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Infection-induced colitis in mice causes dynamic and tissue-specific changes in stress response and DNA damage leading to colon cancer

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Helicobacter hepaticus-infected Rag2−/− mice emulate many aspects of human inflammatory bowel disease, including the development of colitis and colon cancer. To elucidate mechanisms of inflammation-induced carcinogenesis, we undertook a comprehensive analysis of histopathology, molecular damage, and gene expression changes during disease progression in these mice. Infected mice developed severe colitis and hepatitis by 10 wk post-infection, progressing into colon carcinoma by 20 wk post-infection, with pronounced pathology in the cecum and proximal colon marked by infiltration of neutrophils and macrophages. Transcriptional profiling revealed decreased expression of DNA repair and oxidative stress response genes in colon, but not in liver. Mass spectrometric analysis revealed higher levels of DNA and RNA damage products in liver compared to colon and infection-induced increases in S-chlorocystosine in DNA and RNA and hypoxanthine in DNA. Paradoxically, infection was associated with decreased levels of DNA etheno adducts. Levels of nucleic acid damage from the same chemical class were strongly correlated in both liver and colon. The results support a model of inflammation-mediated carcinogenesis involving infiltration of phagocytes and generation of reactive species that cause local molecular damage leading to cell dysfunction, mutation, and cell death. There are strong correlations among histopathology, phagocyte infiltration, and damage chemistry that suggest a major role for neutrophils in inflammation-associated cancer progression. Further, paradoxical changes in nucleic acid damage were observed in tissue- and chemistry-specific patterns. The results also reveal features of cell stress response that point to microbial pathophysiology and mechanisms of cell senescence as important mechanistic links to cancer.

mass spectrometry | biomarkers

Chronic inflammation is a major risk factor for many human diseases, including cancer. Epidemiological studies suggest that at least 20% of all cancers are caused by chronic inflammatory conditions, such as the causative role of Helicobacter pylori and hepatitis B virus infections in gastric and liver cancer, respectively, and the strong association of inflammatory bowel disease (IBD) and colon cancer (1, 2). Although the underlying molecular mechanisms linking inflammation and cancer are poorly understood, many of the steps have been defined. In the case of infection, the inciting event is tissue injury, with local release of cytokines and other chemotactic factors causing infiltration and activation of macrophages and neutrophils to produce large quantities of other cytokines and chemokines as well as reactive chemical species that cause mutagenic and cytotoxic damage (3, 4). For example, up-regulation and activation of inducible NO synthase (iNOS) and NADPH oxidase in macrophages produces nitric oxide (NO) and superoxide (O2•−), respectively. Subsequent reactions yield a battery of reactive species, including the potent nitrosating agent nitrous anhydride (N2O3) and the oxidizing and nitrating agents nitrogen dioxide radical (NO2•), peroxynitrite (ONOO•−), and nitrosoperoxycarbonate (ONOOCO2−) (5). Moreover, activated neutrophils are considered the primary source of the potent oxidant and halogenating agent hypochlorous acid (HOCl) via activation of myeloperoxidase (MPO) (6–8).

These chemical mediators of inflammation can damage all classes of cellular molecules, including DNA, RNA, protein, lipids, and metabolites, by both direct and indirect mechanisms. For example, DNA damage caused by halogenation, deamination, and oxidation, as well as by nucleobase adduct formation, can potentially induce de novo mutations and epigenetic changes initiating tumor development (1, 2, 9). Several recent studies involving mouse models of inflammation have provided evidence supporting such a scenario (10–12). Similar pathways disrupt the function of RNA, as proposed for major human degenerative diseases such as Alzheimer’s disease, Parkinson’s disease, and atherosclerosis (13, 14), whereas protein damage caused by oxidation, nitration, and halogenation can impair function and stability, including those of tumor suppressors, oncogenes, and factors that maintain fidelity during DNA replication (4, 15, 16).

To better understand the complex pathophysiology of inflammation, we undertook a comprehensive analysis of chemical and biological end points in an established model of human IBD: Rag2−/− mice infected with Helicobacter hepaticus (17, 18). This Gram-negative spiral bacterium colonizes the liver and intestinal crypts of the cecum and the colon of several mouse strains, establishing a life-long infection (19, 20). H. hepaticus does not typically cause disease in immunocompetent mice, but infection in susceptible inbred strains can lead to hepatitis whereas immunodeficient mice develop chronic colitis (21–23). For example, mice lacking the recombine-activating gene-2 (Rag2) do not possess functional lymphocytes, and H. hepaticus infection results in chronic colitis and colon cancer (17). H. hepaticus infection in Rag2−/− mice emulates many aspects of human IBD, with colonic and cecal infiltration of macrophages and neutrophils presumably causing increased production of reactive chemical species at sites of infection (18).


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To test this hypothesis and identify potential biomarkers of the pathophysiology of inflammation, we characterized the time course of colon and liver pathology in H. hepaticus-infected Rag2−/− mice with a battery of quantitative molecular, histological, and biochemical analytical tests. For example, we developed an isotope-dilution liquid chromatography-coupled tandem mass spectrometry (LC-MS/MS) method (SI Appendix, Fig. S1) to quantify 14 different DNA and RNA damage products anticipated to represent the full spectrum of inflammation chemistries. This included the 2-deoxyribo- and ribonucleoside forms of base halogenation (5-chlorocytosine, 5-CI-C), nitrosative deamination (xanthine, X; hypoxanthine, H), alkylation by lipid peroxidation products (1, 4-N2-ethenoadenine, εA; 3,N4-ethenocytosine, εC), and oxidation (8-oxo-7,8-dihydorguanine, 8-oxo-G; spiroiminodihydantoin, Sp; guanidinohydantoin, Gh) (see SI Appendix, Fig. S2). Data from these chemical analyses were correlated with quantitative scores from histopathological, immunohistochemical, and gene expression analyses. These analyses revealed a variety of unique features of infection-induced colitis and carcinogenesis.

Results

Histological Analysis of Colon and Liver from H. hepaticus-infected Rag2−/− Mice. To monitor and characterize disease progression, colon and liver tissues were harvested at two different time-points following H. hepaticus infection: 10 and 20 weeks post-infection (w.p.i.). Quantitative PCR analysis revealed that all infected Rag2−/− mice were positive for H. hepaticus, whereas the sham controls were negative. The tissue samples were subjected to a wide range of analyses, starting with histopathology and immunohistochemistry. Although there were minimal or no lesions in large intestine and liver in age-matched control groups, H. hepaticus-infected mice showed significant tissue pathology at both time-points (Fig. 1, SI Appendix, Fig. S3). At 10 w.p.i., changes in the large intestine included inflammation, edema, epithelial defects, crypt atrophy, hyperplasia, and dysplasia, which were similar and most severe in the ileocecal junction and proximal colon (SI Appendix, Fig. S3 A and B). The most prominent histopathological changes consisted of thickened colonic mucosa, infiltration of neutrophils and macrophages, and combinations of surface epithelial tattering or erosion and mucosal gland dilation or distortion with intraluminal cell debris. Where colonic dysplasia was a feature, mucosal glands were variably distorted and lined by crowded epithelial cells, often containing increased numbers of mitoses and reduced goblet cell differentiation. These changes were significant in the proximal and transverse colon but not statistically significant in the distal colon (SI Appendix, Fig. S3B). At 20 w.p.i., lesions in the large intestine were most severe in the ileocecal junction and proximal colon and were similar to those observed at 10 w.p.i. (Fig. 1A, SI Appendix, Fig. S3 A and B). However, progression of dysplasia to intramural carcinoma was evident in some animals at 20 w.p.i., with carcinoma development characterized by invasion of neoplastic glands below the muscularis mucosae layer (SI Appendix, Fig. S3C). Glands were irregularly shaped and lined by atypical, tall columnar epithelial cells with indiscernible borders, cellular and nuclear pleomorphism, visible mitotic figures, and loss of goblet cells. In the liver, there was significant evidence of portal and lobular hepatitis but not interface hepatitis, at both 10 and 20 w.p.i. (SI Appendix, Fig. S3 C and D).

In terms of disease progression, the histological activity indices (HIAs) of colitis (cumulative score of inflammation, edema, epithelial defects, crypt atrophy, hyperplasia, and dysplasia) showed significantly lower levels in the proximal colon at 20 w.p.i. compared to 10 w.p.i. but tended to increase in the distal colon at 20 w.p.i. (P = 0.06) (Fig. 1C and D). In agreement with the HAI in the proximal colon, the hepatitis index (HI) was significantly lower at 20 w.p.i. than at 10 w.p.i. (Fig. 1D). Consequently, the HAI of the proximal colon and the HI tended to correlate in H. hepaticus-infected mice (SI Appendix, Table S1).

To define the extent of the innate immune cell infiltrate in this infection model of colitis, we performed immunohistochemical analyses using F4/80 and MPO staining as markers for macrophage and neutrophil infiltration, respectively (Fig. 2, SI Appendix, Fig. S4). At both time-points, F4/80-positive and MPO-positive cells accumulated significantly and sustainably in the cecum and the proximal colon of H. hepaticus-infected Rag2−/− mice. However, between 10 and 20 w.p.i., we observed a significant decrease in MPO-positive cells in the cecum, with a trend toward a decrease in the proximal colon (P = 0.07), whereas there was a trend towards increased macrophage infiltration at 20 w.p.i. in the cecum and the proximal colon (P = 0.07). This is consistent with the view that neutrophils are recruited early during the inflammatory response, whereas macrophages persist longer at sites of inflammation (24).

LC-MS/MS Quantification of Inflammation-Related DNA and RNA Damage. Toward the goal of defining the chemical mechanisms underlying inflammation-induced carcinogenesis, we used LC-MS/MS (SI Appendix, Fig. S1) to quantify the nucleoside forms of DNA and RNA damage products thought to represent the full spectrum of inflammation-related chemistries: nucleobase deamination (X and I), halogenation (5-CI-C), oxidation (8-oxo-G, Sp, Gh), and alkylation (εA, εC) (SI Appendix, Fig. S2). This platform extends previously published DNA protocols (25, 26) to include the guanine secondary oxidation products Sp and Gh, the halogenated nucleobase 5-CI-C, and a spectrum of RNA damage products (SI Appendix, Tables S2 and S3). Special attention was given to minimize adventitious damage by addition of

Fig. 1. H. hepaticus-infected mice develop chronic colitis, colon carcinoma, and sporadic hepatitis. (A–C) HAIs of colitis were calculated by summing individual scores of inflammation, edema, epithelial defects, crypt atrophy, hyperplasia, and dysplasia for (A) cecum, (B) proximal colon, and (C) distal colon. (D) HIs were calculated by summing individual subfeature histopathology scores from lobular, portal, and interface inflammation, and the number of lobes that contained ≥5 inflammatory lesions. Statistical analysis for all panels is presented as box plots showing the median value (line), the interquartile range (box), and Tukey whiskers embracing data within 1.5-fold of the interquartile range; all data outside the range of the Tukey whiskers are presented as individual data points.
deaminase inhibitors (coformycin, tetrahydrouridine), a metal chelator (deferoxamine), and an antioxidant (butylated hydroxytoluene) during sample preparation (SI Appendix, Fig. S1). LC-MS/MS parameters were optimized for maximal sensitivity resulting in limits of quantification (LOQ) of 1–10 fmol, which amounts to 0.4–8 lesions per $10^8$ nucleotides (nt) in 100 μg of DNA or RNA, with basal levels of all DNA and RNA damage products above the LOQ in liver and colon from uninfected mice (Fig. 3, SI Appendix, Figs. S5 and S6; SI Appendix, Tables S2 and S3).

This analytical platform was applied to liver and colon tissues from H. hepaticus-infected and control Rag$^{2−/−}$ mice at 10 and 20 w p.i. (SI Appendix, Tables S2 and S3). Analysis of DNA and RNA damage products in control mice revealed several important and unique findings. First, basal damage levels were significantly higher in RNA than in DNA for base deamination (X, I), oxidation (8-oxo-G), and chlorination (5-Cl-C) but not for lipid peroxidation-derived adducts (Fig. 3). Second, background levels of the guanine secondary oxidation products Sp and Gh were detected in the range of 1–7 lesions per $10^8$ nt in DNA from the liver and colon, which is nearly 100 times lower than the parent 8-oxo-dG (Fig. 3). Sp and Gh have previously only been observed in vitro and in prokaryotes (reviewed in ref. 27). Third, the levels for all guanine-derived oxidation products were significantly lower in the colon compared to the liver, both in DNA and RNA (Fig. 3). The most prominent differences were observed for Sp and Gh, which were close to the LOQ in colon and significantly higher than the LOQ in the liver. Fourth, the levels of chemically related damage products showed strong and highly significant positive correlations, as demonstrated by correlations between Sp and Gh with their parent molecule 8-oxo-dG and between the lipid-peroxidation-derived etheno-adducts εA and εC in DNA and RNA, respectively (SI Appendix, Fig. S7). These observations satisfy assumptions about the chemical origins of these lesions.

The analytical methods were next used to define changes caused by H. hepaticus infection. Although no significant differences in damage levels were observed in the liver from infected mice, analyses of colon tissue revealed several significant findings, the most prominent of which was the increase in halogenated lesions. Statistically significant increases in 5-Cl-dC (50%) and 5-Cl-rC (33%) were observed at 20 w p.i., along with significant increases in dI (26%) in DNA (Fig. 4). Unexpectedly, we observed transient but significant decreases in εdA (35%), εdC (44%), and 8-oxo-G in RNA (25%) at 10 w p.i. (Fig. 4). The levels of other damage products in DNA and RNA remained unchanged in the infected mice compared to controls (SI Appendix, Tables S2 and S3).

Statistical analyses revealed several significant correlations between the nucleic acid damage and histopathology. Not surprisingly, simple correlation analysis of levels of DNA and RNA lesions with corresponding colonic HAI values revealed significant positive correlations for 5-Cl-dC, dI (Spearman coefficients $r > 0.5$, $P < 0.05$ (SI Appendix, Table S4), with weak tendencies...
In terms of genes that distinguished H. hepaticus infection, 42% and 61% of the genes in the colon showed significantly different expression levels at 10 and 20 w p.i., respectively (Fig. 5, SI Appendix, Fig. S9B and Table S8), whereas 3.4% and 28% of the genes in the liver showed significantly different expression levels, respectively (Fig. 5, SI Appendix, Fig. S9C and Table S8). In infected colons at 20 w p.i., there was significant up-regulation of 80% of genes encoding proteins involved with the generation of reactive chemical species, including inducible nitric oxide synthase (Nos2), xanthine-oxidoreductase (Xdh/XO), and several genes coding for NADPH oxidases, such as Nox1, Cyb, and Ncf2 (Fig. 5, SI Appendix, Table S8). Moreover, we observed up-regulation of genes thought to protect against oxidative stress, such as cell cycle checkpoint gene Cdkn2a, and epithelium-specific glutathione peroxidase 2 (Gpx2), thioredoxin reductase 1 (Txnrd1), and glutathione reductase (Gsr). Paradoxically, several genes of the stress response group (36%), such as Sod and Gst, and most of the genes involved in DNA repair (64%) showed significantly lower expression in H. hepaticus-infected colon compared to uninfected controls (Fig. 5, SI Appendix, Table S8). All major DNA repair pathways were represented in the down-regulated group: base excision repair (BER: Nbh1, NEIL3, and MUTYH), nucleotide excision repair (NER: Xpa, Xpc, Ercc6), mismatch repair (MMR: Mlh3, Msh3, Msh6), and double-strand break repair (DSBR: Prkdc, Mre11, Brcal). With the exception of Ercc3 and Mlh1, all genes of the NER and MMR groups were significantly down-regulated at 20 w p.i. Importantly, none of the housekeeping genes (i.e., 18S rRNA, Actb, Gapdh, Hprt1) were significantly altered by infection, which rules out a general suppression of gene expression in inflamed tissues. Interestingly, markers of cell proliferation, such as Pcn and Ki67, did not change in spite of histological evidence for cell proliferation in the infected colon. Ki67 expression was previously observed to increase only in H. hepaticus-infected mice in the infection-induced hepatocellular carcinoma tissue, with no increase in normal liver tissue (30).

Although some changes in gene expression were common to the liver and colon, such as up-regulation of Xdh/XO at 20 w p.i. (Fig. 5), there were several major infection-induced differences in the liver. Of note, expression of MPO was very highly induced in the liver, which may be related to inducible MPO expression in monocyes, because neutrophilic MPO is expressed only during early stages of neutrophil maturation in the bone marrow (7). Further, in direct contrast to the colon, H. hepaticus infection caused up-regulation of all major DNA repair pathways in the liver, including oxidative dealkylation of nucleobases (Alkhh1 and Alkhh3), BER (Lig1, NEIL3, and Ung), NER (Ercc6), MMR (Pms2), and DSBR (Brcal, Blm, and Exo1) (Fig. 5B). This is consistent with compensation for inflammation-induced DNA damage, which is further supported by up-regulation of several cell cycle checkpoint regulators, such as Tcp53, Atr, Gadd45a, and Cdkn2a, in liver tissues from infected mice. On the other hand, several genes involved in stress response and glutathione metabolism, such as Cat, Gsta1, and Gsta4, were expressed at significantly lower levels in the liver of infected mice compared to controls (Fig. 5).

Discussion

IBD affects millions of people worldwide and significantly increases the risk of colon cancer, with >20% of IBD patients developing colitis-associated cancers within 30 y of disease onset with a mortality rate >50% (4). Although dysregulation of host-microbiome interactions, multifactorial genetic predisposition, and environmental factors are associated with IBD (4, 31), the molecular and chemical mechanisms linking colonic inflamma-
tion with cancer remain unclear, with potential involvement of tumor-initiating DNA damage, epigenetic changes, and interference with tumor-suppressive mechanisms (32). A central feature of current presumptive models of inflammation-mediated carcinogenesis involves infiltration of activated phagocytes at sites of infection, with subsequent generation of reactive oxygen, nitrogen, and halogen species that cause molecular damage leading to cell dysfunction, mutation, and cell death. Our comprehensive analysis of the H. hepaticus-infected Rag2−/− mouse model of colitis-associated carcinogenesis (17, 18) supports this model, with strong correlations among histopathology (Fig. 1), phagocyte infiltration (Fig. 2), and the increased level of HOCl-induced 5-Cl-dC (Fig. 4). The major results of these studies are summarized in Table 1 and Fig. 6, which reveals several unique features of the chemical and biological pathology linking inflammation and cancer in the mouse model of infection-induced colitis: strongly correlated histopathology and molecular damage suggestive of a major role for neutrophils in the inflammatory process, paradoxical changes in nucleic acid damage that are both tissue- and chemistry-specific, and features of cell stress response that are consistent with both known microbial pathophysiology and mechanisms of cell senescence.

Histopathological Correlates of Colitis and Hepatitis in H. hepaticus-Infected Rag2−/− Mice. H. hepaticus colonizes the liver and colon of various mouse strains and is linked with the development of chronic colitis and hepatitis in several murine models (20). Although the complex population of immune cells in the intestinal lamina propria is thought to balance immune tolerance of luminal microbiota with pathogen defense (31), the lack of...
regulatory T cells in Rag2−/− mice disturbs the balance, with infection-induced inflammation leading to cancer (33). The present study demonstrated that H. hepaticus-infected Rag2−/− mice develop typhlocolitis (i.e., inflammation of the cecum and the colon) that evolved into colon cancer by 20 w p.i. (Fig. 1, SI Appendix, Fig. S3). Consistent with a diagnosis of typhlocolitis, the HAI s of the cecum and the proximal colon correlated moderately but with high statistical significance (r = 0.35, P < 0.01) in H. hepaticus-infected animals (SI Appendix, Table S1) and with infiltration of neutrophils and macrophages. Although we observed a progressive increase in the severity of disease in the cecum and the distal colon, histologic activity indices decreased significantly between 10 and 20 w p.i. in the proximal colon. This illustrates the spatiotemporal dynamic nature of colitis development in this model. Carcinomas were observed at the ileoceccolic junction in infected mice (10% of infected mice at 10 w p.i. and 20% at 20 w p.i.), which was one of the regions affected most by inflammation and histopathological changes (SI Appendix, Fig. S3A). This resembles neoplasia in human IBD patients, where carcinomas arise from colitis-associated dysplastic epithelial foci (20).

This study also demonstrates that H. hepaticus-infected Rag2−/− mice temporally develop hepatitis (Fig. 1B, SI Appendix, Fig. S3 C and D) that tended to correlate with the severity of colitis (r = 0.24, P = 0.07; HI versus HAI proximal colon) (SI Appendix, Table S1). In general, the hepatitis was mild to moderate at both time points, but sporadic incidents of severe hepatitis were consistently observed at 10 w p.i. The finding that the severity of colitis and hepatitis tended to correlate (SI Appendix, Table S1) suggests that both disease states are functionally and mechanistically linked, as had been observed in other models (e.g., ref. 34).

**Determinants of the Spectrum of DNA and RNA Damage Products in Inflamed Colon and Liver.** One of the major conclusions of our analysis of 14 different DNA and RNA damage products is that the goal of developing biomarkers from damage products caused by reactive species generation at sites of inflammation is hampered by our lack of understanding of the factors that govern the steady-state levels of damage products in an inflamed tissue. These include the nature of the agent causing the inflammation, the tissue involved, and the response of the affected cells. Although the methods used in the present studies allow analysis of damage in the total population of cell types present in inflamed colon and liver tissues, the results of the DNA and RNA damage analyses provide some unique insights into the factors governing molecular damage in inflamed tissues.

Consider first the complicated local chemistry of molecular damage in inflammation, starting with reactive halogen species generated in vivo by MPO. Although our previous pathologist-focused studies of H. hepaticus infection in the Rag2−/− revealed that the use of an iNOS inhibitor to block NO production caused a reduction in colonic inflammation, hyperplasia, and dysplasia, a more significant reduction in colon pathology was achieved by depleting Gr−/− neutrophils (18). Coupled with the fact that the major source of MPO is neutrophils (7), these observations point to neutrophil-dependent MPO-activity as a major contributor, along with macrophage-generated NO, to the colitis and colon cancer arising in H. hepaticus-infected mice. Our observations provide immunohistochemical and chemical confirmation of the role of reactive halogen species, with higher levels of 5-Cl-dC and 5-Cl-dG in H. hepaticus-infected colons and a persistent and high frequency of MPO-positive cells at sites of inflammation (Figs. 2 and 4). These results are all consistent with other observations of major roles for neutrophils in colitis, including the protective effect of selective neutrophil depletion and scavengers of neutrophil-derived HOCl (7, 35, 36) and the observation of neutrophil accumulation and increased MPO activity in intestinal lesions in IBD patients (7, 37–39). Further, chlorination-derived damage products have also been associated with other inflammatory conditions, such as higher levels of 5-chlorouracil in human atherosclerotic tissue (40) and dihalogenated 2′-deoxyctydine.
in lungs and livers of mice treated with proinflammatory lipopolysaccharide (41). It has been observed that 5-Cl-dC mimics 5-methyl-dC and induces inappropriate DNM1T methylation within CpG sequences (9, 42, 43), which may result in altered cytosine methylation patterns that could affect the expression of tumor suppressor genes and initiate carcinogenesis (9, 42, 43).

Localized oxidation chemistry also appears to play an important role in the spectrum of DNA and RNA lesions. Oxidants generated by normal respiration and activated phagocytes are predicted to cause oxidation of guanine in DNA to form 8-oxo-dG. Many studies support this model, with measurements of 8-oxo-dG from 0.5 to 4 lesions per 10^6 nt (10, 44, 45) agreeing with our analyses in the liver and colon (Fig. 3). Indeed, Sipowicz et al. demonstrated a two-fold increase in 8-oxo-dG in DNA from livers of A/JNCr mice infected with *H. hepatica*, with evidence for oxidative stress induced by increased cytochrome P450 activity (46). However, 8-oxo-dG itself is approximately 1,000-fold more susceptible to oxidation than dG and in vitro studies have demonstrated that 8-oxo-dG oxidation results in secondary oxidation products such as Sp and Gh (26, 47–49). The only previous in vivo evidence for the formation of 8-oxo-dG secondary oxidation products comes from studies of oxidatively stressed *Escherichia coli* (50), in which Sp was detected at levels of approximately 10–100 lesions per 10^6 nucleotides. Using a more sensitive analytical method, we have now been able to detect background levels of both Sp and Gh in both the liver and colon in mice (Fig. 3), at levels some 100 times lower than those observed in *E. coli*. The in vivo derivation of Sp and Gh from 8-oxo-dG in the liver and colon is supported by the significant positive correlations between the adduct levels (*SI Appendix, Fig. S7*). The lack of significant infection-induced increases in the steady-state levels of 8-oxo-dG, Sp, or Gh suggests that either reactive chemical mediators of inflammation present in this study do not cause oxidative DNA damage, or that efficient DNA repair balances the rate of oxidant-induced formation. It is important to point out, however, that the background steady-state level of Sp and Gh can contribute to the mutagenic burden in cells given that Sp is at least an order of magnitude more mutagenic than 8-oxo-dG (51).

Now consider the complications of DNA repair. The fact that we observed either no change or down-regulation of most DNA repair gene expression in infected colon at 10 and 20 w.p.i., respectively, does not preclude a significant DNA repair capacity with extant proteins or translational control of gene expression. Indeed, the significant reduction in the levels of the etheno adducts in DNA in infected colon at 10 w.p.i. (Fig. 4), and the suggestion of decreased levels of etheno adducts in the liver, is consistent with up-regulated DNA repair activity. In the case of etheno adducts, this is possibly related to alkyladenine DNA glycosylase (Aag), the loss of which has been shown to increase etheno adduct levels in colon DNA (11). To illustrate the complexity of damage and repair in inflammation, several studies showed higher levels of etheno adducts in inflammatory conditions in mice and humans (10, 11, 52, 53), whereas other reports noted lower levels of etheno adducts under pathological conditions (54, 55). In peripheral blood lymphocytes of colorectal carcinoma patients, levels of etheno adducts as well as DNA repair rates were lower than in healthy subjects (54). Further, Kirkali et al. observed lower levels of the purine oxidation products 4,6-diamino-5-formamidopyrimidine, 2,6-diamino-4-hydroxy-5-formamidopyrimidine (*FapyG*), 8-oxo-dG, and (5′S)-8,5′-cyclo-2′-deoxyadenosine in tumor tissue compared to surrounding normal tissue in patients with colorectal cancer (56).

Finally, our studies reveal tissue-specific differences in the levels of molecular damage products (Fig. 3). Kirkali et al. observed higher levels of DNA damage products in rectum compared to the colon (56), which corroborates our observations of tissue-specific differences in formation or repair of DNA damage.

Colitis-Associated Changes in Gene Expression Reflect Bacterial Pathophysiologv in *H. hepatica*-Infected Rag2^-/-^ Mice. Our comprehensive analysis of histopathology and molecular damage was complemented by an analysis of infection-induced changes in gene expression in colon and liver for 95 genes involved in the generation of reactive chemical species, stress response and glutathione metabolism, DNA repair, and cell cycle regulation. These changes in gene expression serve as biomarker signatures for both *H. hepatica* infection in general and for organ-specific effects of the infection, as suggested by principal component analysis (*SI Appendix, Fig. S8*). The scores plot clearly distinguishes normal and infected colon and liver, with the tissue-specific effects apparent in the first component and infection-specific changes distinguished in the second component (*SI Appendix, Fig. S8A*).

Whereas the loading plot in *SI Appendix, Fig. S8B* shows the genes that account for each of the four groupings apparent in the scores plot, several rather dramatic shifts in specific functional gene categories are more clearly illustrated in the volcano plots shown in Fig. 5 and *SI Appendix, Fig. S9*. For example, virtually all DNA repair pathway genes were significantly down-regulated in the infected colon at 10 and 20 w.p.i., including genes for MMR, BER, NER, and DSBR (Fig. 5C). This is somewhat paradoxical given the absence of infection-induced changes in the levels of most DNA damage products and the reduction in etheno adducts. However, as noted earlier, reduced gene expression may not correlate with the levels of functional repair proteins. The interesting complication here is that the DNA repair genes suppressed in the colon all showed increased expression in the liver (Fig. 5D), though to a smaller extent.

Our observation of *H. hepatica*-induced down-regulation of DNA repair transcripts is consistent with studies of *H. pylori* infection, which has been shown to suppress DNA repair gene expression in infected cells in culture and in infected mouse and human tissues (57, 58). Whereas Machado et al. observed that infection caused reduced expression of base excision repair and mismatch repair genes and concomitant increases in genomic instability (58), Kim et al. demonstrated that *H. pylori* coculture with gastric cancer cells lines caused reduced levels of both mRNA and protein for both MutS and MutL mismatch repair protein, with heat-sensitive bacterial products responsible for the reductions (57). Further, Toller et al. found that co-culture of *H. pylori* with mouse and human cell lines led to increased levels of DNA double-strand breaks (59). These observations of increased genomic instability and double-strand breaks are consistent with the DNA-cleaving activity of the cytolethal disintegrating toxin (CDT) produced by *H. hepatica*. In addition to inducing cell cycle arrest and cell distortion, CDT is essential for the carcinogenic effects of *H. hepatica* infection in liver cancer models (60), presumably by inducing single-strand breaks that lead to replication-dependent double-strand breaks and subsequent recombination-induced mutations.

Another category of genes showing differential expression in the liver and colon (Fig. 5, *SI Appendix, Fig. S9*) are stress response genes. Considering the normal, uninfected condition, the gene expression results shown in *SI Appendix, Fig. S9A* support the notion that colon is more resistant to stress than the liver, because several genes involved in oxidative defense pathways were expressed at significantly higher levels in the normal liver (e.g., *Cat*, *Sod*, *Gst*). On the other hand, several DNA repair genes, mainly involved in BER, homologous recombination, and MMR as well as genes involved in cell cycle checkpoint regulation, exhibited significantly higher expression levels in the normal colon, which may be related to the higher exposure of colon tissue to food-borne toxics. Interestingly, MUTYH and NEIL3, which were expressed at higher levels in the normal colon, are directly involved in oxidative stress repair, as MUTYH removes dA mispaired with 8-oxo-dG form the DNA backbone.
and NEIL3 is potentially involved in the repair of Sp and Gh (28, 29). The latter may directly account for lower levels of Sp and Gh in colon tissue compared to liver, as part of the generally lower levels of DNA lesions in the colon.

In terms of changes associated with *H. hepaticus* infection, several transcripts involved in DSBR and BER were up-regulated in the liver at 20 w p.i. (Fig. 5), which would be consistent with either hepatic infection by *H. hepaticus* or hepatic DNA damage caused by humoral mediators from the infected colon. Another example involves expression of GPx2, an intestinal epithelium-specific glutathione peroxidase that reduces hydroperoxides by oxidation of glutathione, which was significantly up-regulated in *H. hepaticus*-infected colons at both time points (Fig. 5A, SI Appendix, Fig. S9B). This may be functionally important in that the stability of glutathione peroxidases is mainly regulated by their mRNA stability (61). GPx2 was implicated previously in the prevention of inflammatory colitis in the mouse (61, 62). In accordance with our results, it was shown that GPx2 is up-regulated in human colorectal cancer tissue (63, 64) and consequently may serve as a marker for inflammatory bowel disease (65).

**Colitis-Associated Changes in Gene Expression Reflect Inflammation-Induced Senescence in *H. hepaticus*-Infected Rag2^−/−^ Mice.** Many of the changes in gene expression observed in *H. hepaticus*-infected Rag2^−/−^ mice are consistent with an infection-induced cell senescence, the growth arrest occurring when mitotically capable cells are subjected to any of a variety of stresses, such as the oncogenic stress of inflammation (66). One of the hallmarks of senescence is increased expression of apoptosis-inducing p16 (Cdkn2a) (66), the expression of which is increased significantly in both liver and colon at 20 w p.i (Fig. 5). p16 is up-regulated in premalignant lesions and lost in invasive colon carcinomas by Cdkn2a hypermethylation (67). The fact that *H. hepaticus*-generated CDT also induces p16 expression (68) suggests that a senescence response in the colon epithelium could result from both the infection itself and the resulting genotoxic stress of the inflammatory reaction. Associated with the increased p16 in these studies is reduced expression of DNA damage response activators ATM and ATR and their downstream partners *Chek1* and *Hus1*. Loss of ATM/ATR could result in senescence and genomic instability leading to cell death (69, 70). Furthermore, both the hyperplasia associated with infection (SI Appendix, Fig. S3A) and the infiltration to activated phagocytes (Fig. 2) leads to a predominance of terminally differentiated cells, which may resemble a state of senescence. Our results support the notion that up-regulation of p16 is an inflammation-induced tumor suppressor mechanism that causes cell cycle arrest and senescence in heavily damaged cells.

In summary, the results of our comprehensive analysis of the biological and chemical pathology of *H. hepaticus*-induced colitis and colon cancer and *H. hepaticus*-induced hepatitis in the Rag2^−/−^ mouse model provides a rigorous confirmation of a model of inflammation-mediated carcinogenesis involving activation of phagocytes at sites of infection, with subsequent generation of reactive oxygen, nitrogen, and halogen species that cause molecular damage leading to cell dysfunction, mutation, and cell death (Fig. 6). We have demonstrated strong correlations among histopathology, phagocyte infiltration, and molecular damage chemistry that suggest a major role for neutrophils in the inflammatory process and that reveal dynamic and unexpected changes in nucleic acid damage that are both tissue- and chemistry-specific. The results also reveal features of the cell stress response that point to microbial pathophysiology and mechanisms of cell senescence as important mechanistic links to cancer.

**Materials and Methods**

**Chemicals.** All chemicals were of the highest available quality and purchased from Sigma-Aldrich unless indicated otherwise. LC-MS grade acetonitrile was obtained from EMDS Chemicals. Water purified through a Milli-Q system was used throughout the study.

**Animal Husbandry, *H. hepaticus* Infection, and Tissue Collection.** All experiments were performed in accordance with protocols approved by the Massachusetts Institute of Technology Committee on Animal Care and with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. Rag2^−/−^ mice (129S6/SvEvTacRag2^−/−^) were used throughout the study and housed in a specific pathogen-free barrier facility. Animals were infected with *H. hepaticus* (strain 381, ATCC51449) grown and confirmed as pure culture as described previously (17, 19). Two groups of mice aged 4–6 w received 0.2 mL of fresh inoculums (infected group) or sterile media (control group) by gavage every other day for a total of three doses. Tissues from the first group of mice was harvested at 10 w p.i., whereas the second group of mice was euthanized at 20 w p.i. and tissue collected at necropsy. Tissues were removed by standard necropsy procedures. For histopathological evaluation the liver and entire large intestine, including the ileoceccolonic junction and colon, were harvested from each mouse. For DNA, RNA, and protein damage analyses, separate groups of mice (n = 5–7 animals per group) were analyzed, respectively, in order to obtain sufficient amounts of tissue. For RNA and DNA analyses, liver lobes and whole colons were immediately submerged in RNAlater solution (Qiagen), placed on dry ice, and stored at −80°C until further processing. Cecum and colon samples were analyzed by quantitative PCR (qPCR) using *H. hepaticus*-specific primers to confirm infection status (71).

**Histopathologic Evaluation.** All tissues were fixed in 10% formalin, paraffin-embedded, and stained with hematoxylin and eosin. The degree of lesion severity: 0 (absent), 1 (mild), 2 (moderate), 3 (marked), and 4 (severe). For the liver, an HI was calculated by combining individual scores for lobular, portal, and interface hepatitis, as well as the number of lobes (out of a total of four) that contained ≥5 inflammatory lesions. Hepatitis B was defined by a hepatitis index equal to or greater than 4. For the large intestine, lesions of inflammation, edema, epithelial defects, crypt atrophy, hyperplasia, and dysplasia were scored at separate locations (ileoceccolonic junction, proximal colon, and distal colon). A HAI was calculated by combining individual scores of inflammation, edema, epithelial defects, crypt atrophy, hyperplasia, and dysplasia.

**Immunohistochemistry.** Macrophages and neutrophils were identified in intestinal tissue sections by avidin-biotin complex immunohistochemistry using antibodies specific for myeloperoxidase (1:75; MPO; RB-373-A; Thermo Scientific), a neutrophil-specific marker, and F4/80 (1:150; MF48015; Caltag Laboratories), a macrophage-specific marker using previously established protocols (72). Briefly, ileocecal and colonic sections from 10 randomly selected animals (five males, five females) per group for each different time point were analyzed. Five randomly selected non-oriented sections per mouse (20 μm thick) sections were evaluated by a board-certified veterinary pathologist who was blinded to sample identity. The severity of lesions in the liver and large intestine was scored, using an ascending scale from 0 to 4, based on the degree of lesion severity: 0 (absent), 1 (mild), 2 (moderate), 3 (marked), and 4 (severe). For the liver, an HI was calculated by combining individual scores for lobular, portal, and interface hepatitis, as well as the number of lobes (out of a total of four) that contained ≥5 inflammatory lesions. Hepatitis B was defined by a hepatitis index equal to or greater than 4. For the large intestine, lesions of inflammation, edema, epithelial defects, crypt atrophy, hyperplasia, and dysplasia were scored at separate locations (ileoceccolonic junction, proximal colon, and distal colon). A HAI was calculated by combining individual scores of inflammation, edema, epithelial defects, crypt atrophy, hyperplasia, and dysplasia.

**Synthesis of Isotope-Labeled Internal DNA and RNA Standards.** Uniformly ^15^N-labeled internal standards of DNA damage adducts, d1,d, 8-oxo-dG, d8A, and d8C were synthesized as described previously (25). RNA damage products r1, r8, 8-oxo-rG, r8A, and r8C were synthesized accordingly, using uniformly ^15^N-labeled ribonucleosides (rA, rG, rC) as starting materials (Cambridge Isotope Laboratories). Uniformly ^13^C, ^15^N-labeled internal standards of Sp and Gh were synthesized as described previously (26). Internal standards of 5-chloro-2-deoxycytidine and 5-chloro-cytidine were synthesized according to a protocol as previously described (73). Standards were purified by HPLC, characterized by LC-MS, and quantified by UV absorbance using published extinction coefficients (25, 26).

**DNA and RNA Extraction.** To avoid adventitious nucleobase damage, deaminase inhibitors coformycin and tetrahydrodouridine, and antioxidants deferoxamine and butylated hydroxytoluene were added throughout sample workup, as described previously (25, 74). DNA from approximately 200 mg of colon and liver tissue was extracted using the DNA Isolation Kit for Cells and Tissues (Roche) with modifications as published previously (25). RNA from approximately 100 mg of liver and colon tissue was extracted using the RNeasy Midi Kit (Qiagen) following the manufacturer’s instructions with the
HPLC Prepurification of DNA and RNA. DNA (200 μg) was hydrolyzed to nucleosides by a combination of nuclease P1, DNase I, phosphodiesterase I (USB-Affymetrix), and alkaline phosphatase in the presence of deaminase inhibitors and antioxidants as described previously (25). The following amounts of internal standards were added to the digestion mixture: 1 pmol [5-3H]-dC, 1 pmol [5-3H]-dU, 1 pmol [5-3H]-dA, 100 fmol [3-3H]-3′-N2-oxo-dG, 100 fmol [13C5]-5′-S-CdC, 5 pmol [13C5]-G, and 2.9 pmol [13C5]-N6-Sp. A method for hydrolysis of RNA to ribonucleosides was adapted from a recently developed protocol. RNA (100 μg) was dissolved in 30 μL of sodium acetate buffer (30 mM, pH 6.8) containing 3.3 μM zinc chloride, 33 μM/mL coformincin, 167 μM tetrahydrodoline, 8.33 mM dexamethasone, 1.67 μM butylated hydroxytoluene and each of following internal standards: 10 pmol [5-3H]-Val, 1 pmol rL, 1 pmol [5-3H]-8-oxo-gG, 1 pmol [13C5]-N4-N2-oxa-rA, 1 pmol [13C5]-3′-N2-oxo-rC, and 1 pmol [13C5]-S-CdC. The RNA was hydrolyzed by incubation with 4 U of nuclease P1 (4 μL of 1 U/μL) at 37 °C for 3 h, followed by dephosphorylation by addition of 40 μL of sodium acetate buffer (30 mM, pH 7.8), 34 U of alkaline phosphatase (2 μL of 17 U/μL) and 0.2 U of phosphodiesterase I (USB-Affymetrix) at 37 °C, overnight. Enzymes were subsequently removed by microfiltration (Nanosep K10 Omega, Pall) and the filtrates concentrated under vacuum.

HPLC Preparation of DNA and RNA products. Hydrolyzed DNA preparations were dissolved in 100 μL water and individual 2′-deoxyribonucleosides were resolved and collected using an Agilent 1100 series HPLC system at empirically determined retention times (SI Appendix, Fig. S5 and Table S5) as described previously (25). The nucleoside-containing fractions were desiccated under vacuum, redissolved in 45 μL water, and analyzed by LC-MS/MS.

Based on a recently developed protocol, hydrolyzed RNA preparations were dissolved in 100 μL water and individual ribonucleosides were resolved and collected by HPLC using a Phenomenex Synergi C18 reversed-phase column (250 × 4.6 mm, 4 μm particle size, 80 Å pore size). Gradient elution was performed with a combination of acetonitrile in 8 mM sodium acetate buffer at a flow rate of 0.5 mL/min with a column temperature gradient of 5–45 °C (details are given in the SI Appendix, Table S6). Elution times for damage products are shown in SI Appendix, Fig. S5 and Table S7. The nucleoside-containing fractions were dried under vacuum, redissolved in 45 μL water, and analyzed by LC-MS/MS.

LC-MS/MS Detection of DNA and RNA products. All measurements other than of Sp and Gh were performed using an Agilent 1100 series HPLC system interfaced with an API3000 tandem quadrupole MS (AB Scieix) with a turbine ion spray source. Based on our published (25) and recently developed protocols, DNA and RNA samples were resolved on a Hypersil Gold aQ C18 reverse-phase column (150 × 2.1 mm, 3 μm particle size; Thermo Fisher) using 0.1% acetic acid in water (A) and 0.1% acetic acid in acetonitrile (B) with chromatographic parameters as specified in SI Appendix, Tables S5 and S7. The MS was operated in positive ion mode with the first and third quadrupoles fixed at 37 °C overnight. Enzymes were subsequently removed by microfiltration (Nanosep K10 Omega, Pall) and the filtrates concentrated under vacuum.

Gene Expression. Analyses were performed using customized qPCR arrays (Qiagen SABiosciences), which included 95 DNA repair and stress response genes (SI Appendix, Table S8) on an ABI 7900HT qPCR system (384-well block; Life Technologies). All steps were performed according to the manufacturer instructions with the following specifications: RNA quality and quantity was verified using a RNA 6000 Nano LabChip on an Agilent 2100 Bioanalyzer. Reverse transcription (RT) (1 μg/mL RNA) was performed with 400 ng of total RNA. Genomic DNA and reverse transcription controls were included in each PCR run to check for DNA contamination and impurities in RNA samples that could affect reverse transcription, respectively. Ct values were normalized to four different control genes (Actb, Gapdh, Hprt1, and 18SrRNA) for comparison of control and infected samples. For comparison of gene expression of liver and colon, samples C values were normalized to Gapdh, Hprt1, and 18SrRNA, because Actb showed significant inter tissue variation. Data were analyzed using the PCR Array Data Analysis Web portal (http://www.sabiosciences.com/pcrarraydataanalysis.php) according to the ΔΔCt method.

Statistical Analysis. Statistical analyses were performed using Graph Pad Prism software (GraphPad Software). Ct/s scores were compared using Mann—Whitney U-test for non-parametric data. Because data of adduct levels were not normally distributed, non-parametric Mann—Whitney rank sum test was performed to assess differences between two groups. The Wilcoxon signed-rank test was used for statistical analysis when values in control groups were consistently below the limit of quantification or showed no variance. Correlation analyses were performed using Spearman’s rank order correlation analysis. Analysis of gene expression data was performed using two-tailed Student’s t-test, with P < 0.05 considered statistically significant. SIMCA-P+ (Umetrics, Kinnelon) was used for multivariate data analysis.

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