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Malaria parasite infection compromises colonization resistance to an enteric pathogen by reducing gastric acidity

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Malaria parasite infection weakens colonization resistance against Salmonella enterica serovar (S.) Typhimurium. S. Typhimurium is a member of the Enterobacteriales, a taxon that increases in abundance when the colonic microbiota is disrupted or when the colonic mucosa is inflamed. However, here, we show that infection of mice with Plasmodium yoelii enhances S. Typhimurium colonization by weakening host control in the upper GI tract. P. yoelii–infected mice had elevated gastric pH. Stimulation of gastric acid secretion during P. yoelii infection restored stomach acidity and colonization resistance, demonstrating that parasite-induced hypochlorhydria increases gastric survival of S. Typhimurium. Furthermore, blockade of P. yoelii–induced TNF-α signaling was sufficient to prevent elevation of gastric pH and enhance S. Typhimurium colonization during concurrent infection. Collectively, these data suggest that abundance in the fecal microbiota of facultative anaerobes, such as S. Typhimurium, can be increased by suppressing antibacterial defenses in the upper GI tract, such as gastric acid.

INTRODUCTION

Facultative anaerobic bacteria of the order Enterobacteriales (1) are minority species commonly present in the fecal microbiota of healthy adults (2). An expansion of this taxon in the fecal microbiota is a signature of dysbiosis (3), which is associated with a disruption of the colonic microbiota by antibiotics (4) or linked to inflammation of the colon in patients with ulcerative colitis (5) or colorectal cancer (6). Pathogenic members of the Enterobacteriales, such as Salmonella enterica serovar (S.) Typhimurium, Citrobacter rodentium, or Yersinia enterocolitica, use their virulence factors to trigger colitis, thereby altering the intestinal environment to increase the availability of respiratory electron acceptors, which fuel pathogen growth to escalate fecal shedding (7–11). Infection of mice with the parasite Toxoplasma gondii triggers intestinal inflammation and bacterial dysbiosis characterized by an expansion of commensal Enterobacteriales in the fecal microbiota (12, 13). T. gondii infection induces a dysbiosis dominated by Enterobacteriales by triggering an influx of macrophages, which produce nitric oxide that is converted in the gut lumen into nitrate, thereby fueling growth of commensal Enterobacteriales through nitrate respiration (14).

We recently reported that infection with a malaria parasite, Plasmodium yoelii, increases the abundance of S. Typhimurium in the fecal microbiota of mice (15), which might contribute to the increased risk of malaria patients to develop invasive bloodstream infections with nontyphoidal Salmonella (NTS) serotypes, such as serovar (S.) Typhimurium (16, 17). Infection of mice with P. yoelii triggers expression of inflammatory cytokines and inflammatory infiltrates of macrophages and T cells in the cecal mucosa (15), but it remains unknown whether parasite-induced intestinal inflammation is responsible for increasing fecal shedding of S. Typhimurium during coinfection. Here, we used a coinfection model to interrogate potential interactions by which malaria increases susceptibility to intestinal colonization by Salmonella.

RESULTS

P. yoelii infection increases intestinal colonization of S. Typhimurium

In experimental malaria models such as P. yoelii nigeriensis (henceforth referred to as P. yoelii), infection of mice can alter the gut environment (15, 18) and reduce colonization resistance against S. Typhimurium (15, 19). However, at later time points after infection, S. Typhimurium can overcome colonization resistance because its virulence factors, two type III secretion systems (T3SS-1 and T3SS-2), trigger intestinal inflammation (9). To disentangle the contribution of concurrent malaria from the contribution of S. Typhimurium virulence factors to weakening colonization resistance, groups of mice were infected with the S. Typhimurium wild type or an invA spiB mutant that lacks T3SS-1 (due to a mutation in invA) and T3SS-2 (due to a mutation in spiB) at various time points following P. yoelii inoculation (outlined in Fig. 1A). When challenged at 6 days post–P. yoelii infection (dpp) or later, a S. Typhimurium invA spiB colonized the ceca and colons of P. yoelii–infected mice at 10- to 100-fold higher levels than in control mice (Fig. 1, B and C, and fig. S1A). This was independent of S. Typhimurium virulence, as both wild-type and the invA spiB mutant displayed enhanced colonization (fig. S1B). The 6-dpp time point correlated with a peak in parasite expansion in the blood (fig. S2A). By this time, mice infected with P. yoelii had consistently developed severe anemia (fig. S2B) and showed substantial weight loss (fig. S2C), while splenomegaly increased steadily throughout the parasite infection (fig. S2D). Therefore, 6 days following P. yoelii inoculation was determined to be the earliest reliable time point to interrogate underlying causes of the malaria-associated
defect in colonization resistance and was used as the primary model in subsequent experiments.

**Impact of nitrate- and oxygen-associated growth on boosting S. Typhimurium colonization**

Because macrophages recruited during *T. gondii* parasite infection fuel growth of commensal Enterobacterales through nitrate respiration (14) and previous work shows that *P. yoelii* infection also recruits macrophages to the cecal mucosa (15), we hypothesized that concurrent malaria might increase the luminal availability of nitrate to fuel a S. Typhimurium expansion. To test this idea, we compared the fitness of the S. Typhimurium wild type with a mutant lacking nitrate reductase activity (*napA narG narZ cyxA* mutant) by infecting mice with a 1:1 mixture of both strains. Genetic ablation of nitrate respiration did not reduce the fitness of S. Typhimurium in *P. yoelii*–infected mice (Fig. 1, D and E), despite increased intestinal burden overall in the coinfected group. This finding indicated that nitrate utilization was dispensable for increased fecal recovery of S. Typhimurium during concurrent malaria.

Alternatively, it has been shown that inflammation also benefits S. Typhimurium via increasing the bioavailability of oxygen. Depletion of *Clostridia*, either through S. Typhimurium virulence-associated inflammation or antibiotic administration, results in decreased microbiota production of the short-chain fatty acid (SCFA) butyrate, which typically fuels epithelial colonocytes responsible for maintaining luminal hypoxia (9). Reduced butyrate availability switches colonocyte primary metabolism away from oxygen-intensive breakdown of SCFAs, allowing increased diffusion of oxygen into the colonic lumen. This provides an additional respiratory electron acceptor for growth of facultative anaerobes such as *Salmonella* and *E. coli* using cytochrome oxidases (9, 20, 21). As our previous work found that *P. yoelii* infection can affect the microbiota, including a relative reduction in *Clostridia* in the feces (15), we hypothesized this could be opening the oxygen respiratory niche in the colon and promoting *Salmonella* growth. To test this idea, we compared the fitness of the S. Typhimurium wild type with a mutant lacking the high-affinity cytochrome *bd* oxidase (*cydA* mutant). However, although *cydA* provided an approximately 10-fold fitness advantage in *P. yoelii*–infected mice, a similar fitness advantage was observed in mice not exposed to *P. yoelii* (Fig. 1, F and G), suggesting that increased oxygenation of the intestinal lumen in coinfect ed mice did not contribute to the heightened S. Typhimurium levels. Overall, these results

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**Fig. 1. Enhanced colonization by S. Typhimurium in *P. yoelii*–infected mice is independent of nitrate and oxygen respiratory growth by *Salmonella*.** (A) Schematic outlining the coinfection model used in (B) and (C). *P. yoelii*– and mock-infected mice were challenged at 2, 4, 6, or 8 days after infection (as indicated) with 10⁷ CFU *Salmonella* Typhimurium *invA spiB KanR* (peroral). i.p., intraperitoneal. *Salmonella* loads in the colon (B) and cecum (C) were determined 24 hours after *Salmonella* infection. Data are pooled from two separate experiments with three to six mice per time point (n = 7 to 10 total per group). (D to G) Mock- and *P. yoelii*–inoculated mice (n = 5 per group) were gavaged at 6 dpp with approximately 5 × 10⁸ CFU wild-type S. Typhimurium (WT) and 5 × 10⁸ CFU of either a S. Typhimurium *napA narZ narG cyxA* mutant (*3NR cyxA*, D and E) or a S. Typhimurium *cydA* mutant (F and G). (D) Fecal S. Typhimurium CFUs and (E) competitive indices (WT/*3NR cyxA*) at 24 hpi. (F) Fecal S. Typhimurium CFUs and (G) competitive indices (WT/*cydA*) at 24 hpi. Bars represent the geometric mean. Each symbol or linked pair of symbols represents data from one animal. **P ≤ 0.01, ***P ≤ 0.001, and ****P ≤ 0.0001. NS, not significant.
indicate that increased access to respiratory electron acceptors was not a major contributing factor to the defect in colonization resistance during *P. yoelii* infection.

**P. yoelii** infection does not rapidly increase intestinal *S. Typhimurium* replication

The absence of *Plasmodium*-enhanced respiratory growth defects led us to question whether reduced colonization resistance was actually coupled with more rapid growth of *S. Typhimurium* in coinfectected mice. To evaluate this, mice were inoculated with *S. Typhimurium* carrying the lactose-addicted plasmid pAM34 (22), which allows for the short-term assessment of bacterial growth in low-lactose environments such as the adult mouse intestine (23). At 24 hours after challenge, *S. Typhimurium* populations throughout the gut exhibited equivalent replication rates in both *P. yoelii* coinfectected and mock-infected mice, despite increased bacterial burdens in the coinfectected group (Fig. 2, A and B). Intriguingly, we found that *S. Typhimurium* colonization was already greater in coinfectected mice by 4 hours after challenge, also independent of the in vivo growth rate (Fig. 2, C and D). This rapid difference in *S. Typhimurium* loads without a concomitant increase in replication indicated that boosted growth of the bacteria is not responsible for the enhanced colonization of *P. yoelii*-infected mice.

**P. yoelii** infection affects gastric antibacterial defense

Because differences in *S. Typhimurium* loads arose rapidly after bacterial inoculation of mice, we hypothesized that *P. yoelii* infection may compromise intrinsic “bottleneck” defenses against bacterial colonization that act early following pathogen ingestion. As we also observed that coinfectected mice often exhibited higher *S. Typhimurium* levels throughout the upper gastrointestinal (GI) tract (fig. S3A), it seemed likely that loss of colonization resistance against *S. Typhimurium* in coinfectected mice could be occurring in the stomach or small intestine.

One of the most severe initial bottlenecks against colonization by many enteric pathogens is imposed by the highly acidic (pH 1.5 to 3) environment encountered in the stomach (24). While *S. Typhimurium* thrives at neutral pH (25), it is capable of surviving short-term (60 min) exposure to acidity as low as pH 4 in vitro (Fig. 3A). However, inoculated bacteria are rapidly killed at pH 3.5 or lower, with no bacteria recovered from exposure to media at pH 2.5 or below (Fig. 3A). A similar degree of acid sensitivity is shared by many Enterobacteriales, including both pathogenic and nonpathogenic varieties (fig. S4, A to E). This finding corroborates previous work indicating that *S. Typhimurium* and other enteric pathogens are highly susceptible to killing by the acid levels present in the stomach (24). Thus, deficiencies in maintaining gastric acidity could result in greater survival of ingested *S. Typhimurium* into the intestines.

When measuring the acidity of the stomach lumen directly using a pH-sensing microelectrode, *P. yoelii* infection was associated with a significantly higher gastric pH (mean pH = 4.02 ± 1.07) compared to mock-infected mice (mean pH = 2.46 ± 0.32) by 6 dpp (Fig. 3B). Onset of hypochlorhydria during *P. yoelii* infection correlated with the loss of colonization resistance, as it was not detectable in mice earlier than 6 dpp (fig. S3B). Stomachs of mice at 6 dpp were generally smaller than mock-infected controls (fig. S5A), likely resulting in part from reduced food consumption (fig. S5B) that paralleled observed weight loss patterns during parasite infection (fig. S2C). As meal intake can be stimulatory for gastric acid secretion (26), we predicted that the observed pH differences were a result of acute reductions in food intake during malaria. Fasting mock-infected mice overnight was sufficient to reduce stomach weight (fig. S5C) but did not alter the gastric pH of the fasted mice compared to fed mice with ad libitum food access (fig. S5D). Fasting also did not normalize colonization resistance against inoculated *S. Typhimurium* between fasted mock- and *P. yoelii*-infected mice (fig. S5E), suggesting that more than acute decreases in food intake during *P. yoelii* infection is necessary to explain the defects in gastric acidity and susceptibility to *S. Typhimurium* colonization.

**Malaria-induced reduction in gastric acidity affects *S. Typhimurium* colonization**

We next wanted to determine whether the observed hypochlorhydria is actually responsible for the increased survival of coinfectected *S. Typhimurium*. Gastric acid secretion is normally regulated through a combination of stimulatory and inhibitory signaling to gastric parietal cells, leading to translocation of the gastric proton pump to the apical membrane and active acid secretion into gastric lumen (26). One of the most potent prosecretion signals is histamine, released locally by enterochromaffin-like cells in the gastric glands, binding parietal cell H2 receptors (H2R) (26). To assess whether the high gastric pH during *P. yoelii* infection was responsible for the defect in colonization resistance, mice were treated with the specific H2R agonist dimaprit dihydrochloride to stimulate gastric acid secretion 1 hour before *S. Typhimurium* challenge. H2R agonist administration acutely reduced the stomach pH in *P. yoelii*-infected animals to mock-infected levels (Fig. 3C). Furthermore, H2R stimulation...
in *P. yoelii*-infected mice rescued colonization resistance against *S. Typhimurium* (Fig. 3, D and E) without affecting other aspects of the parasite infection, including parasitemia (fig. S6A) and anemia (fig. S6B).

In complementary experiments, mice were treated with the proton pump inhibitor omeprazole to block acid secretion into the stomach lumen before *S. Typhimurium* challenge. Omeprazole treatment in control mice resulted in a rise in both the gastric pH (Fig. 3F) and *S. Typhimurium* burden (Fig. 3, G and H) relative to vehicle-treated mice, as has been previously observed (24). However, omeprazole administration in *P. yoelii*-infected animals did not further increase *S. Typhimurium* loads over *P. yoelii*-infected mice treated with Dimaprit or vehicle on 6 dpp, then challenged with *S. Typhimurium invA spiB Kan* (peroral) for 4 hours. (F to H) The stomach pH (F), cecal (G), and colon (H) *S. Typhimurium* loads of mock- and *P. yoelii*-infected mice treated with Omeprazole or vehicle on 3, 4, and 5 dpp and challenged on 6 dpp with *S. Typhimurium invA spiB Kan* (peroral) for 24 hours. (C to H) Combined data from separate experiments with *n* = 6 to 8 mice per group. Bars represent the geometric mean, and symbols represent data from individual mice. *P* ≤ 0.05, **P** ≤ 0.01, ***P** ≤ 0.001, and ****P** ≤ 0.0001.

**Fig. 3.** *P. yoelii*-infected mice develop hypochlorhydria that affects colonization of *S. Typhimurium.* (A) *S. Typhimurium* survival in PBS at various pHs. Circles represent individual replicates while the curve is the nonlinear regression (4PL). Horizontal lines indicate 100 and 50 percent survival, while the vertical line marks the pH associated with 50 percent survival. (B) Gastric pH of mice 6 days after *P. yoelii* or mock infection. (C to E) The stomach pH (C), cecal (D) and colon (E) *S. Typhimurium* loads of *P. yoelii*– and mock-infected mice treated with Dimaprit or vehicle on 6 dpp, then challenged with *S. Typhimurium invA spiB Kan* (peroral) for 4 hours. (F to H) The stomach pH (F), cecal (G) and colon (H) *S. Typhimurium* loads of mock- and *P. yoelii*-infected mice treated with Omeprazole or vehicle on 3, 4, and 5 dpp and challenged on 6 dpp with *S. Typhimurium invA spiB Kan* (peroral) for 24 hours. (C to H) Combined data from separate experiments with *n* = 6 to 8 mice per group. Bars represent the geometric mean, and symbols represent data from individual mice. *P* ≤ 0.05, **P** ≤ 0.01, ***P** ≤ 0.001, and ****P** ≤ 0.0001.

Altered expression of gastric signaling peptides during malaria

To explore potential mechanisms underlying the rise in gastric pH, we analyzed the effect of *P. yoelii* on the expression of genes associated with regulation of acid secretion by parietal cells in the stomach (26). Notably, in the stomach tissues of *P. yoelii*-infected mice, expression of the parietal-stimulatory hormone gastrin (*Gast*) was reduced two- to fourfold (Fig. 4A), while transcripts for the inhibitory hormone somatostatin (*Sst*) were increased nearly twofold relative to mock (Fig. 4B). This result suggested that alterations in typical signaling mechanisms governing acid secretion could be contributing to hypochlorhydria.

Parietal cells use specialized H⁺- and K⁺-dependent adenosine triphosphatases (H⁺,K⁺-ATPases; ATP4) concentrated intracellularly on tubulovesicles that localize apically when stimulated to actively secrete protons into the stomach lumen, generating the low pH (26, 27).
Intriguingly, expression of genes encoding the proton pump (Atp4a/b) was slightly reduced in the stomach during malaria, with Atp4b notably reduced twofold relative to mock mice (Fig. 4, C and D). Gastric acid secretion is not typically regulated by altering the expression of proton pump components, so we explored whether this reduction in transcript abundance instead paralleled a reduction in parietal cell abundance, the primary cells expressing Atp4a/b. Blinded histopathology of hematoxylin and eosin (H&E)–stained gastric tissues was imaged by light microscopy (fig. S7A) and fluorescence imaging (fig. S7B) to more accurately quantify highly eosinophilic cells. Eosin is autofluorescent and parietal cells stain strongly with eosin (46). Fig. 4E, suggesting that elevated pH may result from reduced mRNA abundance instead paralleled a reduction in parietal cell abundance, the primary cells expressing Atp4a/b.

Proinflammatory signaling during malaria reduces gastric acidity

We next investigated how the immune response to P. yoelii may be affecting gastric acid production. In the course of assessing gastric gene expression, we observed elevated transcripts for the inflammatory cytokine tumor necrosis factor–α (TNF-α, encoded by Tnf) in the stomach tissue of P. yoelii–infected mice (Fig. 4F). TNF-α treatment was reported to reduce secretagogue responses by rabbit parietal cells (29), and TNF-α administration induced apoptosis of rat parietal cells (30). Moreover, it is known that both clinical and experimental malaria can be associated with elevated circulating TNF-α (31–33), a response that is thought to help control parasite replication (31, 34) but can also be associated with severe or cerebral malaria (32, 33, 35). In line with these findings, circulating TNF-α was also higher with P. yoelii infection than in control mice (Fig. 5A). Thus, we hypothesized that TNF-α responses to the parasite might be interfering with gastric acid secretion.

In mice treated with TNF-α neutralizing antibody, mock- and P. yoelii–infected groups displayed equivalent gastric pH (Fig. 5B and fig. S8A), suggesting that TNF-α signaling is necessary for development of hypochlorhydria with P. yoelii. In the context of S. Typhimurium, TNF-α blockade also equalized initial [3 hours postinfection (hpi)] colonization between mock- and P. yoelii–infected mice (Fig. 5C and fig. S8B), compared to groups treated with an isotype control. Longer challenge time points were not assessed, as TNF-α responses are also necessary for appropriate host defenses to limit systemic S. Typhimurium infection (36, 37). Despite TNF-α blockade, mice infected with Plasmodium still developed anemia (Fig. 5D) and displayed even greater circulating parasitemia (Fig. 5E). This latter impact on parasite burden is actually evidence of effective antibody blockade, as early TNF-α responses to Plasmodium are associated with limiting parasite replication in the host (34, 35). TNF-α blockade did not significantly affect food consumption or weight loss in infected mice, compared with isotype-treated controls (fig. S8, C to E). P. yoelii–infected mice given anti–TNF-α antibody still showed an increase in gastric TNF-α transcripts (fig. S8F), but TNF-α blockade led to a reduction of pro–interleukin-1β expression (fig. S8G), indicating a potential dampening of the proinflammatory cytokine response. Overall, these results link proinflammatory TNF-α...
responses by the host during *P. yoelii* infection with loss of the intrinsic gastric acid defense, thereby increasing susceptibility to secondary infection by enteric pathogens.

DISCUSSION

The expansion of facultative anaerobic bacteria in the fecal microbiota has been associated with disruption of the colonic microbiota by antibiotics (4) and additionally can occur during inflammation of the colon in patients with ulcerative colitis (5) or colorectal cancer (6). While in these instances, increase in the relative abundance of facultative anaerobes such as Enterobacterales has been linked to alterations in the colonic environment, our results demonstrate that changes to stomach acid, an antimicrobial defense of the upper GI tract, can affect the composition of the fecal microbiota. Thus, our findings on how malaria affects colonization resistance to *S. Typhimurium*, a foodborne microbe, have implications for understanding factors affecting the composition of the microbiota in the lower GI tract, as assessed by fecal microbiota profiling. For example, abundance of bacteria from the oral cavity in the fecal microbiota has been linked to multiple pathogenic processes throughout the digestive tract. Increased prevalence of oral bacteria such as *Fusobacterium* has been found in the fecal microbiota of patients with systemic inflammation caused by inflammatory bowel disease (38), liver cirrhosis (39), and HIV infection (5). This spread of oral bacteria to other sites in the digestive tract is clinically important, especially because some members of the oral microbiota have been linked to colonic (40) and pancreatic cancers (41). Notably, patients with both AIDS (42) and liver cirrhosis (43) exhibit hypochlorhydria, although the underlying mechanisms are unknown. Our results suggest that in addition to lowering the barrier to enteric pathogen colonization, inflammation-induced hypochlorhydria may also contribute to spread of oral microbiota to other sites in the digestive tract. Therefore, interpretation of alterations to the fecal microbiota may need to consider habitat filters in the upper GI tract, such as the acidic barrier of the stomach.

Susceptibility to enteric pathogen infection is regulated by both host- and microbiota-associated defenses. Host physiology and immunity influence the distinct habitats of the GI tract, thereby filtering for the most suitable microbial communities throughout (21, 44). The microbiota then serves, in part, to resist pathogen colonization by training host immunity and preempting access to resource and nutrient niches necessary for potential pathogens to thrive in the gut (44–46). However, many enteric pathogens have evolved virulence systems to avert microbiota-mediated colonization resistance by engineering different growth niches to bloom in the intestine (7, 47). Such mechanisms still typically require sufficient quantities of the pathogen to reach the lower intestines to trigger a substantial host response to alter the gut environment (48, 49). Given this limitation, intrinsic host defenses that rapidly restrict survival of ingested microbes—particularly in the upper GI tract—are necessary to maintain maximal defense against infection. This type of resistance is largely provided by the innate production and secretion of compounds with antibacterial properties, such as host-derived antimicrobial peptides (45), various components of bile (23, 45), and gastric acid (24, 50). It is ultimately the combined gauntlet of antimicrobial defenses and nutrient limitation that must be avoided or averted by pathogens to infect the host and cause disease.

Our prior research showed that mice infected with *Plasmodium*, a pathogen that replicates in the bloodstream during the erythrocytic infection state, display increased susceptibility to colonization by NTS (15). The findings presented here advance our understanding of how malaria affects colonization resistance to pathogens by linking the boosted implantation of *Salmonella* to a previously unreported, prolonged reduction in gastric acidity associated with inflammatory signaling during infection with the malarial parasite.

Low stomach pH in healthy animals serves as an important bottleneck in the colonization of many bacterial pathogens (24), including *S. Typhimurium* and related bacteria (Fig. 3A and fig. S4, A to E). Drugs affecting gastric acidity in humans, such as proton pump inhibitors, have been noted as risk factors that can both alter the gut microbiome and increase susceptibility to intestinal infection with pathogens including *Salmonella, Campylobacter*, and *Clostridioides difficile* (45, 51–54). In our model, high stomach pH during malaria was associated with increased *S. Typhimurium* abundance after infection, and exogenous stimulation of gastric acid secretion before *S. Typhimurium* inoculation rapidly restored resistance to colonization, demonstrating that parasite-induced hypochlorhydria was responsible for increased susceptibility to the secondary infection.

Typically, parietal cells in the gastric glands are stimulated by a combination of histamine, gastrin, and acetylcholine released from nearby cells, leading to apical translocation of the parietal’s H⁺/K⁺-ATPase proton pump, thereby allowing active acid secretion into the stomach lumen (26). Somatostatin serves as the major hormonal inhibitor of gastric acid secretion via direct action on parietal cells and indirectly through inhibition of histamine release by enterochromaffin-like cells. Supporting the notion that *Plasmodium* infection alters normal signaling for gastric acid release, we found increased somatostatin and reduced gastrin expression in gastric tissues from parasite-infected mice (Fig. 4, A and B), which suggests reduced parietal cell stimulation as a mechanism underlying elevated gastric pH. This mechanism is consistent with our findings that parietal cell abundance was unchanged (Fig. 4E) and that stimulation with an H2R agonist resulted in acid production (Fig. 3D).

In the context of gastric *Helicobacter pylori* infection, local production of proinflammatory cytokines, including TNF-α, can influence and inhibit gastric acid secretion by parietal cells (26, 29, 55). Our results extend this concept to a parasite infection, as antibody blockade of TNF-α signaling during *Plasmodium* infection was sufficient to maintain normal gastric acidity (Fig. 5B) and colonization resistance (Fig. 5C). TNF-α is known to also be involved in both mouse and human innate immune responses to malaria (33, 56) and appears to contribute to limiting parasite replication in host cells (34), supporting our data that indicates mice treated with TNF-α blocking antibody develop higher circulating parasitemia (Fig. 5E). This inflammatory cytokine response may represent a shared mechanism between our model and susceptibility to disseminated salmonellosis associated with human malaria. It remains unclear whether the hypoacidic environment could actually be affecting the rest of the microbiota, either directly or indirectly (e.g., through altered digestion or vitamin availability), to produce the shifts in the microbial community composition observed previously (15, 19), or whether these changes are the result of additional GI impacts of *Plasmodium* infection.

In sub-Saharan Africa, malaria is associated with a higher risk for systemic NTS infections in children (16, 17, 57). While prior work in mouse models helped illuminate how elements of the immune response to *Plasmodium* can promote systemic infections by *S. Typhimurium*
(58, 59), our current findings suggest the additional mechanism of *Plasmodium*-associated hypochlorhydria by which underlying malaria increases susceptibility to colonization by *S. Typhimurium*. This generalized reduction in initial bacterial killing could particularly benefit the multidrug-resistant ST313 isolates now circulating in Africa, which exhibit genomic degradation and reduced capacity for inflammatory activation (57). Induction of inflammation is well recognized as part of the usual NTS strategy for colonizing the host (7–9, 44, 47), but avirulent *S. Typhimurium* incapable of inducing a substantial intestinal inflammatory response displayed equivalent colonization enhancement by *P. yoelii* (fig. S1B). This suggests that in the context of malaria, reduced killing of the bacteria in the stomach and upper intestines could compensate for genetic deficits in NTS pathogenesis identified in the now circulating strains. Moreover, malaria-associated suppression of colonization resistance could help clarify why such apparent deficits are not eliminated by selective pressure in the region, allowing development toward an extraintestinal pathogenic lifestyle and ultimately increasing the odds of the high-risk bloodstream NTS infections.

**MATERIALS AND METHODS**

**Mice and experimental coinfection model**

Female C57BL/6 mice (6 to 8 weeks old; stock no. 000664) were purchased from the Jackson Laboratory for infection experiments. For generating *Plasmodium*-parasitized blood stocks, CD-1 mice were purchased from Charles River Laboratory. Mice were housed under specific pathogen-free conditions and used for experiments at 8 to 11 weeks of age. For most experiments, at least five mice were used in each group, with multiple cages of mice (two to five mice per cage) used for each group to limit the possibility of cage effects. All animal experiments were approved by the Institution of Animal Care and Use Committee at the University of California, Davis (UC Davis).

**Plasmodium infections**

*P. yoelii* parasite stock was obtained from the Malaria Research and Reference Reagent Resource and was maintained and expanded by passage through CD-1 mice. Blood from multiple CD-1 mice infected with parasite was collected by cardiac puncture, pooled, and mixed 1:2 (v/v) with freezing solution (10% glycerol and 90% Alsever’s solution [Sigma-Aldrich]) for storage in liquid nitrogen. For mock infections, blood was collected from uninfected CD-1 mice and similarly preserved.

C57BL/6J mice were infected with *P. yoelii* parasite upon reaching 8 to 11 weeks of age. Parasitized blood stocks were diluted to 1 × 10^8 red blood cells/ml with 0.9% saline. Mice were then inoculated intraperitoneally with 0.1 ml of diluted parasitized blood. For mock infections, uninfected control blood was diluted with an equivalent volume of saline, and 0.1 ml was injected intraperitoneal into the mice. Parasite infection was tracked through a combination of weight loss, food consumption, blood cell counts, and parasite burden in the blood. Anemia (reduced circulating blood cell counts) and parasite burden were assessed from blood collected from tail snips. For circulating blood cell counts, tail blood was diluted 1:1000 in phosphate-buffered saline (PBS), and cell concentration was assessed using a TC20 Automated Cell Counter (Bio-Rad Laboratories Inc.). Counts were normalized to the average counts from the mock-treated animals collected and measured at the same time. Parasitemia was determined by examination of Giemsa-stained (Harleco) thin blood smears to enumerate the percentage of red blood cells containing detectable *P. yoelii* parasites.

**Salmonella strains**

A complete list of *Salmonella* strains used in this study can be found in table S1. Unless otherwise indicated, bacteria were routinely grown on lysogeny broth (LB) agar or MacConkey agar plates and cultured aerobically at 37°C in LB supplemented with antibiotics at the following concentrations for selection when appropriate: nalidixic acid (Nal), 0.05 mg/ml; carbenicillin (Carb), 0.1 mg/ml; kanamycin (Kan), 0.1 mg/ml; and chloramphenicol (Cm), 0.05 mg/ml. Construction of most mutant strains used in this study has been described in prior work (9, 22, 60–64). *S. Typhimurium invA spiB Kan^R^ (FF559, IR715 ΔinvA ΔspiB phoN::KSAC) was generated by P22 transduction of phoN::KSAC from IR715 phoN::KSAC into IR715 ΔinvA ΔspiB (SPN487), in the manner previously described for the production of *S. Typhimurium invA spiB Cm^R^ (FF183) (60). *S. Typhimurium invA spiB Cm^K^ + pAM34 (GTW58) was produced by transforming the plasmid pAM34 (22) into FF183 by heat shock, and maintained through culturing in LB with carbenicillin and 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG).

**Salmonella infection and colonization readouts**

Single colonies of *S. Typhimurium* grown on selective agar plates were inoculated into LB supplemented with the appropriate antibiotics for selection and incubated with shaking (200 rpm) at 37°C for 14 to 18 hours. Cultures were pelleted by centrifugation (10 min, 4000g, 4°C) and washed with sterile LB without antibiotics. Pelleted *S. Typhimurium* was resuspended in LB and adjusted to the appropriate bacterial density for infections. For single-strain infections, mice were inoculated orally by a pipette tip with 0.02 ml of *S. Typhimurium* at a density of approximately 5 × 10^8 colony forming units (CFU)/ml. This high inoculum dose was used to help limit variability in colonization between mice in the groups and to reduce nondetection of *Salmonella* in some mice that can occur at lower doses for a more accurate comparison of intestinal burden. For competitive infections, the strains were prepared separately, adjusted to 1 × 10^10 CFU/ml, then mixed 1:1 (v/v) and inoculated as 0.1 ml by oral gavage. Inocula were serially diluted and plated for CFU to confirm accuracy of the concentration and the input strain ratio for calculations in competitive infections. *S. Typhimurium* inoculations occurred in the morning, between 06:00 and 12:00. Mice were euthanized at the indicated time points or when they became moribund. Mouse euthanized early due to health concerns were excluded from analysis.

Intestinal contents (approximately 20 to 100 mg) of euthanized mice were collected from the relevant portions of the intestines into 1 to 2 ml of PBS and homogenized by vortex. Samples were then serially diluted in PBS and plated on appropriate selective agar to assess CFU loads. If no *S. Typhimurium* could be recovered after challenge, then the load was set to the detection limit for statistical comparisons (100 CFU/g intestinal content). In the *S. Typhimurium* challenges lasting 4 hours or less, mice were excluded from the analysis if *S. Typhimurium* could be detected in the small intestine, but no CFUs were detected in the cecal or colon content, indicating that the inoculum had not yet reached the large intestine at the point of collection. In that case, using the detection limit as the CFU load in the cecum or colon for statistical analysis would be inaccurate. In the competitive infections, intestinal content was plated on media selecting for both inoculated strains (LB + naladixic acid) and the
mutant strain alone (LB + chloramphenicol). Wild-type S. Typhimurium (IR715) burden was determined by subtraction of the mutant numbers from the overall (LB + nalidixic acid) numbers, and competitive index was calculated as the wild-type:mutant load, corrected for the input ratio of the inoculum.

In vivo pAM34 growth assays
To compare the approximate in vivo rate of S. Typhimurium in P. yoelii–infected and mock-infected animals, mice were perorally inoculated with 1 × 10^9 CFU S. Typhimurium invA spiB Cm^R + pAM34 (GTW58). As has been previously described (22), replication of the pAM34 plasmid is under control of the LacI repressor region, which allows it to be replicated and maintained by Salmonella when grown in high concentrations of lactose or a lactose analog such as IPTG but cannot replicate and is rapidly lost as the bacteria divide in low-lactose environments such as the adult mouse. Thus, loss of the plasmid at a population level correlates with the overall replication of the bacteria in the low-lactose environment, which allows for assessment of the replication rate (23).

To prepare inocula of S. Typhimurium invA spiB Cm^R + pAM34, single colonies grown on LB supplemented with carbenicillin (0.1 mg/ml) and 1 mM IPTG (for plasmid maintenance) were inoculated into LB broth (supplemented with IPTG alone) and incubated with shaking (200 rpm) at 37°C for 12 hours. Then, the plasmid was diluted to better assess the early replication rate by subculturing the inoculum (1:100) in fresh LB without IPTG and allowed to grow for an additional 3 hours before harvesting the bacteria by centrifugation and infecting the mice perorally, as previously described. Immediately following infection of the mice, the inoculum was serially diluted in PBS and plated on MacConkey and MacConkey + Carb + 1 mM IPTG to quantify the ratio of pAM34-containing S. Typhimurium to the total inoculated S. Typhimurium count. At 4 or 24 hours after inoculation, mice were euthanized, and intestinal contents were collected and plated on MacConkey to quantify S. Typhimurium burden and determine the fraction of the population maintaining pAM34. In parallel to the mouse infections, serial dilutions of the inoculum were cultured in LB for an additional 4 to 24 hours then plated on MacConkey and MacConkey + Carb + IPTG to generate a standard curve correlating population-level pAM34 plasmid loss to number of replications, as has been previously described (23).

In vitro pH bacterial survival assays
To assess bacterial sensitivity to low pH, PBS was acidified to different pH levels between 2 and 5 with hydrochloric acid and aliquoted into separate tubes. S. Typhimurium invA spiB Kan^R and strains of Escherichia coli strain Nissle 1917, E. coli O157:H7 EDL933 (ATCC 43895), C. rodentium DBS100 (ATCC 51459), Shigella flexneri M90T (ATCC BAA-2402), and Y. enterocolitica subsp. enterocolitica (ATCC 700823) were grown overnight in LB broth and prepared for inoculating mice as described above, except washed and resuspended using PBS instead of LB, to an optical density at 600 nm (OD_{600}) of 1.0, and plated for quantification of initial CFUs. The inoculum was then diluted 1:100 into the different low-pH PBS tubes and incubated at 37°C with shaking for 1 hour. Samples were serially diluted in fresh PBS (pH 7.4) and plated on MacConkey agar to quantify remaining bacterial concentrations.

Gastric pH assessment
Following euthanasia, stomachs of mice were removed from the body cavity. The microelectrode PH-N and reference probes (Unisense) were immediately inserted into the untreated gastric antral lumen for assessment of pH, and measurements on the Unisense Multimeter (in millivolts) were recorded and converted to pH values using a standard curve generated from concurrent readings of reference standards at pH 2, 4, and 7. The probes were rinsed with 70% ethanol and distilled water between measurements.

Histamine H2R agonist (dimaprit) administration
Mice were inoculated with P. yoelii–infected or control blood as previously described. In the morning of day 6 after infection, mice received a single dose of the histamine H2R agonist dimaprit dihydrochloride (Tocris Bioscience) or a mock treatment with the vehicle. Dimaprit was administered intraperitoneally as 0.1 ml of dimaprit dihydrochloride (40 mg/ml) dissolved in sterile saline, and the vehicle (saline) was administered in an equivalent volume. One hour later, mice were challenged with S. Typhimurium for 4 hours and then assessed for gastric pH and intestinal S. Typhimurium burden.

Omeprazole administration
Mice were inoculated with P. yoelii–infected or control blood as previously described. In the afternoon on days 3, 4, and 5 after infection, mice began receiving daily treatments with either the proton pump inhibitor omeprazole (Sigma-Aldrich) or a mock treatment with the vehicle. Omeprazole was administered intraperitoneally as 0.1 ml of a suspension (30 mg/ml) in 1% Tween 80 (Sigma-Aldrich) in Dulbecco’s phosphate-buffered saline (Gibco), and vehicle was administered in an equivalent volume. At 6 dpp, mice were challenged with S. Typhimurium for 24 hours and then assessed for gastric pH and intestinal S. Typhimurium burden.

Anti–TNF-α antibody administration
Mice were inoculated with P. yoelii–infected or control blood as previously described. On days 3 and 5 after infection, mice received a dose of either Ultra-LEAF Purified anti-mouse TNF-α antibody (clone: MP6-XT22, BioLegend Inc.) or Ultra-LEAF Purified Rat IgG1, α isotype control antibody (BioLegend Inc.). Both antibodies were diluted to 2 mg/ml in dPBS (Gibco) and administered as 0.25 ml per mouse, intraperitoneally, for a dose of 0.5 mg per mouse (approximately 25 mg/kg body weight). At 6 dpp, mice were either euthanized for relevant measurements or challenged with S. Typhimurium for 3 hours then euthanized for gastric pH and intestinal S. Typhimurium burden.

RNA extraction and real-time PCR expression analyses
Following lumen content removal, tissues were snap-frozen in liquid nitrogen at necropsy and stored at −80°C. RNA was isolated from tissues using TRI Reagent (Molecular Research Center). Briefly, whole tissues were suspended in TRI Reagent and homogenized by glass bead–beating for 1 min then treated with chloroform and centrifuged to separate the phases. The aqueous phase was removed and mixed with 95% ethanol, then applied to a silica membrane column (EconoSpin,Epoch Life Science), and washed with 3 M sodium acetate. The column was treated with PureLink DNase (Invitrogen) to remove genomic DNA contamination and washed twice with 10 mM Heps in 70% ethanol, and RNA was eluted in RNase-free water for quantification.

For analysis of mRNA expression, 1 µg of total RNA per sample was reverse-transcribed into complementary DNA (cDNA) in a 50 µl of reaction using MultiScribe Reverse Transcriptase (Thermo Fisher Scientific). The cDNA was then amplified in a real-time PCR instrument (Cepheid) using primers specific to the gene of interest.

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and circulating TNF-α were determined by Mann-Whitney tests. In all comparisons, \( P < 0.05 \) was considered statistically significant.

**Software**

The following software were used: Microsoft Excel for Mac, Prism 8 for macOS (GraphPad Software), QuantStudio Real-Time PCR System version 1.3 (Applied Biosystems), EVOS FL Auto 2 Imaging System (Thermo Fisher Scientific), and Image/FIJI version 2.0.0-rc-69/p1.520 (https://imagej.net/Fiji/Downloads) (66, 67).

### SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/7/27/eaab6232/DC1

### View/request a protocol for this paper from Bio-protoal.

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