA_UpR | TCACCTCAAATTGTCGCCGCCGCTGACGACATATCATATCCGATCCTGAC | This study |
| LocA_DnF | GCTATGAAAACTCGGCTTATTGAGCATCAAACGGCTCTTATATAAGC | This study |
| LocA_DnR | ACACATGACTTATTGGA | This study |
| **ArsR D52N CONFIRMATION PRIMERS** | | |
| Up1F | GGGGATTTTTTGAGCGTTGAG | Servetas et al., 2016 |
| Dn4R | CGCAAACGGCCAATGATCAC | Servetas et al., 2016 |
| **ΔarsS CONFIRMATION PRIMERS** | | |
| HP0165_del_ver_F | TGAAAGCATTGCGATTGAGA | Servetas et al., 2016 |
| HP0165_del_ver_R | AAAACGGCTTTGATGCCTAA | Servetas et al., 2016 |
| **hom EXPRESSION VECTOR CONTRUCTION PRIMERS**<sup>B</sup> | | |
| G27homB_Fxbal | agagagCTAGAATGAGAAAAACTATTCGACCTCGCTTTTTATATTTC | This study |
| G27homB_RPstl | agagagCTGACGGCGACGCTCAGAAAAACCCAC | This study |
| G27pfla_A_Fknl | agagagGTGACGCGCCATTTCGCTCTCTATT | This study |
| G27pfla_A_RXba | ATGAGAAAACTATTCATCCCACTTTTATTATTC | This study |
| **PROMOTER SEQUENCING PRIMERS** | | |
| A1 (F1-jhp0648/HP0709) | TAATTTGCGACAAAACATC | Oleastro et al., 2006 |
| Ar (R1-jhp0650/HP0711) | ATCCAGGCGCTAATGAC | Oleastro et al., 2006 |
| BI (F1-jhp0868/HP0935) | AAGAAGTAGGCGCTGCCGAGAAGTTG | Oleastro et al., 2006 |
| Br (R1-jhp0871/HP0936) | GGTTGCCCTTTGGGCTTGGA | Oleastro et al., 2006 |
| **qRT-PCR PRIMERS**<sup>C</sup> | | |
| G27_16S RT-F | ATGGATGCTAGTTGTTGGAGGGCT | Gilbreath et al., 2012 |
| G27_16S RT-R | TTAAACCACATGCTCCACCGCTTG | Gilbreath et al., 2012 |
| G27homB_FWD Set 4 | CCCGAGACAGCAATTGATAG | This study |
| G27homB_REV Set 4 | ACACCCCTTGCGTGTAGT | This study |
| **EMSA PRIMERS**<sup>D</sup> | | |
| rpoB EMSA-F | CCAAAAGAGGTTAAAGGAGCG | Carpenter et al., 2009 |
| rpoB EMSA-R | CCTCTCTGTGCTTCTCTAC | Carpenter et al., 2009 |
| hp0166EMSA_5′ | AAAACGATCAAAGGAATTGT | This study |
| hp0166EMSA_3′ | CAATTCGCCTCTCGATG | This study |
| EMSA_G27F | CGTGAATGAAAGCGCTTTTAG | This study |
| GFpLuse_AR | CGCGCTCTAGAGGTATCTCCTTTTTAGTTAGT | This study |

<sup>A</sup> Regions of overlap sequence for SOE PCR are underlined.

<sup>B</sup> Restriction sites are italicized; lower case nucleotides are nonspecific nucleotides added to enhance restriction digest.

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with FideliTaq PCR Master Mix (USB) and cloned into pGEM T-easy. Transformants were detected by blue-white screening and confirmed by PCR and KanR. One colony was archived

The overall conservation of the 184 nucleotides upstream of the homB genes from locus A and locus B are shown. The boxed ATG is the start codon for the homB gene. The +1 transcriptional start site (black arrowhead) and the −10 sequence (underlined) are both indicated. Larger letter, the more highly conserved the corresponding base pair. To indicate nucleotides where substitutions/deletions are present, the letter(s) at these positions do not reach the full height. The overall conservation of each nucleotide position is indicated by the colored bar: green = 100%, tan = 30–99%, red < 30%.

FIGURE 1 | WebLogo of the homB promoter region. An alignment and subsequent WebLogo were generated from sequences from 30 homB loci from 29 strains.

The G27 ArsR-D52N ΔhomB strain was constructed in a similar manner. First, The ArsR-D52N KanR SOE cassette from DSM1446 was PCR amplified with Up1F and Dn4R (Table 2). The PCR product was used to transform DSM1442. Colonies were selected on HBA+Kan25. The D52N mutation in ArsR was confirmed by PCR amplification of arsR followed by Apol digestion of the amplicon. One colony with the correct digestion pattern was confirmed by sequencing and archived as DSM1543.

Construction of homB Over Expression Strains

Plasmids containing homB fused to the ureA or flaA promoters were generated to assess the effect of increased homB expression on biofilm formation. These plasmid were constructed using the pTM117 plasmid described by Carpenter et al. (2007). First, pDSM463 (pTM117+ureA promoter) was digested with XbaI and PstI to drop out the 730 bp GFP gene. homB was amplified from G27 with G27_homB_FXbaI and G27_homB_RPstI (Table 2). The homB amplicon was digested with PstI and XbaI and was then ligated with the similarly digested pDSM463. The ligation was transformed into Top10 E. coli and transformants were selected on LB+Kan25. KanR colonies were screened by PCR for the presence of homB. Upon confirmation, one colony was archived as DSM1532. pDSM1532 (pUrea-homB) was used to transform H. pylori G27 (DSM1) and transformants were selected on HBA+Kan25. homB over-expression was confirmed by qRT-PCR (primers listed in Table 2) for several KanR colonies. Similar levels of homB expression were observed among all tested colonies (data not shown) and one strain was archived as DSM1533 (G27 pUrea-homB). To construct the pflaA-homB vector, pflaA was first amplified with G27pflaA_FKpnl and G27pflaA_RXbaI from the pDSM3 plasmid (Table 2). The
amplicon and pDSM1532 were digested with KpnI and XbaI. Digestion of pDSM1532 resulted in the release of the ureA promoter, which was then replaced with the similarly digested flaA promoter. Ligation products were transformed into Top10 E. coli and selected for on LB+ Kan25. KanR colonies were screened for the correct promoter by PCR and sequencing. One control colony was archived as DSM1538. DSM1 was transformed with the pflaA-homB vector, pDSM1538, and the resulting transformant was archived as DSM1539 (G27 pFlaA-homB). Similarly, pDSM215, pDSM1532, and pDSM1538 were transformed into G27 homB (DSM1442) and a single transformant for each was archived: DSM1547 (G27 ΔhomB pTM117), DSM1546 (G27Δ homB pUreA-homB) and DSM1540 (G27Δ homB pFlaA-homB), respectively. Plasmid and strain descriptions can be found in Table 1.

**H. pylori Growth Conditions for RNA Isolation**

Briefly, bacterial cultures were patched from freezer stocks onto HBA plates. After 48 h of plate growth, sterile swabs were used to resuspended bacteria in 5 mL of H. pylori liquid media. After 24 h, cultures were standardized to an OD600 of 0.05 (OD control culture) in fresh liquid media and allowed to grow for a specified length of time depending on the experiment. To analyze growth dependent expression, one 60 mL OD control culture was separated into 2-125 mL flasks, each containing 30 mL, and cells were collected from alternating flasks at each indicated time point. At each time point, 5 mL of bacterial culture was removed, captured on 0.45 µm filters were stored at −80°C until RNA preparation. Shock experiments were carried out as previously described (Merrell et al., 2003; Carpenter et al., 2015). Briefly, OD control cultures were prepared in a similar manner as described above. 10 mL OD control cultures were collected after 20 h of growth, and separated into two equal portions. One portion was immediately collected on a filter and represented the T0 pre-shock sample. The second portion was resuspended in 5 mL of media, which represented one of three environmental shocks: low pH (pH 5), excess nickel (10 µM NiSO4; Sigma-Aldrich), or limited iron [200 µM 2-’-dipyridyl (DPP)]. pH shock and excess nickel were maintained for 90 min, and iron limitation was carried out for 60 min. All cultures were incubated under microaerobic conditions with shaking at 100 rpm as described under the growth condition section. After the respective incubation period, cultures were collected by the filter method, flash frozen and stored at −80°C.

To examine homB expression in biofilm and planktonic cells, biofilms were cultured as previously described; the method is detailed below and described in Servetas et al. (Servetas et al., 2016). Samples were collected for RNA isolation after 24, 48, and 72 h of biofilm growth. At each time point the biofilm and planktonic cells were processed separately. Planktonic cells from three technical replicate wells were pooled, harvested by filtration, and stored at −80°C. Following removal of planktonic cells, each well was washed three times with 2 mL of warm liquid media. After washing, 1.5 mLs of TRIzol reagent (Gibco/BRL) was added to the well, and samples were incubated for 20 min at room temperature to lyse the biofilm cells. The TRIzol from three wells was pooled and stored at −80°C. A total of 4–5 biologically independent biofilm/planktonic samples were cultured and analyzed.

**RNA Isolation, cDNA Synthesis and Quantitative Real-Time PCR**

RNA isolation was conducted as previously described (Thompson et al., 2003). RNA integrity was determined by visualization on a 2% agarose gel. cDNA synthesis and Quantitative Reverse-Transcription PCR (qRT-PCR) were conducted as previously described with a few modifications (Gilbreath et al., 2012). Each cDNA synthesis reaction was carried out using the Quantitect reverse transcriptase kit (Qiagen) and 500 ng of RNA. A control reaction, to ensure there was no genomic DNA contamination, was carried out for each sample by excluding the reverse transcriptase enzyme (NoRT). Following cDNA synthesis, qRT-PCR was performed using an intercalating dye, SYBR green, for homB expression and the 16S rRNA internal reference gene. qRT-PCR primers can be found in Table 2. The qRT-PCR reaction mixture was comprised of 1x SYBR green RT-PCR master mix, 3 pmol of each forward and reverse primer, and 1 µl of template (cDNA or the NoRT control). Reactions were brought to a final volume of 20 µl with DNase free water (IDT). The following 3-step cycling protocol was used: a 10 min hold at 95°C (initial denaturation) was followed by 40 cycles of 10 s at 95°C (denaturation), 10 s at 58°C (annealing), and 10 s at 72°C (extension). SYBR green fluorescence was measured at the end of the extension step. Relative gene expression of each gene compared to the 16S internal control was calculated by the 2−ΔCT method; relative fold difference in gene expression was calculated using the 2−ΔΔCT method. Gene expression studies represent a minimum of three biologically independent experiments.

**Purification of ArsR**

Recombinant ArsR (rArsR) was purified from DSM1177 (Harvey et al., 2014). ArsR expression from DSM1177 was induced as previously described (Harvey et al., 2014). Briefly, a 20 mL culture of DSM1177 was grown overnight in Luria-Bertani (LB) (Difco) broth containing 2xM (100 µg/mL) and Kan20 (20 µg/mL) broth. This culture was subcultured 1:50 into 500 mL of prewarmed LB containing Amp100 and Kan20, and grown with shaking (200 rpm) at 37°C until the OD600 reached between 0.5 and 0.7. At this point, expression was induced using 1mM isopropyl-β-D-thiogalactopyranoside (IPTG) and the culture was placed in a 28°C incubator and grown overnight with shaking at 200 rpm. Following overnight culture, the 500 mL culture was pelleted in two 250 mL volumes. rArsR purification was carried out as previously published (Wen et al., 2006) with several modifications. The pellet was resuspended in native binding buffer (50 mM NaH2PO4, 500 mM NaCl, 10 mM Tris-HCl, pH 8.0) and stored at −80°C until the purification process. Pellets were thawed on ice, treated with protease inhibitor tablets (IDT) and DNase I (Sigma), and then lysed by French press. Lysate
were spun twice to remove cellular debris before loading to a Ni-NTA column (Qiagen) for purification by gravity flow. Approximately 1 mL of Ni-NTA (0.5 mL packed column volume) was used to prepare a column according to the manufacturer directions. The lysates were added to the column and then the flow through was re-loaded to the packed bed volume. The column was then washed sequentially with native binding buffer containing increasing concentrations of imidazole: 40 mL 10 mM imidazole, 40 mL of 20 mM imidazole, 10 mL 30 mM and 10 mL 40 mM imidazole. ArsR was then eluted using 4 mLs of 50 mM imidazole followed by 4 mL 100 mM imidazole. Fractions from wash and elution steps were checked on SDS-Page gel by total protein staining and Western blot analysis to assess purity. The 50 mM fraction was pooled, concentrated and buffer exchanged into native binding buffer using Amicon 10 kD MWCO filtration devices and then quantified. The final concentration of rArsR was calculated to be 327.3 µg/mL.

**ArsR Electromobility Shift Assay (EMSA)**

The promoter regions for *H. pylori* G27 homB, arsR (positive control), and rpoB (negative control) were amplified using the primers indicated in Table 2. Promoter region amplicons were end labeled with [³²P] ATP (Perkin Elmer) as previously described (Gancz et al., 2006). Briefly, 150 ng of each amplicon was combined with 5 µl of [³²P] ATP and T4 polynucleotide kinase (NEB). To remove unincorporated nucleotides, reactions were cleaned with the MinElute Reaction Clean-up Kit (Qiagen) and eluted with 30 µL EB. 60 µL of EMSA binding buffer (25 mM NaPO4 (pH 7), 150 mM NaCl, 0.1 mM MgSO4, 1 mM DTT, 0.6 ug/mL BSA, 200 ng/µl salmon sperm, 4% glycerol) was then added to the labeled templates.

Prior to each EMSA trial, ArsR was phosphorylated in vitro using a lithium potassium acetyl phosphate high energy phosphate donor (Sigma-Aldrich) (Carpenter et al., 2015; Kinoshita-Kikuta et al., 2015). rArsR was diluted 1:2 in phosphorylation buffer (0.3 M Tris-HCl, 50 mM KCl, 10 mM MgCl2, 1 mM DTT, and 40 mM acetyl phosphate). The phosphorylation was carried out at 25°C for 60 min. Immediately following phosphorylation, ArsR−P was serially diluted in EMSA binding buffer and combined with the appropriate EMSA template. For each template, in addition to a no protein control reaction, a cold competition reaction in which 2x unlabeled promoter template was added in addition to labeled template was conducted. All reactions were allowed to incubate for 30 min at room temp. Following incubation, templates were loaded onto 6% DNA retardation gels (Invitrogen) and run for 1 h at 100 V in 0.5X TBE. Gels were then exposed to phosphor screens overnight and scanned on a Storm 860 Scanner (GE Healthcare), followed by analysis using ImageJ64 (NIH).

**Biofilm and SEM Analyses**

Biofilm assays were conducted as previously described (Servetas et al., 2016). Briefly, biofilms were grown in a 24-well tissue culture treated plate. Cultures were added to each well to an OD₆₀₀ of 0.1 in 1 mL of HBA liquid media. Biofilms were allowed to develop for 24, 48, and 72 h. At each time point, planktonic cells were aspirated, wells were washed twice with PBS, biofilms were fixed with methanol, and then stained with crystal violet. Crystal violet was rinsed away 3x with distilled water and then solubilized to assess relative quantity of biofilm. Solubilized crystal violet was read at OD₆₀₀. Data shown represent a minimum of 3 biological replicates. Scanning electron microscopy (SEM) assays were performed as previously described (Servetas et al., 2016). Fresh liquid media in 12 or 24 well plates was inoculated as a 1:10 dilution with cells from overnight cultures (0.3–0.7 OD₆₀₀). Cell-cultured treated cover-slips were placed at the bottom of each well to capture biofilm formation. Samples were treated and visualized by SEM as previously described (Gaddy et al., 2009). SEM images are representative of three biological replicates.

**Statistical Analysis**

*homB* expression over time was analyzed by two-way ANOVA with Tukey's correction. The comparison of *homB* expression between biofilm and planktonic cells was conducted using a paired analysis two-way ANOVA with Sidak's correction for multiple comparisons. The rate of biofilm formation, which is defined as the first time point where the crystal violet quantification was significantly greater than the blank, was determined by a two-way ANOVA with Dunnett's correction. Differences in amount of biofilm formation between strains at each time point were analyzed using a one-way ANOVA with Tukey's correction for multiple comparisons. All analyses were conducted with GraphPad Prism 6.

**RESULTS**

The *homB* Promoter Region Is Highly Conserved Regardless of Locus

*HomB* can be found at two different chromosomal loci (Oleastro et al., 2009a; Servetas et al., 2018). To assess *homB* promoter variability based on the chromosomal locus, we first used BLAST to determine the overall conservation between the *homB* promoter regions available in Genbank. We found that the promoter regions at both loci were highly conserved. For instance, the BLAST analysis using the entire 228 bp intergenic region upstream of *homB* (locus A) from G27 returned matches with complete coverage that ranged from 100 to 91% identity. When this comparison was extended to include strains that carried a *homB* gene at locus B, approximately 85% coverage with identity scores ranging from 99 to 88% were seen. In some rare cases, even when the locus was empty, parts of the *homB* promoter region could still be found. In at least one strain (sequence ID: CP006610.2) although locus B was empty, parts of the *homB* promoter region (approximately 180 bp upstream from the *homB* translational start site) were aligned. The pairwise alignment found a range of 80–100% identity between individual promoters (Supplementary Figure 1, Figure 1). Visualization of the conservation as a weblogo revealed that in addition to regions such as the −10, which is 100% conserved among the 30 loci,
there were several other highly conserved regions (Figure 1). This high degree of conservation in the intergenic promoter region regardless of chromosomal locus could suggest that this area is conserved to maintain binding sequences for regulatory proteins.

**homB Expression Is Modulated by Growth Phase and Environmental Cues**

There is currently little data regarding the expression of homB; however, immunologic data suggests that homB is expressed during infection (Oleastro et al., 2008). To explore homB expression, qRT-PCR was used to quantify homB transcription levels over a standard 48 h in vitro growth curve of H. pylori G27 (Figure 2A). Relatively low and consistent levels of expression over the first 30 h of growth were observed, followed by an approximately 3-fold increase in expression at 48 h. Although the observed increase at 48 h was not statistically significant, these data suggest that homB expression is increased in stationary phase (Figure 2A). Next, we sought to determine what regulatory factor(s) were responsible for modulation of homB expression. To this end, homB expression was assessed in the absence of each of the following well characterized transcriptional regulators: Fur, NikR, and ArsS (Figure 2B). Fur and NikR both serve as metallo-regulators that respond to changes in iron and nickel, respectively; ArsS, is the sensor component of a two component system, ArsRS, which functions in response to changes in pH. While the pattern of homB expression observed in the Δfur background was similar to the wild-type strain (Figure 2B), homB expression was increased in both the ΔnikR and ΔarsS strains throughout the growth curve. Specifically, in the ΔnikR strain background, an average of a two-fold increase in homB expression was observed at all time-points as compared to wild-type. Furthermore, in the ΔarsS strain, transcript levels of homB were elevated between four- and ten-fold in comparison to wild-type over the course of the experiment. Furthermore, where wild-type homB expression peaked at 48 h, deletion of arsS resulted in a shift in maximal homB expression from the stationary phase to mid-log phase (Figure 2B). In the ΔarsS strain, a significant increase in expression was observed at 20 h compared to 6 h (p = 0.0086), 12 h (p = 0.0075), 30 h (p = 0.0295), and 48 h (p = 0.003) (Figure 2B). Together, these data suggest that NikR and the ArsRS two-component system play roles in regulation of homB expression.

Given the increase in homB expression observed in the ΔnikR and ΔarsS strain backgrounds, we next asked how homB expression changed in response to the environmental cues that are sensed by these regulators, nickel and pH, respectively. In wild-type H. pylori, a significant, approximately five-fold decrease in homB expression following a 90 min shock with 10 μM Ni²⁺ was observed (Figure 3A). Unexpectedly, in the ΔnikR strain background a similar decrease occurred. If NikR were responsible for the decrease in homB expression, one would expect homB expression in the ΔnikR strain to remain unaffected by the presence of nickel; however, no difference in homB expression between wild-type and the

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**FIGURE 2** | homB temporal expression in wild-type and regulator-deficient mutant strains. (A) homB expression profile in wild-type Helicobacter pylori strain, G27. The relative expression of homB as compared to the 16S gene was calculated for each time point shown on the x-axis. The solid line is plotted at the mean of 3 biologically independent experiments; errors bars represent the standard error of the mean. (B) homB expression was monitored over time in strains lacking Fur, NikR, and ArsR. The fold difference

(Continued)
\( \Delta \text{nikR} \) strain was observed (\( p = 0.3081 \)) suggesting \( \text{homB} \) is still regulated even in the absence of NikR. Together, these data suggest that the decrease in \( \text{homB} \) expression in response to excess nickel is NikR-independent. In regards to pH, when wild-type \( H. \text{pylori} \) was subjected to a pH of 5 for 90 min, a 10-fold decrease in \( \text{homB} \) expression as compared to pH 7 was observed (Figure 3B). Conversely, when the \( \Delta \text{arsS} \) strain was similarly exposed to pH 5, little to no decrease in \( \text{homB} \) expression was seen. This finding suggests that \( \text{homB} \) repression at low pH is ArsS-dependent (Figure 3B).

As mentioned, ArsS is the sensor kinase portion of the two-component system, ArsRS. In response to decreasing pH, ArsS phosphorylates ArsR and ArsR→P can then regulate gene expression. Ideally, this regulatory cascade would be assessed by knocking out \( \text{arsR} \); however, this gene is essential and deletion of \( \text{arsS} \) is often used as a proxy. Another means of assessing this regulatory pathway is via the use of a non-phosphorylatable ArsR-D52N mutant strain; this strain is still viable but ArsR is unable to be phosphorylated in response to low pH. Given the dramatic effect that loss of ArsS had on \( \text{homB} \) expression, we next tested the same conditions using an ArsR-D52N mutant strain. In agreement with the increase in \( \text{homB} \) expression (Figure 2D). Conversely, when the \( \Delta \text{arsS} \) strain was similarly exposed to pH 5, little to no decrease in \( \text{homB} \) expression was observed in the wild-type strain following exposure to \( 10 \mu \text{M Ni}^{2+} \). A significant decrease in \( \text{homB} \) expression in the \( \Delta \text{arsS} \) mutant strain was also detected during log phase (Figure 2C). Additionally, when ArsR-D52N was exposed to pH 5 shock, there was no decrease in \( \text{homB} \) expression (Figure 3B). These data further support the hypothesis that \( \text{homB} \) expression is repressed by ArsRS in response to low pH. Also of note, when \( \text{arsS} \) was expressed in trans (complemented) in the \( \Delta \text{arsS} \) strain background (\( \Delta \text{arsS-arsS}^{C} \)), decreased \( \text{homB} \) expression in response to low pH was restored.

To determine if ArsR was directly regulating \( \text{homB} \) expression via interactions with the gene's promoter, ArsR-promoter interactions were assessed by EMSA. Given that ArsR is auto-regulatory, we used the \( \text{arsR} \) promoter region as a positive control (Dietz et al., 2002). In addition, the promoter of \( rpoB \) was used as a negative control to show absence of non-specific binding. As shown in Figure 4, a clear shift in the banding pattern was observed for both the \( \text{arsR} \) and \( \text{homB} \) promoter regions (Figure 4). In comparison, no shift was observed for \( rpoB \) (Figure 4). Of note, a shift in the \( \text{homB} \) promoter was observed even at low concentrations of rArsR→P, whereas the interaction...
with the *arsR* promoter region was only evident at higher concentrations. Finally, interaction with the *arsR* and *homB* promoter regions was specific as the addition of excess unlabeled promoter fragment was able to shift the band migration to the same area as the no protein control.

### HomB Contributes to Hyper Biofilm Formation in ArsRS Mutant Strains

We recently showed that strains with a mutation in either portion of the ArsRS two-component system (Δ*arsS* or ArsR-D52N) form significantly more biofilm and do so at an enhanced...
rate as compared to wild-type H. pylori (Servetas et al., 2016). Given that homB was highly over expressed in both the ΔarsS and ArsR-D52N strains (Figure 2C), we reasoned that increased expression of this OMP could contribute to the previously observed biofilm phenotype. Visual analysis of these samples revealed dramatic strain dependent differences in crystal violet staining on the bottom of the wells (Supplementary Figure 2). Furthermore, crystal violet assay quantification of biofilm formation, showed similar results as in our previous studies (Servetas et al., 2016); significant biofilm formation occurred earlier in the ΔarsS and ArsR-D52N strains (24 h) as compared to wild-type (48 h) (Figure 5). Deletion of homB from the ΔarsS or ArsR-D52N strains negated the early biofilm phenotype; significant biofilm formation was not seen until 48 h in the double mutant strains (Figure 5). In addition, by 48 and 72 h, ΔarsS and ArsR-D52N formed significantly more biofilm than wild-type, respectively, whereas ΔarsS ΔhomB and ArsR-D52N ΔhomB did not (Figure 5, Table 3). Overall there was less biofilm formation in all of the ΔhomB strains as compared to their cognate homB positive strains; however, these differences were not always statistically significant (Figure 5, Table 3).

**homB Expression Is Increased During Early Biofilm Formation and Over Expression Enhances Biofilm Formation**

Given the above results that indicated that homB contributes to hyper-biofilm formation, we next asked if homB expression was altered in wild-type biofilm cells as compared to planktonic cells. To assess this, we compared homB expression in biofilm associated bacteria to the planktonic bacteria from the same culture. At 24 h, we observed a significant increase in homB expression in cells within the biofilm as compared to the planktonic cells (Figure 6); by 48 h, this difference was no longer observed. These findings may suggest that increased homB expression helps to establish early biofilm formation; cells expressing higher levels of homB may adhere and initiate the biofilm process. Of note, significant biofilm formation is not observed with wild-type cells until; missing space between cell and until 48 h, which is when the disparity in homB expression between biofilm and planktonic cells is no longer observed (Figures 5, 6).

Thus far, we have shown that homB is regulated by the ArsRS two component system and that loss of ArsRS regulation leads to an increase in homB expression and a higher level of homB expression at earlier time points (Figure 2). Furthermore deletion of homB reversed the hyper-biofilm phenotype in the ArsR-D52N and ΔarsS backgrounds. Given these observations, we next asked if increased homB expression in either a wild-type or ΔhomB strain was sufficient to induce the enhanced biofilm formation that was observed in the ArsR-D52N and ΔarsS mutant strains. To accomplish this goal, constructs where homB expression helps to establish early biofilm formation; cells expressed between 4-fold to 10-fold higher than the wild-type homB was not sufficient to completely recapitulate the dramatic enhancement of biofilm formation that was observed in the ArsR-D52N or ΔarsS strains in this assay (Figure 8). There was a trend toward increased biofilm formation by G27 pFlaA-homB and G27 pUreA-homB at 48 and 72 h and at 72 h there was significantly the ureA promoter is known to be highly active in H. pylori (Bauerfeind et al., 1997), both promoters resulted in comparable levels of homB expression in the WT background. In contrast, higher levels of homB expression were observed in G27 ΔhomB pUreA-homB compared to G27 ΔhomB pFlaA-homB (Figure 7). Furthermore, despite the lack of a native copy of homB, the levels of homB expression in the G27 ΔhomB pUreA-homB were similar to the levels of homB expression seen in G27 pUreA-homB. These data suggest that there is a maximum level of homB transcription that can be tolerated in H. pylori.

Overall, we were able to obtain strains in which homB was expressed between 4-fold to 10-fold higher than the wild-type strain. We first assessed the biofilm formation of these strains in the crystal violet assay and found that the over-expression of homB was not sufficient to completely recapitulate the dramatic enhancement of biofilm formation that was observed in the ArsR-D52N or ΔarsS strains in this assay (Figure 8). There was a trend toward increased biofilm formation by G27 pFlaA-homB and G27 pUreA-homB at 48 and 72 h and at 72 h there was significantly.

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**Table 3**: Statistical analysis of biofilm formation between strains carrying ArsRS mutations.

| Comparison | P-valueA | Summary |
|------------|----------|---------|
| 24 HOURS   |          |         |
| G27 vs. G27 ΔhomB | >0.9999 | ns       |
| G27 vs. G27 ΔarsS   | 0.0604 | ns       |
| G27 vs. G27 ArsR-D52N | 0.059  | ns       |
| G27 vs. G27 ΔarsS ΔhomB | 0.7348 | ns       |
| G27 vs. G27 ΔhomB ArsR-D52N | >0.9999 | ns       |
| G27 ΔarsS vs. G27 ArsR-D52N ΔhomB | 0.7416 | ns       |
| G27 ArsR-D52N vs. G27 ΔhomB ArsR-D52N | 0.9973 | ns       |
| G27 ΔhomB vs. G27 ΔarsS ΔhomB | 0.6305 | ns       |
| G27 ΔhomB vs. G27 ArsR-D52N ΔhomB | 0.9977 | ns       |
| 48 HOURS   |          |         |
| G27 vs. G27 ΔhomB | 0.8478 | ns       |
| G27 vs. G27 ΔarsS | 0.001  | **       |
| G27 vs. G27 ArsR-D52N | 0.2185 | ns       |
| G27 vs. G27 ΔarsS ΔhomB | 0.873  | ns       |
| G27 vs. G27 ΔhomB ArsR-D52N | 0.9994 | ns       |
| G27 ΔarsS vs. G27 ΔarsS ΔhomB | 0.0355 | *        |
| G27 ArsR-D52N vs. G27 ΔhomB ArsR-D52N | 0.4449 | ns       |
| G27 ΔhomB vs. G27 ΔarsS ΔhomB | 0.1702 | ns       |
| G27 ΔhomB vs. G27 ArsR-D52N ΔhomB | 0.5948 | ns       |
| 72 HOURS   |          |         |
| G27 vs. G27 ΔhomB | 0.8825 | ns       |
| G27 vs. G27 ΔarsS | <0.0001| ****    |
| G27 vs. G27 ArsR-D52N | 0.0434 | *        |
| G27 vs. G27 ΔarsS ΔhomB | 0.2289 | ns       |
| G27 vs. G27 ΔhomB ArsR-D52N | 0.9953 | ns       |
| G27 ΔarsS vs. G27 ΔarsS ΔhomB | 0.0419 | *        |
| G27 ArsR-D52N vs. G27 ΔhomB ArsR-D52N | 0.1782 | ns       |
| G27 ΔhomB vs. G27 ΔarsS ΔhomB | 0.0132 | *        |
| G27 ΔhomB vs. G27 ΔhomB ArsR-D52N | 0.5214 | ns       |

A: ns: p > 0.05; *: p ≤ 0.05; **: p ≤ 0.01; ***: p ≤ 0.001.
more biofilm in the G27 pFlaA-homB and G27 pUreA-homB as compared to the control (Figure 8A, Table 4). Additionally, both G27 pFlaA-homB and G27 pUreA-homB displayed significant biofilm formation by 48 h, in comparison to the empty vector control which did not display significant biofilm formation until the 72 h time point. Similarly, in the G27 ΔhomB background, there was a trend toward increased biofilm formation in the pFlaA-homB and pUreA-homB containing strains (Figure 8B, Table 4). These data suggest that over expression of homB may enhance the overall amount of biofilm formation but, this alone was not sufficient to recapitulate the enhanced biofilm formation seen in this assay when the ArsRS system is mutated.

Biofilm SEMs Reveal Dramatic homB-Dependent Differences in Biofilm Formation

Given that crystal violet staining is not the most sensitive measure of biofilm formation, and given that there are sometimes apparent visual differences in biofilm staining (Supplementary Figure 2), we next used SEM imaging as a qualitative and semi-quantitative approach to directly visualize biofilm formation of the isogenic mutant strains (Figure 9, Supplementary Figure 2). When wild-type H. pylori were viewed at the lower magnification, we observed slight biofilm formation and numerous adherent bacteria (Figure 9A). In comparison, less biofilm and adherent bacteria were observed for the ΔhomB strain (Figure 9A). As expected, striking biofilm formation was observed for both of the ArsRS mutant strains. Deletion of homB in both of the strains drastically reduced biofilm formation (Figure 9A). Furthermore, a decrease in biofilm formation was also observed when arsS was supplemented in trans (ΔarsS-arsS′) (Figure 9A).

SEM analysis also revealed a dramatic difference between the empty vector control strain (G27 pTM117) as compared to the homB over expression strains (Figure 9B). All strains containing vectors that over expressed homB displayed enhanced biofilm formation; these levels were similar to what was seen in the ArsRS mutant strains (Figure 9A). Similarly, addition of either homB expression vector into the G27ΔhomB background resulted in biofilm formation that was similar to the other hyper-biofilm forming strains. Of note, the presence of biofilm matrix, flagella, coccoid H. pylori, and possible outer membrane vesicles (OMVs) were visible at higher magnification (Supplementary Figure 3). Finally, while both episomal homB expression and mutation of the ArsRS system induced aberrant biofilm formation, there did appear to be slight architectural differences between the ArsRS mutant strain biofilms and the biofilms formed when homB was expressed in trans (Figures 8, 9). Despite these differences, the SEM data suggest that homB is intricately involved in the biofilm formation of H. pylori.

DISCUSSION

H. pylori persistently colonizes approximately one half of the world’s population. In order to persist within the gastric niche, H. pylori must adapt to its host and to any changes that occur within that host throughout the course of infection. Given their location, components that make up the cellular envelope, including OMPs, provide key host-pathogen interactions. Comparative studies of sequenced H. pylori genomes have identified at least 60 OMPs. Many of these OMPs are among the auxiliary genes that are present in only a subset of strains. HomB, is one of the auxiliary OMPs that is associated with severe
disease. Herein, we evaluated homB expression patterns, which we reasoned could provide clues about its possible role during H. pylori infection.

Initial investigations found that homB could be present at two possible loci, and that strains could harbor zero, one, or two copies of homB (Oleastro et al., 2008, 2009a, 2010; Kang et al., 2012; Kim et al., 2015). Other OMPs such as the bab alleles (babA, babB, and babC) and the sab alleles (sabA, sabB, and hopQ) also occupy multiple possible loci and can vary in copy number. It is hypothesized that homologous recombination
between loci is one way that *H. pylori* regulates OMP expression, which helps it to adapt to its environment (Oleastro et al., 2010). For instance, homologous recombination between *sab* loci has been demonstrated in *vitro* and results in gene duplication (Talarico et al., 2012). Bab genes can also undergo homologous recombination resulting in the formation of chimeric bab genes (Backstrom et al., 2004; Solnick et al., 2004). Additionally, a dinucleotide repeat in the 5′ coding region of bab locus B leads to phase variation of bab alleles present at this locus (Backstrom et al., 2004; Solnick et al., 2004). Translational phase variation is also observed in SabA and SabB (Goodwin et al., 2008). In contrast to the bab and sab loci, homologus recombination leading to gene duplication or altered regulation has not been demonstrated for homB. We found that, the homB promoter regions were highly conserved regardless of locus, and unlike many other OMPs, examination of the promoter and coding regions of homB revealed no mono/dinucleotide repeats that are typically indicative of slip-strand mispairing or phase variation (Figure 1, Supplementary Figure 1; Salaun et al., 2004). It is worth noting that another member of the Hom protein family, HomA, can also be found at locus A and/or locus B. Prior *in silico* analysis revealed regions of homologous recombination in the middle sections of the homB and homA genes, which account for the variability among these OMPs (Oleastro et al., 2010); however, this recombination would not account for movement of genes between loci or the conservation of the upstream regions. To our knowledge, such high levels of conservation of the promoter regions of genes found in different loci is not observed for other OMP families. Furthermore, it is unclear if the presence of two homB genes versus a single homB gene is the result of a single historical duplication event or if this process is fluid during the course of an infection. Our findings suggest future research is needed to elucidate the molecular mechanisms underpinning the genetic rearrangements of homB at locus A and locus B. Additionally, we note that while the promoter regions appear identical, the expression of homB within a strain that carries homB at locus B has yet to be evaluated. Similarly, homB expression has not been evaluated in strains carrying homB at both loci. Thus, while the data presented here answer some questions about homB expression, there are clearly many questions that remain to be answered.

One reason intergenic DNA may be preserved is to maintain the integrity of protein binding sequences that are important for gene regulation. Herein, ArsRS was shown to be a potent regulator of homB expression. In fact, in response to low pH, homB expression was down-regulated nearly ten-fold in an ArsRS dependent manner. Conversely, despite the fact that homB was repressed under conditions of excess nickel, NikR did not appear to directly mediate this change (Figure 3A). Though not completely clear, the nickel responsive change in homB expression may in fact be due to changes in regulation of *arsR*; the coordinated regulation of *arsR* expression by Fur and NikR was recently reported (Roncarati et al., 2016). That study found that in the presence of iron, Fur can bind to the *arsR* promoter to repress expression; however, in excess nickel conditions, NikR interferes with Fur repression of *arsR*, which leads to increased ArsR (Roncarati et al., 2016). Thus, it is plausible that the increase in homB expression observed in the nikR deletion strain is due to the role that NikR plays in ArsR regulation. Also of note, mutations in the ArsRS system not only changed the level of homB expression but also changed the pattern of homB expression (Figure 2). In the wild-type strain, an increase in homB expression was observed during the transition from late-log to stationary phase. An increase in gene expression during the log to stationary transition period is not specific to homB and this transition in gene expression has been coined as the “log-stat switch” (Thompson et al., 2003). In fact, numerous virulence factors are among the genes shown to increase during the growth phase transition, suggesting that perhaps homB is up-regulated as part of *H. pylori’s* virulence arsenal (Thompson et al., 2003). In the ArsRS mutant strains a dramatic increase in expression was still observed; however, it occurred during log-phase (Figure 2B). These data suggest that while ArsRS was responsible for suppression of homB expression, another regulatory factor may be involved in the increase in homB expression that was observed at T48 and T20 in the wild-type and Δ*arsS* strains, respectively. It is plausible that expression of homB is regulated both positively and negatively by multiple environmental cues, which opens the opportunity for future investigations. In addition, we observed subtle differences between the ArsR-D52N mutant strains and the Δ*arsS* strain. On average, a higher level of homB expression was observed in the ArsRS-D52N mutant strain as compared to Δ*arsS* strain (Figures 2C, 3B). One explanation for this differences in homB expression could be that a low level of ArsR−P actually remains in the Δ*arsS* strain; ArsR−P could be generated via spontaneous or nonspecific phosphorylation and could then bind to the homB promoter. Conversely, the ArsR-D52N strain would never be able to be phosphorylated spontaneously or nonspecifically; the crucial residue is mutated. Given this possibility, we note that herein we have assessed binding of ArsR−P to the homB

| TABLE 4 | Statistical analysis of biofilm formation between strains carrying homB expression vectors. |
| Comparison | P-value<sup>a</sup> | Summary |
|-------------|-----------------|---------|
| **24 HOURS** |                  |         |
| G27 pTM117 vs. G27 pFla-homB | ns 0.6498 |         |
| G27 pTM117 vs. G27 pUreA-homB | ns 0.9023 |         |
| G27 ΔhomB pTM117 vs. G27 ΔhomB pFla-homB | ns 0.9853 |         |
| G27 ΔhomB pTM117 vs. G27 ΔhomB pUreA-homB | ns 0.9841 |         |
| **48 HOURS** |                  |         |
| G27 pTM117 vs. G27 pFla-homB | ns 0.168 |         |
| G27 pTM117 vs. G27 pUreA-homB | ns 0.2375 |         |
| G27 ΔhomB pTM117 vs. G27 ΔhomB pFla-homB | ns 0.8074 |         |
| G27 ΔhomB pTM117 vs. G27 ΔhomB pUreA-homB | ns 0.7257 |         |
| **72 HOURS** |                  |         |
| G27 pTM117 vs. G27 pFla-homB | ns 0.0006 |         |
| G27 pTM117 vs. G27 pUreA-homB | ns 0.0381 |         |
| G27 ΔhomB pTM117 vs. G27 ΔhomB pFla-homB | ns 0.5599 |         |
| G27 ΔhomB pTM117 vs. G27 ΔhomB pUreA-homB | ns 0.0908 |         |

<sup>a</sup>ns: p > 0.05; *p ≤ 0.05; **p ≤ 0.01.
FIGURE 9 | Continued
Servetas et al. ArsRS and homB Contribute to H. pylori Biofilm

FIGURE 9 | SEM analysis of H. pylori biofilm formation. H. pylori biofilms were visualized by SEM. (A) 1000x SEM images of biofilm formation at 48 h by wild-type G27, ΔarsS, and ArsR-D52N as compared to their isogenic ΔhomB strain derivatives and the arsS complemented strain. (B) 1000X images of biofilm formation at 48 h by G27 and G27ΔhomB strains carrying a control vector (pTM117) or vectors containing homB under control of the ureA or flaA promoters.

promoter. However, future studies should examine the structure function relationships between this interaction and should examine the ability of purified ArsR-D52N to bind the homB promoter using a technique such as fluorescence anisotropy. Such studies will allow us to address difference in the binding kinetics between the different forms of ArsR (ArsR, ArsR∼P, and ArsR-D52N) and could provide valuable information on the exact DNA binding site needed for binding and regulation.

In a recent publication we proposed a model in which ArsRS senses the pH at the gastric epithelium and modulates gene expression to promote colonization and biofilm formation (Servetas et al., 2016). Given the location of HomB in H. pylori’s outer membrane, and the increase in homB expression observed following ArsRS mutation, we reasoned that HomB may contribute to biofilm formation. In fact, at 24 h, a significant increase in homB expression was observed in the biofilm associated bacteria as compared to the planktonic cells (Figure 6). Furthermore, homB was not only integral to the hyper-biofilm phenotype observed in the ArsRS mutant strains (Figures 5, 8), but expression of homB in trans also enhanced biofilm formation in wild-type and the ΔhomB strain backgrounds (Figures 7, 8). Surprisingly, even the subtle increase in homB expression in the G27ΔhomB pFlaA-homB strain (Figure 7), appeared to be sufficient to induce biofilm formation (Figure 9, Supplementary Figure 3). While the level of homB expression was not always dramatically increased, the pattern of homB expression over time was changed in all of the hyper-biofilm strains. In wild-type H. pylori, homB expression didn’t change dramatically until 48 h (Figure 2A) whereas in the ΔarsS strain, the ArsR-D52N strain, or in strains where homB was expressed in trans, homB expression was altered (Figures 2, 6). These data suggest that over expression of homB as well as aberrant timing of homB expression may both contribute to the hyper-biofilm phenotype.

While our data indicate that HomB is a crucial component of the H. pylori biofilm, numerous questions about biofilm formation remain. For example, do other OMPs contribute to biofilm development? This question is of particular interest given
that higher levels of biofilm formation were observed in the ΔarsS strain as compared to the ArsR-D52N strain and given that deletion of homB didn’t completely ablate biofilm formation. While the differences in biofilm between the ArsRS mutant strains may be related to homB, ArsRS regulates numerous genes, including other OMPs. Clearly, biofilm formation is a multifactorial process and it is highly likely that multiple genes that contribute to biofilm formation are affected, either directly or indirectly, by the ArsRS two component system. Thus, the difference in biofilm formation between the two ArsRS mutant strains may be related to the compounding of subtle differences in gene expression between the two strains.

To date, a few other OMPs have been referenced in terms of biofilm formation (Wong et al., 2016; Yonezawa et al., 2017). A high-throughput screen identified the presence of another gene within the Hom family, homD, in association with H. pylori strains that formed robust biofilms; however, no molecular work has been done to verify the role of this OMP in biofilm development (Wong et al., 2016). Conversely, the OMP, AlpB, was evaluated more fully and appears to play a considerable role in biofilm formation in H. pylori strain TK1402 (Yonezawa et al., 2017). Furthermore, mutation of a single domain in AlpB, which is predicted to be in an environmentally exposed loop, affected biofilm development by TK1402 (Yonezawa et al., 2017). Notably, differences between the HomB and HomA paralogs are also predicted to reside in environmentally exposed loops (Servetas et al., 2018). Thus, these findings present an interesting avenue to pursue in the identification of functional differences between HomA and HomB. It is also of note that several other OMPs have been identified within the ArsRS regulon; sabA, labA, and hopZ were all recently shown to be subject to ArsRS-regulation dependent on Acio-Pizzarello et al. (2017). Similar to homB, strains lacking Ars show an increase in expression of sabA, labA, and hopZ (Acio-Pizzarello et al., 2017). Thus, it is plausible that these other ArsRS regulated OMPs may contribute to the biofilm formation that remains in the ΔhomB strains (Figures 5, 7, 8).

In summary, the data presented herein demonstrated that aberrant homB expression was necessary and sufficient to induce hyper-biofilm formation by H. pylori G27. Given the heterogeneity of H. pylori OMP profiles seen across H. pylori strains and the differences in the propensity for biofilm formation seen between H. pylori strains, the work presented here highlights areas for future investigation; what are the roles of other OMPs in biofilm development, and what consequences do variable OMP genotypes play on biofilm formation in H. pylori?

**AUTHOR CONTRIBUTIONS**

SS provided the experimental design, conducted experiments, preparation of manuscript. DM provided the experimental design, preparation of manuscript. AK, J-HC sequenced homB locus A and locus B promoter regions from clinical isolates. RD, JG conducted the SEM studies. IW aided in biofilm experiments.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: [https://www.frontiersin.org/articles/10.3389/fmicb.2018.01497/full#supplementary-material](https://www.frontiersin.org/articles/10.3389/fmicb.2018.01497/full#supplementary-material)

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