Helicobacter infection decreases basal colon inflammation, but increases disease activity in experimental IBD

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ABSTRACT

Background: Helicobacter species are best known for their roles in the pathology of gastritis; however, several Helicobacter species also colonize the intestine, and less is known about effects of Helicobacter on the development of intestinal inflammation. To evaluate contributions of Helicobacter in inflammatory bowel disease, we investigated whether and how pre-existing intestinal colonization would affect disease severity and biomarkers of inflammation in experimental IBD.

Materials and Methods: Mice were infected with H. muridarum 2 weeks prior to induction of colitis mediated by 3% dextran sulfate (DSS). Disease activity index, stool blood and consistency, colon length, myeloperoxidase, histopathology, blood and lymphatic vessels, and numbers of dilated mucosal crypts were measured in control, DSS-only, H. muridarum-infected, and H. muridarum-infected + DSS mice. Results: Prior to DSS challenge, H. muridarum-infected mice showed little distal gut injury by several indices of colon inflammation with decreased blood vessel density in the submucosa, and lower lymphatic density in the mucosa and submucosa. However, after DSS colitis, H. muridarum-infected mice exhibited significantly greater disease. Weight change, stool bleeding, diarrhea, and angiogenesis were all increased in H. muridarum-infected mice in DSS colitis compared to DSS controls. Conclusions: Our data show that Helicobacter colonization of the intestine, unlike that of the stomach, lowers basal gut inflammatory scores, but increases disease activity and inflammation in an acute colitis model. Intestinal Helicobacter infection may therefore represent a significant sub-clinical factor which predisposes the gut to inflammatory injury.

Keywords: Helicobacter; Colon; Lymphatics; Crohn’s Disease; Ulcerative Colitis

1. INTRODUCTION

Helicobacter is a genus of Gram-negative bacteria that includes Helicobacter pylori (H. pylori), which is well-adapted for colonization of the human stomach via binding to gastric epithelial cells [1]. In less than 30 years since its discovery, H. pylori has been associated with the causation of gastric ulcers and gastric cancer [2-4]. In addition to H. pylori, various Helicobacter species have been discovered in other mammalian organ systems, including the intestinal tract [5-7].

Inflammatory bowel disease (IBD) includes Crohn’s disease (CD) and ulcerative colitis (UC) [8]. CD is characterized by transmural inflammation/injury at any point along the gut, while ulcerative colitis involves mucosal damage that is limited to the colon [8,9]. Despite their differences in topographical distribution, CD and UC share several characteristics, including malabsorption, protein-losing enteropathy, cachexia, edema, and diarrhea. In addition, blood and lymphatic vessel expansion and remodeling have been reported previously in active IBD [10-12]. Angiogenic mediators, such as vascular endothelial growth factors (e.g., VEGF-A), fibroblast growth factor, and inflammatory cytokines (e.g., TNF-α), contribute to IBD pathophysiology by triggering activation of vascular endothelial cells and expansion of the microvasculature, which leads to increased lymphocyte infiltration and vascular leakage [13,14]. In contrast,
lymphatic vessels are largely important because of their function to remove inflammatory cells and cytokines. Consequently, the lymphatic expansion seen in IBD is likely a compensatory mechanism allowing clearance of inflammatory substances that accumulate during inflammation. Van Kruiningen and Colombel actually propose that dysfunctions of the intestinal lymphatics, such as lymphocytic and granulomatous lymphangitis have a significant role in IBD pathogenesis. Several animal studies show that obstruction of small intestinal lymphatic vessels results in intestinal dysfunction similar to IBD [13, 15]. Therefore, IBD etiology may reflect both increased leukocyte accumulation and cytokine release from activated, angiogenic microvessels but also decreased removal of these substances by dysfunctioning lymphatic networks.

Several models of ileitis and colitis suggest that bacteria are necessary, but insufficient triggers of IBD [16]. Several other studies have reported that Helicobacter may modulate IBD. For example, H. maccaceae has been linked with chronic idiopathic colitis in young rhesus monkeys [17]. Further, a study of children with CD reported PCR evidence for Helicobacter infection in 59% of children with CD vs. 9% of healthy controls [18]. Similarly, Laharie et al. found that evidence of H. pullorum or H. canadensis infection was significantly associated with CD in adults [19]. Of particular interest, the study showed that enterohepatic Helicobacter species predominated, rather than H. pylori [19]. Finally, H. canis, another enterohepatic Helicobacter, has also been detected in duodenal ulcerations in CD [20]. Therefore, it appears that certain Helicobacter species may be related to the pathogenesis of IBD. However, the exact mechanism of Helicobacter involvement is still undiscovered.

In this study, we examined the effect of H. muridarum infection on the development of acute gut injury caused by DSS-treatment. H. muridarum infects the intestines of mice and is typically considered a commensal organism by DSS-treatment. In this case, we also report that changes in disease activity, such as stool blood, weight loss, etc., were significantly increased in mice pre-infected with H. muridarum compared to mice treated with DSS only. Finally, we discovered that initially H. muridarum infection causes a decrease in blood vessel density, but an increase in the density of blood vessels is seen after the stress of the DSS-treatment that is greater than the increase seen in the DSS-treated mice treated without H. muridarum pre-infection. Therefore, it appears that H. muridarum may cause changes in blood vessel density that lead to an increase in disease activity.

2. MATERIALS & METHODS

2.1. Isolation and Propagation of H. muridarum Strain TM1

We obtained gastric homogenates from mice that were believed to have been colonized with candidates H. heilmannii (generously provided by Dr. Jani O’Rourke, University of New South Wales, Australia). Scrapings from the frozen homogenate were inoculated into Ham’s F-12 medium supplemented with 1% FBS, β-cyclodextrin, vancomycin, trimethoprim, and 5-fluorocytosine. Once growth of a highly-motile, spiral-shaped organism was apparent, efforts were begun to isolate the organism in pure culture. Aliquots of the culture were inoculated onto Campylobacter blood agar, plates. A small amount of culture was stabbed into F-12 soft agar [25], and new broth cultures were inoculated using medium drawn from the culture surface to minimize transfer of non-motile organisms. Cefsulodin and colistin were added, in addition to the above antimicrobial agents. H. muridarum-strain TM1 was unable to form colonies on any of the solid media tested, but was able to grow in F-12 soft agar containing 4% FBS. Organisms were taken from the edge of the soft agar growth region and re-inoculated into broth. Limiting dilution was also carried out on broth cultures. All organisms obtained from limiting dilution or soft agar were identical in morphology and displayed the same rapid back-and-forth motility as the organism seen in the original culture.

PCR of 16S ribosomal DNA from pure cultures was carried out using Helicobacter genus-specific primers [26]. The resulting product was sequenced by the Arizona State University DNA Laboratory. The strain was determined to be H. muridarum by comparing the 16S sequence with those of other bacteria using the NCBI Blast search engine. The sequences of the two isolates were identical to each other and 99% identical to the 16S ribosomal RNA gene sequence from H. muridarum ATCC 49282 over the sequenced stretch of 669 nucleotides. The only discrepancy was due to an “N” in our sequence data. There was a single nucleotide mismatch with H. muridarum strain ST1. H. muridarum TM1 was
and usually caused a color change within 30 seconds. The solution turns from yellow to red within 15 minutes. Our urea, 63 µg/ml phenol red). If urease is present, the characteristic shape and motility. A rapid urease test was used to further confirm that recovered organisms were by adding 10 µl of culture to 100 µl of urease detection muridarum. The presence of urease activity was detected by adding 10 µl of culture to 100 µl of urease detection solution (1 mM sodium phosphate buffer, pH 6, 33 mM urea, 63 µg/ml phenol red). If urease is present, the solution turns from yellow to red within 15 minutes. Our H. muridarum strain has extremely high urease activity and usually caused a color change within 30 seconds.

2.2. Infection of Murine Intestines

H. muridarum was grown in 25 cm² tissue culture flasks containing Ham’s F-12 medium supplemented with 1% FBS under conditions described above. After microscopically verifying appropriate morphology and motility, the culture was centrifuged at 4°C at 7840 × g for 15 minutes and the pellet was resuspended in 3 ml 0.9% NaCl. 200 ul aliquots of the concentrated H. muridarum suspension were administered orally to each mouse using a gavage needle.

Infection was confirmed by stool culture two weeks post-infection (data not shown). Fresh fecal pellets were collected and placed in microfuge tubes with F12 medium containing 1% fetal bovine serum and antimicrobial agents capable of suppressing most non-Helicobacter fecal organisms. The tubes were spun at 10,000 × g and then incubated at 37°C for 10 minutes. The upper portion of the supernatant was then removed and added to a well in a 24-well plate containing F12 + 5% FBS and antibiotics, as described above. This procedure enriched for motile bacteria, thus reducing contamination by the bulk of fecal organisms, which are non-motile or weakly motile. Antibiotic selection further reduces growth of unwanted microorganisms. Cultures were monitored daily by phase contrast microscopy for organisms with the criteria established by Cooper et al. [29]. This scoring system includes edema, extent of injury, leukocyte infiltration, crypt abscesses, and loss of goblet cells. In this grading system, inflammation severity was scored on a 0 - 3 scale, (0: none; 1: slight; 2: moderate; 3: severe), the extent of injury was scored on a 0 - 3 scale (0: none; 1: mucosal; 2: submucosal; 3: transmural), and crypt damage was scored on 0 - 4 scale (0: none; 1: basal 1/3 damaged; 2: basal 2/3 damaged; 3: only surface epithelium intact; 4: loss of entire crypt and epithelium). Each value was multiplied by an extent index from 1 - 4 that reflects the amount of involvement for each section (1: 0% - 25%, 2: 26% - 50%, 3: 51% - 75%, 4: 76% - 100%). At least 3 sections from each colon were analyzed to produce each score value. A maximum possible histopathological score for this assay is 40.

2.3. Induction of Experimental Colitis

Four groups of animals were used: control uninfected mice (CU), control H. muridarum-infected mice (MU), DSS-treated uninfected mice (CD), and DSS H. muridarum-infected mice (MD). Experimental colitis was induced in C57B6 mice (n = 5) by supplementing drinking water with 3% DSS as described in Ganta et al. (DSS, MW = 36 - 50 kDa; ICN Biomedical, Costa Mesa, CA); mice in the control groups (n = 5) group received tap water without DSS [28]. Mice were inoculated with H. muridarum ≥ 2 weeks prior to initiation of DSS treatment in order to make sure that infection was well-established.

3% DSS administration produces an erosive distal colitis with an initial onset at 3 - 4 days that is characterized by progressive weight loss, diarrhea, occult blood, leukocyte infiltration, colon shortening, loss of intestinal epithelial barrier, and histopathological changes in colon structure [29,30]. Mice typically lose ~20% body weight by day 10 with continuous administration of 3% DSS. Over 9 days, mice were given unrestricted access to pellet diet (Purina, St. Louis, MO), and tap water (control), or tap water containing 3% DSS in “colitis” groups. Mouse weight, stool form, and occult blood were recorded daily. On day 9, mice were sacrificed by cardiac puncture while under ketamine/xylazine anesthesia (k/x anesthesia). Mouse weight, stool form, and occult blood were recorded daily. On day 9, mice were sacrificed by cardiac puncture while under ketamine/xylazine anesthesia (k/x anesthesia). Body weight change is defined as the percentage difference in body weight for each mouse as a fraction of the starting (day = 0) weight.

Tissues were analyzed for histopathological injury using the criteria established by Cooper et al. [29]. This system includes edema, extent of injury, leukocyte infiltration, crypt abscesses, and loss of goblet cells. In this grading system, inflammation severity was scored on a 0 - 3 scale, (0: none; 1: slight; 2: moderate; 3: severe), the extent of injury was scored on a 0 - 3 scale (0: none; 1: mucosal; 2: submucosal; 3: transmural), and crypt damage was scored on 0 - 4 scale (0: none; 1: basal 1/3 damaged; 2: basal 2/3 damaged; 3: only surface epithelium intact; 4: loss of entire crypt and epithelium). Each value was multiplied by an extent index from 1 - 4 that reflects the amount of involvement for each section (1: 0% - 25%, 2: 26% - 50%, 3: 51% - 75%, 4: 76% - 100%). At least 3 sections from each colon were analyzed to produce each score value. A maximum possible histopathological score for this assay is 40.

2.5. Immunohistochemical Analysis

Colon preservation in formalin, sectioning of the tissue, and immunohistochemistry were completed as described in Ganta et al., 2010 [28]. Staining for vascular endothel-
bial growth factor receptor-3 (lymphatic vessels) and mouse endothelial cell antigen-32 (blood vessels) was performed. Once stained, the vessels in each layer of the entire colon wall were counted.

2.6. Tissue MPO

Tissues were prepared as described previously [32]. Briefly, the mouse colons were dissected and homogenized in phosphate buffer (5 mM; pH 6). The samples were centrifuged at 30,000 × g for 30 minutes at 4°C. The supernatant was discarded, and the pellet was washed again in phosphate buffer. Then the pellet was suspended in 50 mM phosphate buffer (pH 6, 25°C), which contained 0.5% hexadecyltrimethylammonium bromide. The samples were frozen and thawed 3X with sonication between each cycle. Protein concentrations in samples were determined by BCA assay (Bio-Rad). Samples were sonicated for 6 secs, and 20 μl added to the wells (n = 3). 180 μl of phosphate buffer (50 mM, pH6), 0.0167% o-dianisidinedi HCl (ICN.), and 1% H2O2 was added to each 96-well and incubated at 37°C. A450nm was measured at 80 min. The activity is expressed as change in absorbance per 60 minutes per mg of protein.

2.7. Ethical Considerations

Animal protocols were approved by the LSU Institutional Animal Care and Use Committee. All mice were monitored daily for disease activity; any animals showing restricted movement or loss of 20% body weight were humanely sacrificed following terminal ketamine/xylazine anesthesia (k/x = 150/60 mg/kg via intraperitoneal injection) and thoracotomy.

2.8. Statistical Analysis

All statistical analysis was completed using GraphPadInStat 3. Comparisons between the four groups (CU, CD, MU, and MD) were completed using one-way ANOVAs with Tukey-Kramer post-tests. Repeated measures one-way ANOVAs with Dunnett’s post-tests were used on comparisons between day 0 scores and later scores from within the same group. Statistical significance was defined as p < 0.05.

3. RESULTS

3.1. H. muridarum Infection Exacerbates Increases in Disease Activity Index

The Disease Activity Index (DAI) was measured for all four groups of mice (Figure 1(a)). Throughout the 9 days of the study, DAI scores for CU and MU mice were constant at day “0”. The DAI scores for CD and MD mice rose as time passed, and the MD scores were significantly higher than the CU and MU mice scores after day 2. The CD scores became significantly higher than the CU and MU scores on days 6 - 9. Further, DAI scores of the MD mice were significantly higher than the scores of CD mice on days 6 - 8, indicating an earlier worsening of the disease state in MD mice. Finally, the MD mice had a significantly higher DAI score than at day “0” after day 2. MD mice also showed significant disease activity 2 days earlier than CD mice.

Weight loss was measured in CU, MU, CD, and MD mice (Figure 1(b)). The % of original weight was similar for all groups up until day 3. From day 4 onward, the % original weight of CU mice was significantly lower than MU. The MD mice had significantly greater weight loss on days 7 - 9 than CD mice. Both the CD and MD mice exhibited a large weight loss compared to the CU or MU mice. After day 6, the difference between the MD mice and the CU or MU mice was significant, whereas the % original weight of the CD mice was significantly less only on day 9. The MD mice lost a large portion of their body weight, approximately 23% by day nine. By comparison, the CD mice lost approximately 15% of their body weight by day nine. The MD mice also had significantly less weight than on day “0” by day 6 of measurements. However, CD mice did not have significant weight loss until day 7. Therefore, the weight loss was accelerated by approximately one day.

Diarrhea (stool form) (Figure 1(c)), was evaluated to determine effects of treatments on each group of mice. The stool grade for CU and MU mice remained stable for the duration of the study. Conversely, the grade of the MD and CD mice increased steadily throughout the study. By day 4, scores were significantly greater than the CU or MU mice. On days 7 and 8, the MD mice had a significantly higher stool grade than the CD mice. Furthermore, the MD mice did not have higher stool grade scores any earlier than the CD mice. Therefore, the H. muridarum did not affect the time of onset for changes in stool consistency.

Stool blood measurements (Figure 1(d)) showed that the stool blood in CU and MU mice remained stable at “0” throughout the study. Both CD and MD mice showed significantly increased stool blood after day 5 compared to CU and MU mice. At days 6 - 9, the scores of occult blood were significantly higher in MD mice compared to CD mice. The presence of more blood in the stool of MD mice vs. CD mice suggests that these combined stresses increase tissue injury and/or capillary fragility. MD mice had significantly elevated stool blood by day 4. For the CD mice, this occurred on day 5. Once again shows that colitis may progress more quickly in the H. muridarum-infected mice.
Figure 1. (a) Both the CD and MD mice show a steady increase in DAI (p < 0.001) from CU and MU mice on days 6 - 8. The MD mice also show a higher DAI than the CD mice on days 3 (p < 0.01), 6 (p < 0.001), 7 (p < 0.01), and 8 (p < 0.001). (b) CD mice show a decrease (p < 0.001) in weight after day 7, and MD mice show a decrease (p < 0.001) in weight after day 6. MD mice also showed a significantly larger decrease in weight than the CD mice on days 7 - 9. (c) CD and MD mice had higher stool grades than CU and MU mice beyond day 3. MD mice have a higher stool grade than CD mice on days 7 and 8. (d) Both CD and MD mice had higher amounts of blood in their stool compared to the CU and MU mice beyond day 5. The MD mice had a significantly higher amount of blood in the stool compared to CD mice on days 6 - 9.

3.2. *H. muridarum* Infection Has Several Pathological Effects on Colitis Mice

A small decrease (non-significant) difference in colon length was seen in the MU mice compared to the CU mice. Both CD and MD mice showed a significant reduction in colon length compared to the CU and MU mice (Figure 2(a)). The MD mice also had a shorter colon than the CD mice, but the difference was not significant.

Histological analysis of the murine intestinal tissue sections showed that a significantly higher amount of dilated mucosal crypts was present in the CD and MD mice when compared to the CU and MU mice. Interestingly, the MD mice also had approximately 2.5 times the number of the abnormal crypts than the CD mice. Therefore, the MD mice show a great increase in what appears to be a histological marker of normal colitis (Figure 2(b)).

MPO activity was assayed in colon samples of all 4 subsets of mice and expressed as avg ± SEM. The change in absorbance at 450 nm/60 min/mg of sample from CD mice (0.058 ± 0.009) was significantly higher (p < 0.05) than the absorbance levels of the CU mice (0.035 ± 0.008). The change in absorbance of samples from MU (0.034 ± 0.006) and MD (0.03012 ± 0.015) mice were not higher than the absorbance of the control samples. Furthermore, the absorbances of the CU, MU, and MD samples were very similar. Therefore, MPO activity is not increased in either type of *H. muridarum*-infected mouse, but the activity is increased in CD mice (Figure 2(c)).

Histopathological score of the colon tissue was measured for all four different groups of mice after they were
sacrificed on day 9 using H&E-stained tissue sections (Figure 2(d)). The CD and MD mice had significantly higher scores than the CU or MU mice (p < 0.001). However, there was no significant difference between the CU and MU mice or the CD and MD mice (data not shown). Furthermore, no significant differences were seen between these mice on any of the individual subscales: severity of inflammation, extent of injury, and crypt damage (data not shown).

3.3. *H. muridarum* Infection Alters Lymphatic Vessels

Lymphatic vessels were visualized in the mucosa, submucosa, and serosal layers of the colon using VEGF-receptor 3 (VR3) immunohistochemistry (Figure 3(a)). The stained slides were used to determine the abundance of lymphatic vessels in the mucosa (Figure 3(b)), submucosa (Figure 3(c)), and serosa (Figure 3(d)) of each mouse. In the mucosa, MU mice had a significantly lower level of lymphatic vessels than the CU mice (p < 0.05). The MU mice also displayed significantly lower levels of lymphatic vessels than both the CD and MD mice (p < 0.001). Therefore, the infection greatly decreased the amount of lymphatic vessels whenever no disease is induced. The MD mice illustrated a significant increase in lymphatics compared to the CU mice (p < 0.001). This shows that *H. muridarum* infection increases the amount of lymphatic vessels whenever disease is induced via DSS. The increase is possibly a compensatory mechanism for the greatly reduced number of lymphatics that is induced by the infection when disease is not present. An increase in lymphatics in CD mice compared to CU mice was also observed but was not significant. The amount of lymphatic vessels was also greater in MD mice than in CD mice, but the difference was not significant.
In the submucosa, MU mice had significantly lower levels of lymphatic vessels than the CU, CD, and MD mice (p < 0.001). The CU, CD, and MU mice had roughly equivalent numbers of lymphatic vessels in the submucosa. In the serosa, the MD mice had significantly lower levels of lymphatics than the CD (p < 0.001) and MU (p < 0.05). The CD mice had increased lymphatic vessels compared to the CU mice, but the difference was not significant. The MD mice had a greater amount of lymphatics than the CD mice, but this difference was also not significant. Finally, the MD mice showed significantly greater levels of lymphatic vessels than the CU mice (p < 0.001).

3.4. *H. muridarum* Infection Increases Blood Vessel Abundance in DSS Mice

Blood vessels were visualized in the mucosa, submucosa, and serosal layers of the colon using mouse endothelial cell antigen-32 (MECA-32) immunohistochemistry (Figure 4(a)). The stained slides were used to quantify the abundance of blood vessels in the mucosa (Figure 4(b)), submucosa (Figure 4(c)), and serosa (Figure 4(d)) of the colon. In MD mice, the quantity of blood vessels in the mucosa was significantly greater than in the CU, CD, and MU mice (p < 0.001). The quantity of vessels in the mucosa of CU, CD, and MU mice was similar. In the submucosa, MD mice had a significantly greater number of blood vessels than the CU, CD, and MU mice (p < 0.001). Furthermore, the MU mice had significantly lower amounts of blood vessels than the CU mice (p < 0.05). As before with lymphatics, the submucosa also contained the most blood vessels out of the three layers of intestinal tissue.

In the serosa, the CU (p < 0.001) and MU (p < 0.01)
mice had significantly fewer blood vessels than the MD. The CD mice also had fewer blood vessels than the MU mice, but the difference was not significant. Moreover, the CD mice exhibited an increase (not significant) in blood vessels from the CU and MU mice.

4. DISCUSSION

Since the H. pylori—gastritis connection was discovered, many studies have described the role of this bacterium in gastric mucosal injury. In addition to epithelial injury, another important proposed target of H. pylori pathogenesis is the gastric mucosal microcirculation [33]. Kalia et al. described that toxigenic H. pylori increased leukocyte adhesion, platelet aggregation, post-venular capillary vasoconstriction and leakage of labeled albumin [34,35]. Several other studies have demonstrated that H. pylori-induced microvascular disturbances may involve platelet activating factor and nitric oxide (NO), since hexanolamine-PAF and L-arginine prevented increased platelet emboli and thrombi in mucosal capillaries and post-capillary venules [36]. P-selectin also participates in these forms of platelet leukocyte binding [37], and aggregation of the platelets is known to participate in the recruitment of leukocytes [38].

Additionally, mast cells and mast cell-derived histamine may contribute to microcirculatory disturbances associated with H. pylori. Atuma et al. showed that mast cell stabilization (with ketotifen) or the PAF antagonist WEB2086 blocked H. pylori extract induced vasoconstriction [39]. Kalia et al. showed that ketotifen and pyrilamine, an H1 receptor antagonist, prevent increased macromolecular leakage of labeled albumin in response to H. pylori extracts [36]. Kurose et al. found that albumin leakage produced by H. pylori was decreased by mast cell stabilization, or blocking antibodies against CD11b/CD18, ICAM-1 or P-selectin [37].

While Helicobacter colonizes the gastric mucosa and causes local injury, Helicobacter has also been found in...
other areas of the gastrointestinal tract including the liver, gallbladder, and colon [40]. Despite the large amount of research on *H. pylori*-induced gastric mucosal injury, much less has been accomplished on the extragastric effects of *H. pylori*. However, those few studies have illustrated that Helicobacters (*pylori*, etc.) also disturb microcirculatory integrity in acute pancreatitis, and increase the severity of ischemia-induced pancreatitis [33]. In addition, lipopolysaccharide released by *H. pylori* triggers high rates of NO production through iNOS-induction [41], which has been associated with increased vascular permeability in the heart, kidney, liver, and lung [42]. Some investigators have also studied the effects of *H. pylori* in coronary artery disease. Pellicano *et al.* have shown a higher occurrence of *H. pylori* infection in patients with acute myocardial infarction [43]. Prasad *et al.* studied the effect of *H. pylori* pathogen burden on endothelial dysfunction. They discovered that an elevated pathogen burden, (which included *H. pylori* infection plus normal commensal bacteria and viruses), was a significant risk factor for endothelial dysfunction and coronary artery disease [44].


Extra-gastric effects of *H. pylori* have also been studied in the large intestine where it apparently provokes different responses than in the stomach. For example, in a DSS colitis model, Luther *et al.* have shown that *H. pylori* DNA decreases dendritic cell production of proinflammatory cytokines, (IFN-γ and IL-12), which might lessen IBD severity [45]. However, *E. coli* DNA produced an intense inflammatory edema, irritation, bleeding and diarrhea. This reaction to *E. coli* DNA was mediated by toll-like receptor (TLR) 9 expressed on dendritic cells which produced Type 1 IFNs, which in this study was associated with increased vascular permeability in the heart, kidney, liver, and lung [46]. Some investigators have also studied the effects of *H. pylori* in coronary artery disease. Pellicano *et al.* have shown a higher occurrence of *H. pylori* infection in patients with acute myocardial infarction [43]. Prasad *et al.* studied the effect of *H. pylori* pathogen burden on endothelial dysfunction. They discovered that an elevated pathogen burden, (which included *H. pylori* infection plus normal commensal bacteria and viruses), was a significant risk factor for endothelial dysfunction and coronary artery disease [44].

Vijay-Kumar *et al.* reported anti-inflammatory effects of TLR-3 activation mediated by binding of synthetic viral RNA which reduces gut injury in experimental colitis [48]. Because Helicobacter clearly may activate several types of TLRs (TLR-3, -9), the roles of TLRs in gut inflammation are complex. Luther *et al.* revealed that the microbial DNA of *H. pylori* has the ability to block inflammatory stimulation produced by *E. coli* DNA in the colon. Although Luther *et al.* showed that *H. pylori* DNA has anti-inflammatory effects, their findings exclude several alternate mechanisms and mediators active during Helicobacter colonization and do not completely address how Helicobacter colonization might affect the initiation and progression of colitis.

### 4.2. Non-*Pylori* Helicobacter Infection May Increase Gut Injury in IBD

There appears to be some association between Helicobacter and gut injury in IBD. Kaakoush *et al.* found that Helicobacteriaceae were present in 41.5% of children with CD, while only present in 22.5% of normal children (p = 0.0062). Interestingly, these Helicobacter species were not gastric *H. pylori*, but rather other Helicobacter strains found in the liver, gall bladder, and intestine [49]. Enterohepatic Helicobacter has also been found in the intestines of animals [20]. Using a Helicobacter-specific PCR, *Helicobacter canis*, [a species of Helicobacter that colonizes the colon of canines] was recovered from superficial ulcerations of the superior and descending duodenum in human CD [20]. This discovery indicates that Helicobacter colonization of the gut may correlate with increased susceptibility to IBD. In another model, Helicobacter colonization of the liver was associated with lower production of Th1 cytokines (IL-1β, IFN-γ, TNF-α) and Th17 (IL-17A) [40]. In our hands, *H. muridarum*-decreased vascular density in intestinal wall layers under basal conditions, but increased it during active inflammation (DSS colitis). Helicobacter contributions to intestinal injury may be complex and state-dependent.

Because Helicobacter within and outside the gastrointestinal tract appear to alter microcirculatory physiology, changes in microvascular structure and function could contribute to gut pathophysiology as is suggested in Helicobacter-associated atherosclerosis and cardiovascular risk. In this study, we found an increase in blood vessels within the mucosa and submucosa of MD mice that was even higher than the increase seen in CD mice. Furthermore, we discovered that the mucosa and submucosa of MU mice had less lymphatic vessels than the CU mice. This decrease in lymphatic vessels could have played a role in worsening of colitis in the infected mice due to a lessened initial capability to evacuate lymph and immune cells from the area of disease. Therefore, it appears that
alterations in microcirculation do play a role in contributing to the gut pathophysiology of Helicobacter infected mice.

Our study has shown that the DAI in MD mice is increased early on when compared to CD mice. However, by the end of the 9-day study, the difference seemed to have disappeared, and both groups of mice had similar levels of disease activity. Furthermore, when the mice colons were scored for histopathological changes after sacrifice at day 9, there was no difference in the scores of MD and CD mice. This seems to indicate rapid disease progression in MD mice that culminates in a similar level of disease as the CD mice by the end of disease development. Further studies need to be performed to determine if the histopathological disease state of MD mice is actually worse on earlier days in the study.

Despite evidence from both our study and previous studies that Helicobacter does contribute to IBD gut injury, we saw no increase in the amount of MPO activity in the colon of either the MD or the MU mice. On the other hand, the CD mice saw approximately a doubling of activity. Interestingly, this suggests that neutrophil infiltration of the diseased colon in the colitis mice infected with H. muridarum is less than the infiltration in regular colitis mice. This change may be related to the decreased submucosal blood vessel abundance seen in the MU mice. The decrease in blood vessels could lead to a decrease in the ability to recruit neutrophils to the disease area. Therefore, the gut would be unprepared to handle the type of corrosive injury that occurs when DSS is administered. Even without the increase in neutrophils, greater disease activity index scores are seen in the MD mice than in the CU, CD, and MU mice. Therefore, further studies are needed to determine which immune cells are actively involved in the immune responses in these MD mice.

4.3. Is Helicobacter Infections a Risk Factor for Colorectal Cancer?

In MD mice, the colon shows significantly more dilated mucosal crypts than were observed in CD colons. Crypt dilation is not uncommon in mouse models of colitis, but the number of abnormal crypts was significantly greater than that in the other three groups of mice, including the CD mice. Yoshizawa et al. have previously shown that H. pylori infection alters gland morphology in the stomach, including the dilation of mucous glands [50]. Furthermore, H. pylori has been convincingly shown to play a role in the development of gastric cancer [50]. This is particularly relevant because abnormal mucosal crypts, such as those seen in high numbers in the MD mice, have been identified as precursor markers of colon carcinogenesis [51]. Since the high numbers of abnormal crypts are not seen in the MU mice, it is possible that the additional stress induced by DSS colitis when added to H. muridarum provokes these pre-cancerous histological changes in the colon mucosa. Consequently, H. pylori infection, like H. muridarum, may significantly exacerbate the formation of intestinal cancers in IBD-associated colon cancer, and the appearance of these dilated mucosal crypts may be a warning sign that precedes the onset of colon cancers. However, much more work must be done before we can know if this is actually the case.

Sugimoto et al. have also found an interesting link between H. pylori and carcinogenesis wherein angiotensin II type I receptor mRNA levels were linked to the extent of inflammatory cell infiltration into the gastric mucosa. They also found a significant increase in the amount of angiotensin II type I and type II receptor mRNA levels in gerbils with gastric ulcers. These findings show a possible link between the renin-angiotensin system (RAS) and H. pylori infection. This is important because the RAS system is involved in angiogenesis required for carcinogenesis to proliferation. Therefore, there may be a link between the RAS and gastric (or possibly intestinal) carcinogenesis related to H. pylori infection [52]. Interestingly, chronic H. pylori infection of the stomach was also associated with increased rates of intestinal metaplasia [50]. This finding strongly suggests that gastric colonization with H. pylori produces changes in the intestine, particularly in response to gut injury as we see here. Because H. muridarum-colonization significantly increased crypt dilation, a prominent feature of intestinal metaplasia, the intensification of angiogenesis and metaplastic changes seen may represent an important combination of risk factors for development of subsequent CRC. Because Helicobacter might directly affect the intestine through gut colonization and indirectly influence CRC development through induction of gastrin secretion, (a stimulus for intestinal mucosal epithelial growth and a risk factor for CRC), Helicobacter infection might represent an important risk for hyper intense gut injury and malignancy and needs to be evaluated by prospective correlative trials [53].

5. CONCLUSION

In this study, we infected mice with H. muridarum to study its effects on the onset and severity of IBD. We found that H. muridarum increased disease activity and caused earlier disease onset. We also discovered differences in lymphatic and blood vessel densities between the infected and non-infected mice. All of these observations point to the possibility of an active and important role for Helicobacter colonization in the onset and worsening of IBD that involve microvasculature alterations.
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REFERENCES


**ABBREVIATIONS**

Dextran Sodium Sulfate (DSS), Inflammatory Bowel Disease (IBD), Myeloperoxidase (MPO), Disease Activity Index (DAI), Control Uninfected Mice (CU), Control *H. muridarum*-infected Mice (MU), DSS-treated Uninfected Mice (CD), and DSS *H. muridarum*-infected Mice (MD)