

1 **Chemotaxis allows bacteria to overcome host-generated reactive oxygen**  
2 **species that constrain gland colonization**

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12 *Running title: Interplay of ROS and bacterial chemotaxis*

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15

16 **Classification:** Biological sciences; microbiology

17

18 **Abstract:**

19 The epithelial layer of the gastrointestinal tract contains invaginations, called glands  
20 or crypts, which are colonized by symbiotic and pathogenic microorganisms and may  
21 function as designated niches for certain species. Factors that control gland  
22 colonization are poorly understood, but bacterial chemotaxis aids occupation of these  
23 sites. We report here that a *Helicobacter pylori* cytoplasmic chemoreceptor, TlpD, is  
24 required for gland colonization in the stomach. *tlpD* mutants demonstrate gland  
25 colonization defects characterized by a reduction in the percent of glands colonized,  
26 but not in number of bacteria per gland. Consistent with TlpD's reported role in  
27 reactive oxygen species (ROS) avoidance, *tlpD* mutants showed hallmarks of exposure  
28 to large amounts of ROS. To assess the role of host-generated ROS in TlpD-dependent  
29 gland colonization, we utilized mice that lack either the ability to generate epithelial  
30 hydrogen peroxide or immune cell superoxide. *tlpD* gland colonization defects were  
31 rescued to wild-type *H. pylori* levels in both of these mutants. These results suggest  
32 that multiple types of innate immune generated ROS production limit gland  
33 colonization and that bacteria have evolved specific mechanisms to migrate through  
34 this gauntlet to establish in the glands.

35

36 **Significance statement:**

37 Microbial colonization of the gastrointestinal tract occurs at distinct sites within the  
38 tissue including glandular structures found in the stomach and intestine. Multiple  
39 lines of evidence suggest that glands supply niches that promote chronic microbial  
40 colonization, a process that is critical for symbiotic and pathogenic bacteria to

41 maintain themselves. In this report, we show that host-produced reactive oxygen  
42 species (ROS) constrain gland colonization by the gastric pathogen *Helicobacter pylori*.  
43 A bacterial cytoplasmic chemoreceptor, TlpD, allows *H. pylori* to avoid ROS and  
44 enhances *H. pylori*'s ability to colonize a broad swath of glands. We propose that hosts  
45 limit gland access and spread by producing ROS, and bacteria counter with  
46 chemotactic responses that allow navigation through this gauntlet.

47

#### 48 **Introduction:**

49 The epithelium of the gastrointestinal (GI) tract contains invaginations, called  
50 glands in the stomach and crypts in the intestine, which are thought to serve as a  
51 niche for particular microbes and in turn, promote chronic colonization by specific  
52 microbial species. Our knowledge of the factors that control the colonization of these  
53 structures is incomplete. Host factors that have been implicated in controlling gland  
54 colonization include the production of mucus (1), the production of antimicrobial  
55 peptides (2), and the presence of resident immune cells in the lamina propria (3).  
56 Gland colonization, therefore, requires microbes to bypass these defensive strategies.

57 Bacteria too appear to have special adaptations to the gland niche. These  
58 include the ability to use certain carbohydrates (4) and perform chemotaxis (5-7).  
59 The chronically-colonizing gastric pathogen *Helicobacter pylori* is one such microbe  
60 that requires chemotaxis for gland colonization (5-7). Chemotaxis permits bacteria to  
61 sample their environment via chemoreceptors that use ligand-binding signals to alter  
62 the autophosphorylation of a complexed histidine kinase CheA. Ultimately, this  
63 pathway alters flagellar motility to allow bacteria to follow or repel themselves from

64 gradients of specific signals (8). *H. pylori* expresses four chemoreceptors, three of  
65 which (TlpA, TlpB, and TlpC) are embedded the inner membrane, and one that is fully  
66 cytoplasmic (TlpD). The relevance of individual chemoreceptors on overall gastric  
67 colonization has been gauged previously by the level of colonization defect that a  
68 particular mutant displays. Among individual *H. pylori* chemoreceptor mutants, *tlpD*  
69 mutants display the most severe colonization attenuation in two animal models of  
70 infection (9, 10). The exact nature of the *tlpD* mutant colonization deficit, however,  
71 has remained unclear, as has the role of specific signals and chemoreceptors in gland  
72 colonization.

73 TlpD has been linked to a chemotactic response to multiple stress-related  
74 signals including electron transport chain inhibitors (11), acid (7), and reactive  
75 oxygen species (ROS) including hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) or the superoxide  
76 generators metronidazole and paraquat (12). One hypothesis is that these signals are  
77 connected because they all affect oxidative stress experienced in the cytoplasm (12).  
78 Gastric *Helicobacter* are known to encounter host-generated ROS derived from both  
79 epithelial and immune cells during infection and must cope with this stress to  
80 successfully colonize (13, 14).

81 ROS are produced by both gastric epithelial cells and innate immune cells, and  
82 include hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), superoxide (O<sub>2</sub><sup>-</sup>) and hypochlorous acid (HOCl)  
83 (13). To counter these stresses, *H. pylori* possesses a suite of ROS detoxification  
84 systems including catalase, superoxide dismutase, and peroxiredoxins (15). *H. pylori*  
85 mutants lacking these systems are sensitive to ROS and are also attenuated in the  
86 host (15). ROS production limits colonization at epithelial surfaces in the stomach and

87 intestine (14, 16); in agreement with this idea, mouse mutants that lack the epithelial  
88 DUOX enzyme produce less H<sub>2</sub>O<sub>2</sub> and allow elevated colonization by a relative of *H.*  
89 *pylori*, *Helicobacter felis* (14). ROS production may serve to drive microbes away from  
90 the epithelial surface, as microbial adherence to intestinal epithelial cells promotes  
91 H<sub>2</sub>O<sub>2</sub> production and hosts respond to *H. pylori* infection with elevated ROS (15, 16).  
92 However, it is not clear how ROS affects colonization within the glands.

93         To define the contribution of TlpD in gastric colonization, we first determined  
94 its effect on bacterial distribution in the stomach. We found that *tlpD* mutants showed  
95 specific deficits in colonizing a broad swathe of gastric glands, and displayed  
96 hallmarks of exposure to elevated ROS. This result raised the possibility that gland  
97 colonization defects could be due to an inability of *tlpD* mutants to successfully  
98 navigate in response to ROS, an idea that was further supported by the observation  
99 that *tlpD* mutants achieved normal numbers per gland in the glands they colonized.  
100 To assess whether host-generated ROS impacted *H. pylori* colonization, we compared  
101 the colonization and distribution of wild type (WT), *tlpD* and nonchemotactic *cheY*  
102 mutants in mice deficient in either epithelial dual oxidases (*Duoxa*<sup>-/-</sup>) or phagocytic  
103 NOX2 NADPH oxidase (*Cybb*<sup>-/-</sup>). Infection of either *Duoxa*<sup>-/-</sup> or *Cybb*<sup>-/-</sup> mutant mice  
104 rescued the gland colonization defects of *tlpD* mutants noted in WT hosts. Our results  
105 suggest that ROS production impacts *H. pylori* gland transit, and that TlpD-mediated  
106 chemotactic responses are needed to thread this restricted gland access.

107

108 **Results**

109 ***tlpD* mutants have minor colonization defects but achieve normal per gland**  
110 **loads**

111 To begin our analysis of TlpD's role in colonization, we orally infected WT C57BL6  
112 mice with WT, *tlpD*, or *cheY* mutant variants of *H. pylori* that all expressed GFP. *cheY*  
113 encodes the central chemotaxis signaling proteins, so mutants that lack it are fully  
114 non-chemotactic, while mutants that lack *tlpD* lose only responses sensed by that  
115 receptor and thus are partially chemotactic. After two weeks of infection, the total  
116 colonization levels in tissue of the stomach corpus and antrum were determined. *tlpD*  
117 mutants showed colonization defects in the antrum and corpus of WT mice (Fig. 1A),  
118 similar to that previously reported (9). These results suggested that *tlpD* GFP+ *H.*  
119 *pylori* behaved similarly to *tlpD* infections lacking GFP described previously (9), and  
120 encouraged the analysis of gland colonization by the mutant.

121 We next sought to examine TlpD's role in gland colonization. To monitor gland  
122 colonization, we employed the bacterial localization in isolated glands (BLIG)  
123 approach in which gastric glands are isolated from the infected corpus or antrum  
124 tissue, epithelial cells labeled with Hoechst DNA stain, and glands examined for the  
125 presence of GFP+ *H. pylori* by fluorescent microscopy (5). Bacteria within glands were  
126 counted manually, and two parameters of gland colonization were compared  
127 between *H. pylori* strains. The first parameter was gland bacterial load, the number of  
128 bacteria per infected gland. Our calculation of gland bacterial load excludes non-  
129 infected glands. The second parameter was gland occupancy, the percent of glands  
130 infected.

131 In WT mice, WT *H. pylori*-colonized the glands of both the corpus and the  
132 antrum to similar levels, averaging 10 bacteria/infected gland as reported previously  
133 (Fig. 1B) (5). Loss of TlpD did not affect gland load in the antrum but caused a ~1.8-  
134 fold increase in gland load in the corpus compared with WT (Fig. 1B). Full loss of  
135 chemotaxis (*cheY* mutants) also resulted in elevated gland loads of 2- to 3-fold in both  
136 the corpus and the antrum relative to WT *H. pylori* (Fig. 1B). These results suggest  
137 that chemotactic defects did not impair growth within glands, and if anything,  
138 resulted in elevations in bacterial gland load.

139

140 **The loss of *tlpD* or chemotaxis results in a reduction in gland occupancy**  
141 **throughout the stomach in WT hosts**

142 Because *tlpD* and *cheY* mutants appeared to have altered gland phenotypes,  
143 we next analyzed gland occupancy to determine the percentage of glands infected by  
144 *H. pylori*. This frequency likely reflects both the initial population of glands infected  
145 by *H. pylori* as well as the ability to spread and colonize new glands. In WT mice, WT  
146 *H. pylori* colonized 40-50% of corpus and antral glands by two weeks of infection, and  
147 were found in similar proportions in both regions (Fig. 1C). *tlpD* mutants showed an  
148 ~3-fold reduced occupancy in both the corpus and antrum relative to WT *H. pylori*  
149 (Fig. 1C). *cheY* mutant gland occupancy was also decreased relative to WT *H. pylori*,  
150 with significant reductions in both the corpus and the antrum (Fig. 1C). These results  
151 suggest that chemotaxis generally and TlpD specifically is required for *H. pylori* to  
152 occupy new glands.

153

154 ***tlpD* mutants show hallmarks of elevated ROS exposure relative to WT *H. pylori***

155 We reported recently that TlpD mediates chemotactic repellent responses to  
156 multiple ROS (12). Combining this information with our data above suggested that  
157 *tlpD* mutant gland colonization defects could be due to an inability of these mutants  
158 to sense and repel themselves away appropriately from ROS. We therefore asked  
159 whether *tlpD* mutants experienced differential oxidative stress *in vivo*. For this  
160 approach, we used quantitative real-time PCR of mRNA isolated from infected mouse  
161 tissue. We examined the expression of the catalase gene (*katA*) mRNA by *H. pylori*  
162 strains, whose expression has been shown to be sensitive to several oxidative  
163 stresses (17, 18). We determined that this gene was modestly upregulated *in vitro* in  
164 our strains following exposure to 1 mM H<sub>2</sub>O<sub>2</sub> for twenty minutes (Fig. 2A). This result  
165 suggested that *katA* mRNA could serve as a reasonable proxy for H<sub>2</sub>O<sub>2</sub> exposure *in*  
166 *vivo*. We next assessed whether the expression of *katA* mRNA differed between WT,  
167 *tlpD*, or *cheY* *H. pylori* during infection of WT mice. *tlpD* mutants expressed  
168 significantly more *katA* mRNA than WT *H. pylori* in the antrum, and modestly more in  
169 the corpus (Fig. 2B). These results suggest that *tlpD* mutants experience elevated  
170 oxidative stress during infection. Conversely, *cheY* mutants did not express elevated  
171 catalase mRNA (Fig 2B). This outcome suggests that the loss of TlpD specifically leads  
172 *H. pylori* to be exposed to conditions that are different than those encountered by WT,  
173 consistent with high exposure to oxidative stress.

174



175 **Gland colonization defects of *tlpD* are rescued in hosts deficient in H<sub>2</sub>O<sub>2</sub>**  
176 **production by gastric epithelial cells.**

177 The results presented above suggest that TlpD helps to mitigate exposure of *H.*  
178 *pylori* to oxidative stress in the mouse. In order to follow up on oxidative stress and  
179 its role in TlpD-mediated colonization, we next infected two mutant mouse hosts that  
180 were deficient in the production of H<sub>2</sub>O<sub>2</sub> and O<sub>2</sub><sup>-</sup> production. The first of these lacks  
181 the dual oxidase (Duox) heterodimeric enzyme complex by virtue of loss of the  
182 *Duoxa*-encoded subunit (14). Duox is expressed by gastric epithelial cells and  
183 generates extracellular H<sub>2</sub>O<sub>2</sub> that may serve to limit physical interactions between  
184 microbes and the epithelial surface (16). Duox has been implicated in limiting the  
185 colonization of a related *Helicobacter* species in the stomachs of mice (14).

186 To examine whether Duox impacted *H. pylori* colonization, *Duoxa*<sup>-/-</sup> mice were  
187 infected as done with WT mice for two weeks, at which point the mice were sacrificed  
188 and colonization of WT, *cheY*, and *tlpD* GFP+ *H. pylori* was compared. All *H. pylori*  
189 strains colonized the *Duoxa*<sup>-/-</sup> mutants to levels that were a bit elevated but not  
190 significantly different from those in WT mouse hosts (Fig. 3A). Gland loads were also  
191 generally similar between WT and *Duoxa*<sup>-/-</sup> glands, across WT and *tlpD* mutant *H.*  
192 *pylori* in both locations, and *cheY* mutants in the corpus (Fig. 3B). There was a  
193 modest increase in gland load in the antrum of the *tlpD* mutant and a very large  
194 decrease in loads of the *cheY* mutant, suggesting the effect of Duox was greatest in the  
195 antrum.

196 We next assessed how the loss of *Duoxa*<sup>-/-</sup> would alter gland occupancy. WT *H.*  
197 *pylori* gland occupancy was seemingly unaffected by the loss of *Duoxa*<sup>-/-</sup>, as ~50% of

198 glands were infected in this background as well as in WT mice (Fig. 3C). Interestingly,  
199 the *tlpD* mutant showed an increase in gland occupancy compared to its levels in a  
200 WT mouse, moving from <15% occupied to over 40% (Fig. 3C). Indeed, the *tlpD*  
201 mutant achieved gland occupancy levels in the corpus and antrum that were not  
202 different from WT *H. pylori* (Fig. 3C). In contrast, the *cheY* mutant was not rescued,  
203 suggesting the loss of *Duoxa*<sup>-/-</sup> rescue is specific to signals sensed by TlpD and  
204 requires chemotaxis. This apparent rescue in *tlpD* gland occupancy suggests that the  
205 loss of H<sub>2</sub>O<sub>2</sub> production by gastric epithelial cells allows for *tlpD* mutants to move  
206 more readily into new gastric glands in both the corpus and the antrum.

207

208 **Gland colonization defects of *tlpD* are rescued in hosts deficient in O<sub>2</sub><sup>-</sup>**  
209 **production by phagocytes.**

210 We next assessed the contribution of phagocyte ROS production to *H. pylori*  
211 gland colonization. Phagocyte ROS production was assessed in *Cybb*<sup>-/-</sup> mice that lack  
212 the catalytic subunit of phagocyte oxidase (Phox). *Cybb*<sup>-/-</sup> mice were infected as above  
213 and the same colonization parameters were compared between WT, *cheY*, and *tlpD*  
214 GFP+ *H. pylori*.

215 The overall colonization of the corpus and antrum was seemingly unaffected  
216 by loss of *Cybb* for all three *H. pylori* strains, showing no significant differences from  
217 WT mouse infections (Fig. 4A). Gland loads, on the other hand, were affected in *Cybb*<sup>-/-</sup>  
218 hosts. Both WT and *tlpD* *H. pylori* showed elevated gland loads in the corpus and the  
219 antrum relative to WT BL6 infections, achieving 20-30 bacteria/gland in both regions.  
220 *cheY* mutants did not follow this trend in *Cybb*<sup>-/-</sup> hosts and instead showed reduced

221 gland loads in the corpus and the antrum relative to WT BL6 infections (Fig 4B). This  
222 outcome suggests that superoxide may limit *H. pylori* numbers in a chemotaxis-  
223 dependent way. Lastly, we compared gland occupancy in *Cybb*<sup>-/-</sup> hosts. Strikingly, the  
224 *tlpD* mutant gland occupancy in both the corpus and antrum climbed to levels that  
225 were not different from WT *H. pylori*. This finding suggests that, similarly to *Duoxa*<sup>-/-</sup>  
226 infections, gland occupancy defects of *tlpD* were rescued by loss of host ROS. As seen  
227 with *Duoxa*<sup>-/-</sup> infections, *cheY* gland occupancy did not appear to benefit from the loss  
228 of *Cybb*<sup>-/-</sup> (Fig 4C). These results suggest that the loss of superoxide production by  
229 phagocytes rescues gland colonization defects of *tlpD H. pylori*, as was observed in  
230 *Duoxa*<sup>-/-</sup> hosts. Chemotaxis appears necessary for this rescue, as *cheY* mutants showed  
231 similar gland colonization values observed in WT mice. Taken together these results  
232 suggest that host-generated ROS serves as a barrier for gland colonization by *H. pylori*  
233 that the bacteria overcome via TlpD-mediated chemotactic responses. Furthermore,  
234 *tlpD* colonization defects can be attributed to low gland occupancy in the corpus and  
235 the antrum, which can be rescued to WT levels by disrupting host ROS production.

236

## 237 **Discussion**

238 We report here the *H. pylori* requires repellent ROS chemotaxis to be able to  
239 successfully colonize glands. Factors that control gland colonization throughout the  
240 GI tract are poorly understood, although it appears that an interplay exists between  
241 host and microbe to regulate gland access. Host factors known to limit gland  
242 colonization include mucus production (1), oxygen gradients emanating from the  
243 epithelial surface (19), and antimicrobial peptide production (2). Microbial

244 adaptations that have been reported to aid gland colonization include chemotaxis (5-  
245 7), sugar transport systems (20), and the ability to dampen host immune responses  
246 (21). Therefore it seems reasonable to posit that glands represent a desired but  
247 protected niche for some microbes in the GI tract. Our work demonstrates that ROS  
248 limits gland access, and chemotaxis helps overcome this barrier.

249 Host ROS generation has been implicated in limiting microbial adhesion in the  
250 intestine, and *Duoxa*<sup>-/-</sup> mice showed elevated mucosal penetrance by a subset of the  
251 microbiota (16). Our data suggest that host ROS plays an important role in restricting  
252 gland access in the stomach, and that bacteria can use chemotaxis to overcome this  
253 barrier. Gland colonization defects observed for *tlpD* mutants in WT hosts were  
254 effectively rescued in hosts with ROS production defects. Low gland occupancy could  
255 be due to very low initial colonization, or to low gland-gland spread, as both of these  
256 processes benefit from chemotaxis (5). Our results lend support to a prior report  
257 suggesting that TlpD mediates chemotactic repellent responses to ROS treatments *in*  
258 *vitro* (12), and defines the nature of colonization defects of *tlpD* mutants which have  
259 been described in the past (9, 10).

260 Our work additionally suggests that chemotaxis is not required for growth  
261 once bacteria are in glands, because we observed here that non-chemotactic and *tlpD*  
262 mutants obtained high numbers/gland in WT mice. These results are match those in a  
263 prior report, which presented the average number of bacteria/gland in a manner that  
264 included uninfected glands in that calculation (5). Excluding uninfected glands from  
265 data reported in Keilberg *et al.* concerning gland loads for WT and *cheY* *H. pylori*  
266 would produce similar values as those described in this report (5). It is not yet

267 known what sets gland load, but it has been observed that this number varies over  
268 the course of a mouse infection, climbing to an average of ~15-25 bacteria/gland  
269 within the first month, and then dropping to less than 5 by six months of infection (5).  
270 Our results show that chemotaxis can affect these within-gland levels, somewhat  
271 surprisingly playing a role to limit bacterial numbers. Our data suggest the possibility  
272 that chemotaxis plays a critical role in gland exit, such that without chemotaxis or  
273 TlpD specifically, bacterial numbers rise in the glands but bacteria cannot effectively  
274 leave. This phenotype in turn creates poor gland occupancy.

275         Previous work showed that TlpD drives chemotactic repellent responses *in*  
276 *vitro* (7, 11, 12), and our data is consistent with the idea that it also mediates  
277 chemotactic repellent responses in the host. Specifically, we found that *tlpD* mutants  
278 display hallmarks of high ROS exposure, in agreement with the idea that these  
279 mutants cannot avoid ROS. One role for this response *in vivo* comes from the  
280 observation that hosts upregulate defensive ROS production upon *H. pylori* infection  
281 (15). Thus, *H. pylori* may experience a delay from initially colonizing glands to  
282 experiencing stress imparted by the host. Therefore a repellent response mediated by  
283 TlpD could limit the detrimental effect of these stresses.

284         In conclusion, we have described host ROS generation as an additional host  
285 limitation on gland colonization in the stomach that is overcome by chemotaxis. We  
286 implicate the *H. pylori* cytoplasmic chemoreceptor TlpD in ROS-dependent gland  
287 colonization effects in the host and show that colonization defects noted for a *tlpD*  
288 mutant in WT hosts is relieved in ROS-production deficient hosts. TlpD appears to be  
289 involved in the dispersal of *H. pylori* between glands in a ROS-dependent fashion.

290

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297 design, data collection and interpretation, or the decision to submit the work for  
298 publication.

299

300 **Methods:**

301 **Bacterial strains and culture conditions**

302 WT and *cheY* SS1 GFP+ *H. pylori* described previously were employed for mouse  
303 infections (5). *H. pylori* was cultured on either Columbia horse blood agar (CHBA), or  
304 brucella broth with 10% fetal bovine serum (FBS; Life Technologies) (BB10). CHBA  
305 consisted of Columbia agar (BD) with 5% defibrinated horse blood (Hemostat Labs,  
306 Davis, CA), 50 µg/ml cycloheximide, 10 µg/ml vancomycin, 5 µg/ml cefsulodin, 2.5  
307 U/ml polymyxin B, and 0.2% (wt/vol) β-cyclodextrin. All chemicals were from  
308 Thermo Fisher or Gold Biotech. Cultures were incubated at 37°C under 5 to 7% O<sub>2</sub>,  
309 10% CO<sub>2</sub>, and balance N<sub>2</sub>.

310

311 **Creation of *tlpD* GFP+ *H. pylori* mutants**

312 *tlpD* GFP+ *H. pylori* strain SS1 (KO1614) was constructed by transformation of  
313 Δ*tlpD::cat* SS1 (KO914) (12, 22) with the plasmid pTM115 (5, 23) isolated from *H.*  
314 *pylori* strain SS1, and selected on CHBA plates containing 15 µg/ml kanamycin (5, 9).

315

316 **Animal infections and *H. pylori* colonization calculations**

317 The University of California, Santa Cruz Institutional Animal Care and Use Committee  
318 approved all animal protocols and experiments. *Cybb*<sup>-/-</sup> targeted homozygous null  
319 mice in a B6.129S background were obtained from Jackson Laboratory (JAX stock  
320 #002365, Bar Harbor, ME)(24); *Duoxa*<sup>-/-</sup> mice lacking functional dual oxidase  
321 enzymes by virtue of loss of the *duoxa1*-*duoxa2* maturation subunits (25) were  
322 obtained as heterozygotes on the B6 background from the University of Michigan. All

323 mice were obtained as breeding pairs, and bred at UC Santa Cruz. *Duoxa*<sup>-/-</sup> mice were  
324 generated, screened, and maintained as previously described (14). In brief, *Duoxa*  
325 genotyping was performed by isolating genomic DNA from tail tissue with the Qiagen  
326 DNeasy Blood & Tissue Kit., followed by PCR with a common primer (DA-WT/KO), a  
327 WT allele-specific primer (DA- WT-R), and a knockout allele-specific primer (DA-KO-  
328 R) (14). Genotypes were judged by the presence of the WT allele as a 381-basepair  
329 fragment, and the knockout allele as a size of 568 basepair fragment (25).

330 Six to eight-week-old mice (male and female) were infected intraorally by  
331 allowing the animals to drink a 50 microliter suspension from a pipette tip containing  
332 *H. pylori* grown to mid-exponential phase and concentrated to an optical density at  
333 600 nm of 3.0 (~ 5x10<sup>7</sup>/50 µl) in BB10 medium, as done previously (6). At the end of  
334 an infection period, mice were sacrificed by CO<sub>2</sub> narcosis. The stomach was removed,  
335 opened along the lesser curvature and washed in phosphate-buffered saline (PBS) to  
336 remove food. The corpus and antrum were divided based on tissue coloration, cut  
337 into pieces that were then processed to analyze total bacterial colonization, gland  
338 isolation, or for RNA extraction. For total bacterial colonization, corpus and antral  
339 tissue was weighed, homogenized with the Bullet Blender (Next Advance) with 1.0-  
340 mm zirconium silicate beads, and then plated to determine the number of colony  
341 forming units (CFU) per gram of stomach tissue on CHBA with the addition of 20  
342 µg/ml bacitracin, 10 µg/ml nalidixic acid, and 15 µg/ml kanamycin.

343



344 **Gland isolation and microscopy**

345 Glands were isolated by incubating dissected gastric tissue in Dulbecco's phosphate-  
346 buffered saline (DPBS) (Millipore) plus 5 mM EDTA at 4°C for 2 hours with agitation,  
347 as described previously (5, 26). The tissue was subsequently transferred to DPBS  
348 containing 1% sucrose and 1.5% sorbitol and shaken for thirty seconds. Glands were  
349 labeled with 10 µg/ml Hoechst DNA stain (Life Technologies). Glands were kept on  
350 ice and examined as soon as possible. Ten microliters of shaken tissue were placed on  
351 glass slides and visualized with a Nikon Eclipse E600 microscope with fluorescence  
352 filters for 4',6'-diamidino-2- phenylindole (DAPI), GFP, and RFP. For each time point  
353 and infection, 100 glands each were imaged for the corpus and antrum, and the  
354 number of *H. pylori* cells inside the gland was counted manually for each gland. Gland  
355 load levels were calculated by averaging the number of bacteria observed in  
356 colonized glands per mouse and *H. pylori* strain. Gland occupancy was calculated as  
357 the frequency of glands occupied per mouse host and averaged over at least three  
358 mice. Gland colonization comparisons were made for at least three mice per genotype  
359 and *H. pylori* strain.

360

361 **RNA isolation and qPCR**

362 Gastric tissue was flash frozen in liquid nitrogen, homogenized in TRIzol (Invitrogen)  
363 and RNA was isolated following the TRIzol RNA isolation protocol (GIBCO). DNA was  
364 removed by following the TURBO DNA-free kit protocol (Life technologies). cDNA  
365 was produced with the High-Capacity cDNA Reverse Transcription Kit (Life  
366 technologies) using random primers. qPCR was performed using the SensiFAST SYBR

367 No-ROX kit (Bioline) using the primers listed below. Primer efficiency was calculated  
368 by amplifying serial dilutions of WT *H. pylori* genomic DNA, plotting the Ct values  
369 obtained per dilution and calculating the slope. Efficiencies were derived from the  
370 slope with the equation  $\text{Efficiency} = -1 + 10^{(-1/\text{slope})}$  (27). Relative fold changes were  
371 calculated using the  $\Delta\Delta\text{Ct}$  method with Pfaffl correction for PCR amplification  
372 efficiency, using *16S* and *gapB* as reference genes with primers listed 5'-3' below (27).  
373 16S forward: GGAGGATGAAGGTTTTAGGATTG; 16S reverse:  
374 TCGTTTAGGGCGTGGACT; *katA* forward: AGAGGTTTTGCGATGAAGT; *katA* reverse:  
375 CGTTTTTGAGTGTGGATGAA; *gapB* forward: GCCTCTTGACGACTAACGC; *gapB*  
376 reverse: CTTTGCTCACGCCGGTGCTT.

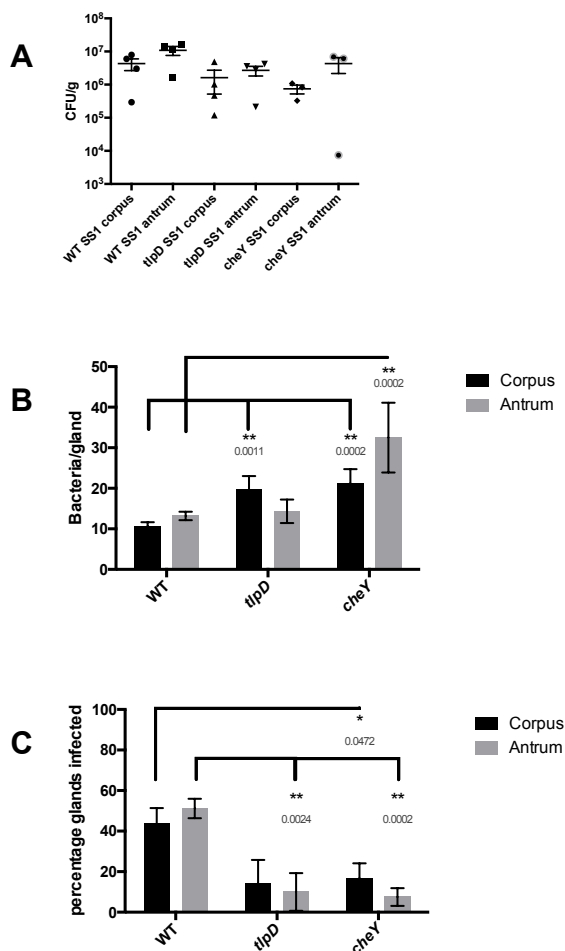
377

### 378 ***In vitro* treatment of *H. pylori* with H<sub>2</sub>O<sub>2</sub>**

379 Overnight cultures of *H. pylori* strains were adjusted to OD<sub>600</sub> = 0.2, split into two  
380 cultures with one receiving treatment with 1 mM H<sub>2</sub>O<sub>2</sub> for twenty minutes. RNA  
381 isolation and qPCR protocols were identical to that described above.

382

383 **Figures**



384

385 **Figure 1. *tlpD* mutants have deficits in gland occupancy in WT mice but not**  
 386 **colonization of total tissue or individual glands**

387 Comparison of colonization of WT mice by *H. pylori* GFP+ SS1 WT, *tlpD*, and *cheY*.

388 Mice were orally infected, and stomachs were collected and analyzed for tissue and

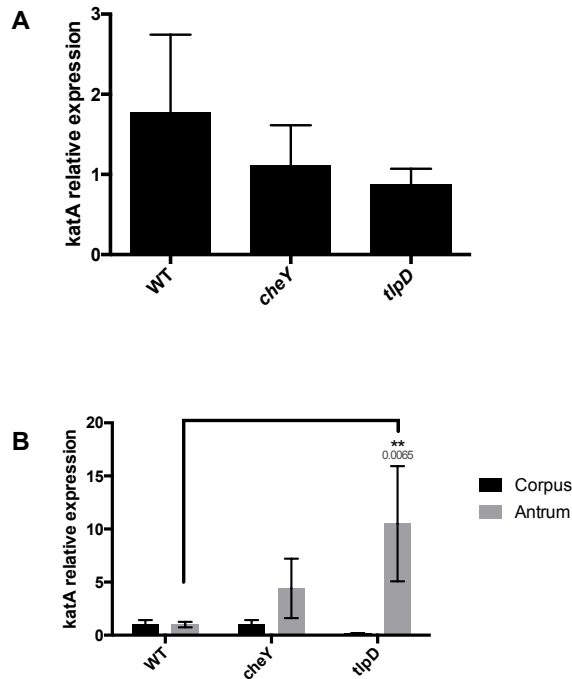
389 gland colonization after 2 weeks of infection. (A) CFU/gram for corpus or antrum

390 regions. *H. pylori* GFP+ SS1 WT (n = 4), *tlpD* (n = 4) and *cheY* (n = 3).

391 in the isolated corpus and antral glands. These numbers are the average number of

392 bacteria counted per gland, excluding uninfected glands. Infected gland numbers are:

393 WT corpus (436 glands from 6 mice), WT antrum (508 from 6 mice), *tlpD* corpus (67  
394 glands from five mice), *tlpD* antrum (48 glands from five mice), *cheY* corpus (58  
395 glands from four mice), *cheY* antrum (24 glands from four mice). (C) Gland occupancy  
396 in the isolated corpus and antral glands, representing the percentage of glands  
397 infected with the indicated *H. pylori* strain. Error bars represent standard error of the  
398 mean (SEM) for all panels. Numbers of mice infected are the same as described for  
399 gland loads. Statistical differences are indicated by \* ( $P < 0.05$ ) and \*\* ( $P < 0.01$ ) as  
400 analyzed by Student T-test.  
401



402

403 **Figure 2. *tlpD* mutants show evidence of ROS exposure *in vivo***

404 Comparison of catalase mRNA expression *in vitro* and *in vivo* between *H. pylori* strains.

405 (A) Mean +/- SEM of fold change increases in *katA* mRNA of *H. pylori* strains exposed

406 to 1 mM H<sub>2</sub>O<sub>2</sub> for twenty minutes, normalized to *gapB*. (B) Comparison of mean +/-

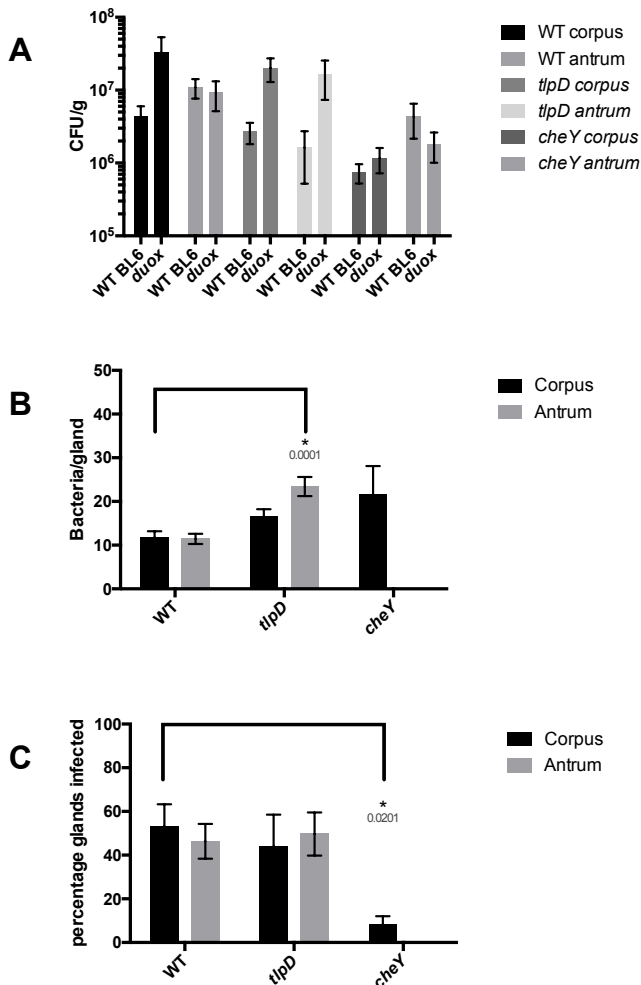
407 SEM of *katA* expression by *H. pylori* strains in three WT mice, normalized to *gapB*.

408 Statistical differences are indicated by \* ( $P < 0.05$ ) and \*\* ( $P < 0.01$ ) as analyzed by

409 Student T-test, with actual p values indicated above the bar. *gapB* expression was

410 insensitive to H<sub>2</sub>O<sub>2</sub> exposure based on comparison to 16S rRNA.

411



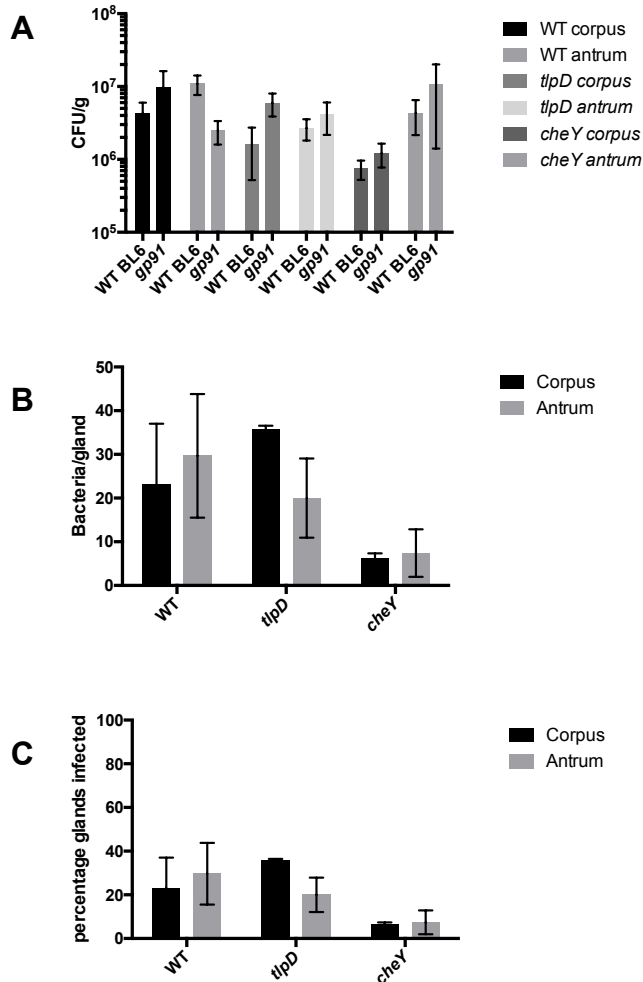
412

413 **Figure 3. Loss of epithelial H<sub>2</sub>O<sub>2</sub> rescues *tlpD* mutant gland defects**

414 Colonization of *Duoxa*<sup>-/-</sup> mice by WT, *tlpD*, and *cheY* GFP+ *H. pylori* SS1 strains at two  
 415 weeks post-infection. Mice were orally infected, and stomachs were collected and  
 416 analyzed for tissue and gland colonization. (A) CFU/gram at two weeks post-infection  
 417 for corpus or antrum regions with WT (n = 4), *tlpD* (n = 5) and *cheY* (n = 5) GFP+ *H.*  
 418 *pylori* SS1. Data for WT mice are the same as in Fig. 1, and are reshown here for  
 419 comparison. (B) Gland loads in the isolated corpus and antral glands, representing the  
 420 average number of bacteria counted per gland, excluding uninfected glands. Infected  
 421 gland numbers are: WT corpus (313 glands from six mice), WT antrum (472 glands

422 from 6 mice), *tlpD* corpus (132 from six mice), *tlpD* antrum (149 glands from three  
423 mice), *cheY* corpus (24 glands from three mice). (C) Gland occupancy in the isolated  
424 corpus and antral glands, representing the percentage of glands infected with the  
425 indicated *H. pylori* strain. Error bars represent SEM for all panels. Numbers of mice  
426 infected are the same as described for gland loads. Statistical differences are indicated  
427 by \* ( $P < 0.05$ ) and \*\* ( $P < 0.01$ ) as analyzed by Student T-test.

428



429

430 **Figure 4. Loss of immune superoxide rescues *tlpD* mutant gland defects**

431 Colonization of *Cybb*<sup>-/-</sup> mice by WT, *tlpD*, and *cheY* GFP+ *H. pylori* SS1 strains at two  
 432 weeks post-infection. Mice were orally infected, and stomachs were collected and  
 433 analyzed for tissue and gland colonization. (A) CFU/gram at two weeks post-infection  
 434 for corpus or antrum regions using WT (n = 6), *tlpD* (n = 14) and *cheY* (n = 6) GFP+ *H.*  
 435 *pylori* SS1 strains. Data for WT mice are the same as in Fig. 1, and are reshown here  
 436 for comparison. (B) Gland loads in the isolated corpus and antral glands, representing  
 437 the average number of bacteria counted per gland, excluding uninfected glands.  
 438 Infected gland numbers are: WT corpus (69 glands from three mice), WT antrum (89



439 glands from 3 mice), *tlpD* corpus (107 glands from three mice), *tlpD* antrum (60  
440 glands from three mice), *cheY* corpus (31 glands from three mice), *cheY* antrum (37  
441 glands from three mice). (C) Gland occupancy in the isolated corpus and antral glands,  
442 representing the percentage of glands infected with the indicated *H. pylori* strain.  
443 Error bars represent SEM for all panels. Numbers of mice infected are the same as  
444 described for gland loads. Statistical differences are indicated by \* ( $P < 0.05$ ) and \*\* ( $P$   
445  $< 0.01$ ) as analyzed by Student T-test.  
446

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