Accumulation of abasic sites induces genomic instability in normal human gastric epithelial cells during *Helicobacter pylori* infection

D Kidane¹, DL Murphy² and JB Sweasy²

*Helicobacter pylori* infection of the human stomach is associated with inflammation that leads to the release of reactive oxygen and nitrogen species (RONs), eliciting DNA damage in host cells. Unrepaired DNA damage leads to genomic instability that is associated with cancer. Base excision repair (BER) is critical to maintain genomic stability during RONs-induced DNA damage, but little is known about its role in processing DNA damage associated with *H. pylori* infection of normal gastric epithelial cells. Here, we show that upon *H. pylori* infection, abasic (AP) sites accumulate and lead to increased levels of double-stranded DNA breaks (DSBs). In contrast, downregulation of the OGG1 DNA glycosylase decreases the levels of both AP sites and DSBs during *H. pylori* infection. Processing of AP sites during different phases of the cell cycle leads to an elevation in the levels of DSBs. Therefore, the induction of oxidative DNA damage by *H. pylori* and subsequent processing by BER in normal gastric epithelial cells has the potential to lead to genomic instability that may have a role in the development of gastric cancer. Our results are consistent with the interpretation that precise coordination of BER processing of DNA damage is critical for the maintenance of genomic stability.

Oncogenesis (2014) 3, e128; doi:10.1038/oncsis.2014.42; published online 24 November 2014

INTRODUCTION

*Helicobacter pylori* colonizes the gastric mucosa of half of the world's population and is a major etiopathogenic factor for chronic antral gastritis, duodenal ulcers and gastric cancer. Chronic inflammation associated with the long-term persistence of *H. pylori* infection leads to release of reactive oxygen and nitrogen species (RONs) from inflammatory cells. RONs can cause DNA base damage, strand breaks and damage to the tumor-suppressor genes and enhanced expression of proto-oncogenes.

Nitric oxide has also been found to inhibit the function of 8-oxoguanine glycosylase (OGG1) to impair the removal of DNA lesions that likely contribute to carcinogenesis. In addition, bacterial products including cytotoxins, lipase, phospholipase or the urease-mediated release of toxic ammonia at the site of inflammation can damage DNA, which may represent an early step in gastric carcinogenesis via development of chromosomal aberrations and DNA mutations. *H. pylori* also induces hypoxia-inducible factor-1α that in turn inhibits mismatch repair to promote infection-associated microsatellite instability and cancer.

The cellular consequences of DNA oxidation by RONs can lead to a number of different types of damage, such as 7, 8-hydroxy-2′-deoxyguanosine (8oxodG), abasic sites (AP) and oxidized deoxyribose sugars, which in turn lead to single-stranded DNA (ssDNA) breaks and double-stranded DNA breaks (DSBs), crosslinking of DNA and mutation. The most common oxidative base modifications resulting from direct attacks by hydroxyl radicals are purine lesions (8oxodG and 8-oxoA) and pyrimidine lesions (thymine glycol and cystosine glycol) in the human body in association with human cancer. Thousands of these lesions may be formed in each cell daily and levels are increased upon exposure to a variety of environmental factors. Oxidized bases, including 8oxodG, are removed predominantly by base excision repair (BER). BER is the major repair pathway of DNA damage induced by RONs and is critical for maintaining genome stability during chronic inflammation that occurs during bacterial infection. BER is initiated by DNA glycosylases that recognize and cleave the damaged bases. The OGG1 bifunctional DNA glycosylase is the major enzyme that catalyzes the removal of 8oxodG paired with C. OGG1 remains bound to its abasic site product and its turnover can be stimulated either by APendonuclease 1 (APE1) or by NEIL1, both of which can process the AP site. After AP site processing and end-remodeling, the single-nucleotide gap is filled by DNA polymerase beta and the nick is sealed to complete repair.

In humans, defective removal of 8oxodG or other types of base lesions may increase susceptibility to *H. pylori*-induced cancer due to mutagenesis. For example, unrepaired 8oxodG mismatches with adenine and results in G:C to A:T transversion mutations. Importantly, every step of BER generates intermediates (AP sites, 5′-deoxyribose phosphate residues and ssDNA breaks), which have been shown to be both mutagenic and toxic to cells. An imbalance between the generation of excess AP sites and inefficient repair has the potential to lead to mutation or DNA replication fork collapse.

Some studies showed that *H. pylori* infection in BER-deficient cells leads to increased levels of inflammation that consequently result in the production of more RONs and tumor-promoting cytokines. Furthermore, Toller et al. found that coculture of *H. pylori* with mouse and human gastric cancer cell lines led to...
increased levels of DSBs. Although *H. pylori* infection induces genomic instability, the underlying mechanism is not clear. Because *H. pylori* induces oxidative base damage, we hypothesized that the processing of this small base damage will lead to the accumulation of BER intermediates. Our results show that *H. pylori* infection significantly increases the number of AP sites in cells. These AP sites arise in replicating DNA and eventually lead to DSB formation. In contrast, downregulation of the OGG1 DNA glycosylase reduces the number of AP sites and DSBs during infection. These data suggest that OGG1 deficiency has a protective role against genomic instability induced by *H. pylori* infection. Our data support the conclusion that *H. pylori* infection induces the accumulation of AP sites in DNA that are further processed into DSBs, resulting in genomic instability and cellular transformation.

**RESULTS**

*H. pylori* induces accumulation of AP sites

To determine whether *H. pylori* infection induces an increased number of AP sites compared with non-infected controls, we infected GES-1 immortal but non-transformed gastric epithelial cells with *H. pylori* at a multiplicity of infection (MOI) of 100 bacteria to 1 cell. We then measured the number of AP sites in DNA extracted from the cells following infection at different time intervals as previously described (Figures 1a and b). For these

![Graph](attachment:image1.png)

**Figure 1.** (a–e) Global levels of AP site accumulation are enhanced in *H. pylori*-infected gastric epithelial cells. (a) A standard curve for estimating AP sites/10^5 nucleotides using the AP sites assay kit (see Materials and methods section). (b) Estimated numbers of AP sites/10^5 nucleotides before (unfilled columns) and after 3, 6, 12, 24, 48, and 72 h of *H. pylori* infection of GES-1 cells (filled columns) and after 1 h treatment of GES-1 cells with 10mM H_2O_2 as a control. (c) There is no fold change difference observed in protein expression levels of APE1 and DNA polymerase beta in *H. pylori*-infected versus non-infected control. We have applied the ImageJ program to measure the integrated density of each band and normalized with the loading control. We then calculated the ratio of adjusted integrated density of the sample from the infected versus non-infected group as the fold change. (d) Representative images of the lengths of DNA tracks before and after APE1 treatment of *H. pylori*-infected GES-1 cells. (e) The distribution of the lengths of the DNA tracks (μm) in cells treated with APE1 after *H. pylori* infection. ns, not significant.
experiments, AP sites were measured using an AP site assay kit (Colorimetric; Abcam, Cambridge, MA, USA) that utilizes an aldehyde reactive probe (ARP) reagent that reacts specifically with an aldehyde group, which is the open ring form of an AP site. We found that *H. pylori* infection generates on average 16 AP sites/10⁵ nucleotides within 24 h compared with ~4 AP sites/10⁵ nucleotides in non-infected controls (P = 0.0001), a fourfold difference. Strikingly, the numbers of AP sites remain high over several hours in cells infected with *H. pylori*, reaching a maximum level of ~22 AP sites/10⁵ nucleotides within the first 3 h of infection, and this is not significantly reduced during the next consecutive 6–48 h of infection (Figure 1b). Remarkably, *H. pylori* infection induces on average a threefold increase in the level of AP sites compared with cells treated with 10 mM H₂O₂, a potent base-oxidizing agent (P = 0.01).

We wished to determine whether AP sites could arise in newly synthesized DNA in cells infected with *H. pylori*. To determine whether this is the case, we infected GES-1 cells with *H. pylori* for 12 h and then added iodo-deoxyuridine (IdU) to label the DNA during the active infection. We then lysed the cells, spread the DNA fibers on glass slides, treated them with APE1 followed by fixation and measured the lengths of the DNA fibers (Figure 1d). Fixation after treatment with APE1 allows the measurement to work. The lengths of the DNA tracks are significantly decreased in APE1-treated DNA fibers derived from *H. pylori*-infected cells (P = 0.001) versus infected cells without APE1 treatment (Figure 1e), confirming that DNA track length in these experiments is dependent upon lesion processing by addition of APE1 to the fibers on the slide. Importantly, we found that the average length of the DNA tracks in APE1-treated DNA fibers derived from non-infected cells is significantly higher than the length of APE1-treated DNA fibers from infected cells (11.8 μm versus 6.4 μm; P = 0.0001; Figure 1e), suggesting that more AP sites arise in actively replicating DNA from infected cells. However, protein expression of APE1 and DNA polymerase beta were not altered after 12 and 24 h of *H. pylori* infection (Figure 1c).

OGG1 downregulation reduces the number of AP sites during *H. pylori* infection

The OGG1 DNA glycosylase removes 8oxodG from DNA to initiate BER, resulting in AP sites. The increase in the number of AP sites observed in GES-1 cells infected with *H. pylori* could be a result of deficient processing by APE1 after excision of the damaged base by OGG1. To determine whether AP site levels observed in infected cells resulted from the removal of damaged bases by OGG1, we used small interfering RNA (siRNA) to downregulate OGG1 in infected GES-1 cells and controls, and then quantified the

Figure 2. OGG1 downregulation suppresses AP site accumulation. (a) Western blot analysis of siRNA mediated downregulation of OGG1, GAPDH (control) and immunoblot with anti-rabbit OGG1 antibody, anti-mouse GAPDH and the loading control, anti-alpha-tubulin. (b) The estimated numbers of AP sites in OGG1 and GAPDH downregulated cells before (unfilled columns) and after (filled columns) infection with for 3, 6, 12, 24, 48 and 72 h. Note that all the experiments were conducted after 72 h of downregulation of OGG1 and GAPDH genes. (c) Representative images of DNA fibers after OGG1 and GAPDH downregulation. Cells were infected with *H. pylori* for 12 h and labeled with IdU for 2 h after *H. pylori* infection. The cells were lysed and the DNA fibers were spread on slides and treated with APE1 for 1 h at 37 °C. (d) Average lengths of DNA tracks in OGG1 and GAPDH downregulated cells with or without APE1 treatment. ns, not significant.
numbers of AP sites in these cells (Figures 2a and b). Down-regulation of OGG1 protects the cells from the accumulation of high levels of AP sites (Figure 2b) compared with controls that are downregulated for GADPH. We found that \textit{H. pylori} infection generates on average 14 AP sites/10^5 nucleotides within 24 h compared with ~4 AP sites/10^5 nucleotides in infected but OGG1-downregulated cells (\(P=0.02\); Figure 2b). Remarkably, \textit{H. pylori} infection induces on average a threefold lower number of AP sites in OGG1-downregulated cells compared with control cells (\(P=0.01\)). Next, we wished to determine the levels of AP sites in newly synthesized DNA in cells that were downregulated for OGG1. We performed these experiments by treating cells with an

**Figure 3.** \textit{H. pylori} infection of normal human gastric epithelial cells induces DSBs. (a) Images of GES-1 cells that were infected with J99 strain of live \textit{H. pylori}, dead bacterial extract or a supernatant of the bacterial cell culture for 24 h. Note that blue represents the nuclei of cells stained with 4',6-diamidino-2-phenylindole and red represents \(\gamma\)H2AX foci. (b) The percent of \(\gamma\)H2AX-positive cells, meaning >10 foci/cell, was estimated from live bacteria-infected GES-1 cells (\(n=50\)), bacterial cell extract (\(n=46\)), supernatant (\(n=60\)) and non-infected (\(n=70\)). (c) Western blot analysis of \(\gamma\)H2AX before and after 12 and 24 h of \textit{H. pylori} infection. The level of \(\gamma\)H2AX increased seven- and eightfold after infection with \textit{H. pylori} for 12 and 24 h, respectively. Note that quantification of bands was carried out using the National Institutes of Health ImageJ program by measuring the integrative density of each \(\gamma\)H2AX band that is normalized with alpha-tubulin band as internal control. (d) Estimated percentage of cells with \(\gamma\)H2AX foci in different cell cycle stages based on FACS analysis after 24 h of infection. Cells were stained with \(\gamma\)H2AX antibody and propidium iodide to assess the levels of DSBs and the cell cycle phase, respectively, and were analyzed by flow cytometry. (e) Percentage of cells with \(\gamma\)H2AX foci in OGG1-deficient cells after 24 h of infection. (f) Percentage of cells with \(\gamma\)H2AX in different stages of the cell cycle in OGG1-deficient versus OGG1-proficient cells after 24 h of infection based on FACS analysis. Note that all images were taken ×63 Zeiss microscope with constant exposure time and any cell with >10 foci of \(\gamma\)H2AX/cell was categorized as a positive. The data were analyzed based on the paired \(t\)-test using GraphPad Prism program. ns, not significant.
siRNA against OGG1 and infected GES-1 cells with *H. pylori* for 12 h. We then labeled replicating DNA for 2 h in cells with an active *H. pylori* infection, lysed the cells, spread the DNA fibers on slides, treated with APE1 and fixed the cells. We then quantified the lengths of the DNA tracks (Figure 2c). For non-infected GES-1 cells, we compared control cells (GAPDH downregulated) versus OGG1-downregulated cells treated with APE1 (10 μM) in infected GES-1 cells that are downregulated for GAPDH (3.9 μM; *P* = 0.0001; Figure 2d). These data suggest that the downregulation of OGG1 results in fewer AP sites that serve as a substrate for APE1. Our results are consistent with the idea that removal of damaged bases by OGG1 leads to the accumulation of DSBs during the G1 phase of the cell cycle,62,63 which could be occurring in the cell repair,64 we performed co-immunostaining of the cells with antisera against PCNA and γH2AX. This revealed that 45% of the γH2AX foci colocalized with PCNA in infected cells (*P* = 0.001, *n* = 78; Figures 4a and b) and colocalization was observed in very few uninfected cells (*n* = 92; Figure 4a). Colocalization of PCNA with γH2AX is likely indicative of DNA damage within cells positioned in S phase.

Replication fork arrest is a source of genomic rearrangements, and the recombinogenic properties of blocked forks likely depend on the cause of blockage. We applied a DNA fiber approach to determine whether infection with *H. pylori* leads to slowed or stalled replication forks. The general schematic for these experiments is shown in Figure 5a. We labeled actively replicating DNA with IdU and then infected cells with *H. pylori* for 12 h, followed by incubation with CldU, as shown in the general schematic in Figure 5a. The replication fork speed after infection was quantified by dividing the length of each fluorescent track by the time of incubation with the halogenated nucleotide, shown in Figures 5b and c. Analyses of replication fork speeds in *H. pylori*-infected versus non-infected cells showed a broad distribution of values ranging from 0.085 to 1.24 kb/min and 0.4 to 2.5 kb/min, respectively, (Figure 5c). Approximately 58% of the forks in infected cells exhibit fork speeds between 0.2 and 0.6 kb/min. In contrast, the distribution of fork speeds in non-infected cells is between 0.6 and 1.2 kb/min (Figure 5c). The average fork speed with infection (mean ± s.e.m.) is 0.42 ± 0.03 kb/min, which is significantly reduced compared with that of non-infected control groups (1.05 ± 0.048 kb/min; *P* = 0.0001; Figure 5d). Furthermore, we find that *H. pylori* infection induces stalled forks in 40% of cells (*n* = 263) versus ~6% of non-infected control cells (*n* = 201; *P* = 0.0001)
Replication protein A-coated ssDNA and appearance of Rad51 increases during infection

Replication protein A (RPA), the major ssDNA-binding protein in eukaryotic cells, accumulates along stretches of ssDNA generated by stalled replication forks and/or DNA damage.65,66 To determine whether ssDNA levels increase during infection, we quantified the number of cells with RPA foci (Figures 6a and b). We find that the number of H. pylori-infected cells with RPA foci significantly increased (n = 143, 63%) compared with non-infected controls (n = 176, 18%; P = 0.001; Figure 6b). Once DSBs are induced by H. pylori,5′-DNA end resection of DSBs is a prerequisite for loading of Rad51 to promote strand exchange activity.67,68 When the cells were analyzed for the presence of RAD51 foci (Figure 6c), 50% of
the infected cells exhibited foci compared with 13% for non-infected controls (13%; n = 67; P = 0.0001; Figure 6d).

H. pylori infection induces chromosomal aberrations
DSBs can lead to chromosomal fragmentation and genomic rearrangements if not repaired in an accurate and timely manner.61 Previous work has documented the existence of chromosomal aberrations but only in gastric cancer cells infected with H. pylori. However, it is not known whether H. pylori infection induces chromosomal aberrations in normal gastric epithelial GES-1 cells, which are the cells that are initially infected with this pathogen. Therefore, we asked whether H. pylori infection induces chromosomal aberrations in normal human gastric epithelial cells. Infection of GES-1 cells with H. pylori for 24 h results in significantly increased levels of chromosomal fusions (64%), chromosomal fragments (48%), chromatid breaks (36%) and chromosome breaks (n = 21%) versus non-infected cells (P = 0.0001; Figures 7a and b). Our observation is significant because the levels of chromosomal aberrations are more pronounced in normal gastric epithelial cells than in previously reported gastric cancer cell lines infected with H. pylori.53 We tested the hypothesis that the genomic instability resulting from H. pylori infection induces cellular transformation. We infected normal gastric epithelial cells with H. pylori for 35 days and monitored focus formation. In the focus formation assay, non-transformed cells will grow to confluence forming a monolayer (Figure 7c), whereas transformed cells will continue to grow after reaching confluence, thereby forming foci (Figure 7c). We find that the number of foci/field are significantly increased in infected cells versus non-infected control cells (Figure 7d; P = 0.0001).

DISCUSSION
The response of gastric epithelial cells to H. pylori infection and their ability to maintain genomic stability via DNA repair mechanisms is essential in preventing gastric cancer initiation and progression. Here, we provide important mechanistic insight regarding the consequences of H. pylori infection of normal human gastric epithelial cells, a physiologically relevant model of gastric cancer. We show that infection of these cells with H. pylori induces the accumulation of AP sites. Our results show that AP sites are generated at a high frequency in genomic DNA during infection. In contrast, downregulation of OGG1 during H. pylori infection protects cells from accumulating AP sites. Our data suggest that H. pylori infection induces 8oxodG lesions in genomic DNA that are removed by OGG1, leading to the accumulation of AP sites, either as a result of continuous generation, deficient repair or an imbalance between generation of AP sites and their repair. Moreover, we show that many of these AP sites are further processed into DSBs, resulting in genomic instability in OGG1-proficient H. pylori-infected cells. We conclude that H. pylori infection induces BER intermediates that likely overwhelm the BER machinery. This alters the dynamics of AP sites in the DNA to drive genomic instability in gastric epithelial cells.

Our results show that processing of AP sites into DSBs occurs during all phases of the cell cycle in OGG1-proficient cells. The cell cycle profiles revealed an increase in the percentage of cells in G1 versus S and G2 after infection. (Supplementary Figure 1) Therefore, we propose that DNA DSBs accumulate as a result of AP site processing during infection by two different mechanisms. The first mechanism is associated with the G1 phase of the cell cycle in which we observe the most significant accumulation of DSBs
We suggest that these DSBs arise as a result of the removal of clustered damaged bases by OGG1. OGG1 glycosylase activity could catalyze the removal of the damaged base, resulting in an AP site that is processed by the OGG1 lyase activity, leading to DSBs if the oxidative DNA damage is clustered. However, OGG1 has weak lyase activity, so once the damaged base is removed by this enzyme, it is possible that other bifunctional DNA glycosylases such as NTH1, NEIL1 or NEIL2, or perhaps APE1 itself, could process the abasic sites, leading to the accumulation of DSBs. Our results show that OGG1 deficiency suppresses the accumulation of AP sites, resulting in a significant reduction of DSBs at the G1 cell cycle stage during H. pylori infection. These data suggest that OGG1 functions in the removal of bases that are damaged during infection, and that an alternative DSB repair pathway may be operative in the presence of OGG1 deficiency.

(Figure 7e). We suggest that these DSBs arise as a result of the removal of clustered damaged bases by OGG1. OGG1 glycosylase activity could catalyze the removal of the damaged base, resulting in an AP site that is processed by the OGG1 lyase activity, leading to DSBs if the oxidative DNA damage is clustered. However, OGG1 has weak lyase activity, so once the damaged base is removed by this enzyme, it is possible that other bifunctional DNA glycosylases such as NTH1, NEIL1 or NEIL2, or perhaps APE1 itself, could process the abasic sites, leading to the accumulation of DSBs. Our results show that OGG1 deficiency suppresses the accumulation of AP sites, resulting in a significant reduction of DSBs at the G1 cell cycle stage during H. pylori infection. These data suggest that OGG1 functions in the removal of bases that are damaged during infection, and that an alternative DSB repair pathway may be operative in the presence of OGG1 deficiency.
OGG1 deficiency protects cells from H. pylori-induced DSBs at the G1 cell cycle stage. The presence of other bifunctional glycosylases, such as NEIL1, do not appear to have a significant backup role in the removal of 8oxodG induced by H. pylori likely as a result of their weak glycosylase activity against 8oxodG. However, the lyase activity of glycosylases that recognize other types of base lesions may process AP sites owing to robust AP lyase activity and may be able to promote BER in OGG1-deficient cells infected with H. pylori. This could contribute to the lower numbers of AP sites in OGG1-deficient cells infected with H. pylori. Our observation of the critical role of OGG1 during H. pylori infection is supported by previously published data, showing that OGG1 deficiency promotes a protective role against inflammation and genotoxicity associated with H. pylori infection.73–77 Our results are consistent with the interpretation that OGG1 has a key role in the removal of RONs-mediated 8oxodG lesions during H. pylori infection. Initiation of BER of clustered oxidized bases on opposite strands of the DNA by OGG1 would lead directly to the formation of DSBs. These resulting DSBs are most likely repaired by the error-prone pathway of non-homologous end-joining during G1, leading to chromosomal aberrations.

The second mechanism of DSB generation occurs during S and G2 phases, as DSBs also accumulate during these phases of the cell cycle. Our results suggest that DSBs arise as a result of replication fork encounter of AP sites that may potentially block G2 phases, as DSBs also accumulate during these phases of the prone pathway of non-homologous end-joining during G1, of DSBs. These resulting DSBs are most likely repaired by the error-prone pathway of non-homologous end-joining during G1, leading to chromosomal aberrations.

Altogether, our data indicate that H. pylori infection induces the accumulation of un repaired BER intermediates that can initiate a cascade of events to generate DSBs. We show that overwhelming the capacity of BER as a result of oxidative DNA damage induced by H. pylori results in the accumulation of AP sites and DSBs in normal gastric epithelial cells, leading to genomic instability, an underlying cause of gastric cancer. Our findings suggest that alteration of the activity of enzymes that function in BER as a result of genetic mutation could significantly impact gastric cancer risk.

MATERIALS AND METHODS

H. pylori strains, culture and preparation of bacterial components

The H. pylori type strain, J99 strain ATCC 700824, was used in this study and was stored at −80°C. The H. pylori strain was inoculated onto Columbia blood agar plates, which consisted of 39 g of Columbia Agar Base (CM0331, Becton Dickinson, Franklin Lakes, NJ, USA) per liter, 7% hemolyzed horse blood (SR0048) and supplemented with one vial of Dent (SR0147, Oxoid, Hampshire, UK). For growing visible colonies, plates were kept in a GasPak (BD GasPak EZ products, Franklin Lakes, NJ, USA) at 37°C from 3 to 10 days until colonies were observed. From the primary growth, single colonies were propagated in blood agar for an additional 48 h, harvested in phosphate-buffered saline (PBS) and inoculated into Hams F12 media supplemented with 5% horse serum and incubated in 5% CO2 incubator. Bacterial protein extraction was conducted using B-PER extraction reagents based on manufacturer’s protocols (Thermo Scientific, Rockford, IL, USA).

Cell lines and cultures

The GES-1 gastric epithelial cell line was obtained from the Chinese Academy of Sciences. GES-1 cells are derived from a human gastric mucosa epithelium and immortalized via SV40 and are non-tumorigenic.86,87 GES-1 cells were maintained in RPMI supplemented with 10% fetal bovine serum, 1% glutamate and 1% penicillin-streptomycin.

Preparation of GES-1 cellular extracts

The cells were suspended in 20 mM HEPES, pH 7.9, 10 mM NaCl, 1.5 mM MgCl2, 10% glycerol 1 mM phenylmethylsulfonyl fluoride (PMSF) and a protease inhibitor cocktail and incubated for 1 h on a rocking platform at 4°C. The whole-cell lysate was collected after centrifugation at 13 000 r.p.m. at 4°C for 20 min and aliquoted and stored −80°C.

Confocal microscopy of nuclear protein localization, antibodies used

For γH2AX foci staining, cells were grown on four chamber slides and infected with H. pylori for 24 h (100:1 bacteria to host cell ratio, MOI=100). After cell media was removed and rinsed with PBS, cells were fixed with methanolic acid (3:1 ratio), incubated for 15 min at −20°C, and permeabilized in PBS containing 0.5% Triton X-100 for 8 min at room temperature (RT). Cells were then incubated with 1:200 diluted rabbit polyclonal anti-γH2AX antibody (Bethyl Laboratory, Montgomery, TX, USA) for 1 h at RT and detected with a secondary fluorescein isothiocyanate (FITC)-conjugated goat anti-rabbit IgG (Jackson Immunoresearch Laboratories, West Grove, PA, USA). Antibody dilutions and washes after incubations were performed in PBS containing 0.5% bovine serum albumin and 0.05% Tween-20. Finally, coverslips were mounted in VECTASHIELD mounting medium with 4’,6-diamidino-2-phenylindole (H-1500; Vector Laboratories, Burlingame, CA, USA).

Flow cytometry

GES-1 cells were cultured in four chamber slides and infected with H. pylori (100:1 bacteria to host cell ratio, MOI=100) for 24 h. The bacterial media was removed and the cells were rinsed with PBS then, cells were harvested by trypsinization, washed once with PBS and pelleted. The pellet was resuspended by adding 70% ice-cold ethanol dropwise while vortexing. Cells were fixed overnight at −20°C. The cells were incubated with primary phospho-γH2AX antibody (Millipore, Billerica, MA, USA; 05-636) 1:500 overnight at 4°C. Following the incubation, cells were washed twice with PBS and incubated with anti-mouse secondary antibody conjugated to FITC 1:500 at 1 h at RT. Cells were washed twice with PBS and resuspended in 500 μl propidium iodide (PI)/RNase staining buffer (BD Pharmingen, Franklin Lakes, NJ, USA). Fluorescence was analyzed by flow cytometry using the BD FACS calibur (Franklin Lakes, NJ, USA) and also by using Flowjo 8.8.6 software (Ashland, OR, USA).

Metaphase spread preparation

For preparation of metaphase chromosome spreads in the absence of infection, GES-1 cells were treated with colcemid (final concentration of 0.1 μg/ml) for 6 h before harvesting. In contrast, GES-1 cells were cultured and infected with H. pylori (100:1 bacteria to host cell ratio, MOI=100) for 24 h including colcemid treatment for the last 6 h. The cells were trypsinized and were washed once with PBS. Mitotic cells were collected and centrifuged at 200 g for 5 min at RT, and the harvested cells were treated with 75 mM KCl at 30 min at 37°C. After centrifugation, the cells were fixed in a mixture of 3:1 ethanol:acetic acid. Ten microliters of cell suspension were dropped onto slides and allowed to dry, followed by rinsing the slides in PBS and staining with 5% Giemsa stain for 8 min. The slides were rinsed with water and were air-dried. Images were acquired with a Zeiss microscope (Peabody, MA, USA).

DNA fiber analysis

For DNA replication analysis, sequential labeling of DNA with IDU and 5-chloro-2'-deoxyuridine (CldU) were performed. A subconfluent, asynchronous population of GES-1 cells was first labeled for 30 min with 25 μM IDU, washed with medium three times and infected for 12 h with H. pylori at an MOI of 100:1 (bacteria to cell ratio; MOI = 100). The cells were then labeled for another 30 min with 250 μM CldU. After incubation, cells were treated with 75 mM KCl at 30 min at 37°C. After centrifugation, the cells were fixed in a mixture of 3:1 ethanol:acetic acid. Ten microliters of cell suspension were dropped onto slides and allowed to dry, followed by trypsinizing the slides in PBS and staining with 5% Giemsa stain for 8 min. The slides were rinsed with water and were air-dried. Images were acquired with a Zeiss microscope (Peabody, MA, USA).
Ham’s F12 media and then pulsed with CDU label for 30 min, and the cells were harvested for the fiber assay. For immunofluorescence staining, the slides were fixed for 10 min with methanol-acetic acid (3:1) and air-dried. The slides were treated with 2.5 % HCl for 30 min, washed with 1X PBS three times and then blocked with 3% bovine serum albumin/PBS for 1 h. CDU was detected by incubating acid-treated fiber spread with rat anti-BrdU monoclonal antibody (Abcam) and CDU was detected using mouse anti-BrdU monoclonal antibody (1:1000; Becton Dickenson) for 1 h at RT. followed by washing three times with 1X PBS and stained with secondary antibody conjugated with sheep anti-mouse Cy3 and goat anti-rat Alexa flour 488 for 1 h at RT. The slides were mounted with Vectashield mounting media and covered with coverslips. Images were acquired with ×63 magnification using a Zeiss microscope and processed and analyzed using the ImageJ program (NIH, Bethesda, MD, USA). The lengths of red (Cy3) or green (AF 488) labeled patches were measured using the ImageJ software (National Institutes of Health; http://rsweb.nih.gov/ij/) and arbitrary length values were converted into micrometers using the scale bars created by the microscope. Fluorescence images were captured using a Zeiss LSM 510 inverted confocal microscope using ×63/numerical aperture 1.4 oil immersion objective, and data analysis was carried on using the ImageJ software. We applied a conversion factor used is 1 μm = 2.59 kb.85

AP site distribution on DNA tracks

A subconfluent, asynchronous population of GE1 cells was infected for 12 h with H. pylori (MOI = 100) then labeled for 2 h with 25 μM IDU. Cells were harvested and placed onto a glass slide and allowed to gently adhere to the glass surface. Next the cells were lysed causing their genomic DNA to be released, and then the DNA is straightened and aligned on the slide using gravitational flow. After air-drying, the slides were treated with APE1 (50 and 100 U) with reaction buffer containing 1 × NEB4 buffer for 1 h at 37 °C. The slides were washed with 1 × PBS three times, and fixed for 10 min with methanol-acetic acid (3:1) and processed for immunofluorescence based on the above described methods. To determine the relative distribution of AP site formation on the length of DNA track, we treated the slides with APE1 (100 U) for 1 h at 37 °C before fixation, then fixed the cells and labeled with secondary antibody conjugated with sheep anti-mouse Cy3. The slides were mounted with Vectashield mounting media and covered with coverslips. Images were acquired with ×63 magnification using a Zeiss microscope, and were processed and analyzed using the ImageJ program.

AP site analysis

GE1 cells were cultured until 70% confluent and infected with H. pylori (with 100:1 bacteria to host cell ratio, MOI = 100), for different time intervals. The bacterial media was removed and the cells were washed three times with PBS, then genomic DNA was isolated using DNAzol genomic DNA isolation reagents that contains guanidine/thiocyanate buffers that contain guanidine/detergent lysis buffer that we purchased from InVitrogen (Grand Island, NY, USA). The AP assay was conducted after the DNA was labeled with ARP. The AP assay kit (Colorimetric; Abcam) utilizes the ARP reagent that reacts specifically with an aldehyde group, which is the open ring form of DNA intercalating agent. After treatment of mRNA containing AP sites with ARP reagents, AP sites are tagged with biotin residues, which can be quantified using an avidin–biotin assay followed by a colorimetric detection. The kit provides the necessary reagents for convenient determination of abasic sites in purified DNA in 96-well plate format. The number of AP sites was measured and calculated based upon a standard curve generated using ARP standard DNA solutions as described previously. DNA Damage AP sites assay kit, Colorimetric, (Abcam).

siRNA for OGG1 downregulation

For the siRNA studies, we use SMARTpool siRNA against OGG1 with target sequence 5′-GCCUUCUUCGUCAUUAGUG-3′ (A-005147-20), 5′-GCCUUCUGGACGACUGUCAU-3′ (A-005147-19), 5′-GGCUUCAAGGCGUCACCT-3′ (A-005147-18) and 5′-CUClCAGGAAUUCCACGGGU-3′ (A-005147-17) were used to downregulate OGG1. In addition, ON-TARGETplus GAPDH control siRNA with target sequence 5′-GUAACCGGAAUUGGCGUCAU-3′ were synthesized by Dharmacon Research Inc. (Lafayette, CO, USA). All DharmaFECT transfection reagent and protocols for downregulation of OGG1 and GAPDH were carried out based on modified Thermo Scientific DharmaFECT Transfection Reagents-siRNA Transfection Protocol.

Statistical analysis

All the reported data were evaluated in a pairwise manner, comparing infected versus non-infected control cells using GraphPad Prism (La Jolla, CA, USA).

Cellular transformation assay

Viable cells (200 cells/well) were seeded into wells of a six-well plate (in triplicate) and cultured in an incubator with 5% CO2. Then, bacteria were added in the ratio (GES-1 cells:bacterial cells) of 1:10 for coculture. GES-1 cells cultured under the same conditions but without coculture with H. pylori were used as controls. The culture continued for 35 days, and the cells were used for experiments and analysis during that period of time. Cells in the plate were washed with PBS after removal of the medium, and then fixed followed by staining with crystal violet for 15 min. The stain was washed off under running tap water, and the plate was allowed to dry. The number of distinctly stained cell growth pattern was counted under microscope.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

We thank Shanghai Jiao Tong University School of Medicine for generous gift of GE1 cells. We also thank Professor Raymond Monnat and Dr Julia Sidova for scientific advice. DK was supported by United States National Institutes of Health (NIH/National Cancer Institute (NCI)) K01 CA158450 and the work was also supported by CA 116753 (to JBS).

REFERENCES

1 Peek RM Jr, Blaser MJ. Helicobacter pylori and gastrointestinal tract adenocarcinomas. Nat Rev Cancer 2002; 2: 28–37.
2 Correa P. Human gastric carcinogenesis: a multistep and multifactorial process—First American Cancer Society Award Lecture on Cancer Epidemiology and Prevention. Cancer Res 1992; 52: 6735–6740.
3 Figureiredo C, Machado JC, Pharoah P, Sereca R, Sousa S, Carvalho R et al. Helicobacter pylori and interleukin 1 genotyping: an opportunity to identify high-risk individuals for gastric carcinoma. J Natl Cancer Inst 2002; 94: 1680–1687.
4 Cerutti PA. Oxidants and cancer. Lancet 1994; 344: 862–863.
5 Feig DL, Reid TM, Loeb LA. Reactive oxygen species in tumorigenesis. Cancer Res 1994; 54: 1890s–1894s.
6 Schreck RR. Tumor suppressor gene (Rb and p53) mutations in osteosarcoma. Pediatric Hematol Oncol 1992; 9: ix–i.
7 Bagchi D, Bhattarcharya G, Stohs SJ. Production of reactive oxygen species by gastric cells in association with Helicobacter pylori. Free Radic Res 1996; 24: 439–450.
8 Farinati F, Cardin R, Degan P, Rugge M, Mario FD, Bonvicini P et al. Oxidative DNA damage accumulation in gastric carcinogenesis. Gut 1998; 42: 351–356.
9 Izzotti A, De Flora S, Cartiglia C, Are BM, Longobardi M, Camoinaro A et al. Interplay between Helicobacter pylori and host gene polymorphisms in inducing oxidative DNA damage in the gastric mucosa. Carcinogenesis 2007; 28: 892–898.
10 Kim JJ, Tao H, Carlone E, Leung WK, Graham DY, Sepulveda AR. Helicobacter pylori impairs DNA mismatch repair in gastric epithelial cells. Gastroenterology 2002; 123: 542–553.
11 Loeb LA, Preston BD. Mutagenesis by apurinic/apyrimidinic sites. Annu Rev Genet 1986; 20: 201–230.
12 Obst B, Wagner S, Sewing KF, Bell W. Helicobacter pylori causes DNA damage in gastric epithelial cells. Carcinogenesis 2000; 21: 1111–1115.
13 Park DL, Park SH, Kim SH, Kim JW, Cho YK, Kim HJ et al. Effect of Helicobacter pylori infection on the expression of DNA mismatch repair protein. Helicobacter 2005; 10: 179–184.
14 Yao Y, Tao H, Park DI, Sepulveda JL, Sepulveda AR. Demonstration and characterization of mutations induced by Helicobacter pylori organisms in gastric epithelial cells. Helicobacter 2006; 11: 272–286.
15 Jaiswal M, LaRussso NF, Nishio N, Nakabeppu Y, Gores GJ. Human Ogg1, a protein involved in the repair of 8-oxoguanine, is inhibited by nitric oxide. Cancer Res 2001; 61: 6388–6393.
16 Jaiswal M, LaRussso NF, Shapiro RA, Billiar TR, Gores GJ. Nitric oxide-mediated inhibition of DNA repair potentiates oxidative DNA damage in cholangiocytic. Gastroenterology 2001; 120: 190–199.
12 Hill JW, Haza TK, Izumi T, Mitra S. Stimulation of human 8-oxoguanine-DNA glycosylase by AP-endonuclease: potential coordination of the initial steps in base excision repair. Nucleic Acids Res 2001; 29: 430–438.

71 Vidal AE, Hickson ID, Boitexs S, Radicella JP. Mechanism of stimulation of the DNA glycosylase activity of hOGG1 by the major human AP endonuclease: bypass of the AP lyase activity step. Nucleic Acids Res 2001; 29: 1285–1292.

72 Shibutani S, Takeshita M, Grollman AP. Translesional synthesis on DNA templates.

73 Dou H, Mitra S, Haza TK. Repair of oxidized bases in DNA bubble structures by human DNA glycosylases NEIL1 and NEIL2. J Biol Chem 2003; 278: 49679–49684.

74 Wiederhold L, Leppard JB, Kedar P, Karimi-Busheri F, Rasouli-Nia A, Weinfeld M et al. AP endonuclease-independent DNA base excision repair in human cells. Mol Cell 2004; 15: 209–220.

75 Dizdaroglu M, Karahalil B, Senturker S, Buckley TJ, Roldan-Arjona T. Excision of products of oxidative DNA base damage by human NTH1 protein. Biochemistry 1999; 38: 243–246.

76 Luna L, Bjoras M, Hoff E, Rognes T, Seeberg E. Cell-cycle regulation, intracellular sorting and induced overexpression of the human NTH1 DNA glycosylase involved in removal of formamidopyrimidine residues from DNA. Mutat Res 2000; 460: 95–104.

77 Touati E, Michel V, Thiberge JM, Ave P, Huerre M, Bourgade F et al. Deficiency in OGG1 protects against inflammation and mutagenic effects associated with H. pylori infection in mouse. Helicobacter 2006; 11: 494–505.

78 Higuchi K, Katayama T, Iwai S, Hidaka M, Horiiuchi T, Maki H. Fate of DNA replication fork encountering a single DNA lesion during oriC plasmid DNA replication in vitro. Genes Cells 2003; 8: 437–449.

79 Shibutani S, Takeshita M, Grollman AP. Translesional synthesis on DNA templates containing a single abasic site. A mechanistic study of the ‘A rule’. J Biol Chem 1997; 272: 13916–13922.

80 Swanson RL, Morey NJ, Doetsch PW, Jinks-Robertson S. Overlapping specificities of base excision repair, nucleotide excision repair, recombination, and translesion synthesis pathways for DNA base damage in Saccharomyces cerevisiae. Mol Cell Biol 1999; 19: 2929–2935.

81 Jeggo P, O’Neill P. The Greek Goddess, Artemis, reveals the secrets of her cleavage. DNA Repair 2002; 1: 771–777.

82 Letavayova L, Markova E, Hermanska K, Vickova V, Vlasakova D, Chovanec M et al. Relative contribution of homologous recombination and non-homologous end-joining to DNA double-strand break repair after oxidative stress in Saccharomyces cerevisiae. DNA Repair 2006; 5: 602–610.

83 Budzowska M, Kanaar R. Mechanisms of dealing with DNA damage-induced replication problems. Cell Biochem Biophys 2009; 53: 17–31.

84 Henry-Mowatt J, Jackson D, Masson JY, Johnson PA, Clements PM, Benson FE et al. XRCC3 and Rad51 modulate replication fork progression on damaged vertebrate chromosomes. Mol Cell 2003; 11: 1109–1117.

85 Wang BJ, Zhang ZQ, Ke Y. Conversion of cadherin isoforms in cultured human gastrointestinal carcinoma cells. World J Gastroenterol 2006; 12: 966–970.

86 Lian H, Cui J, Wang Y, Liu J, Wang J, Shen H et al. Downregulation of Rad51 participates in OTA-induced DNA double-strand breaks in GE-1 cells in vitro. Toxicol Lett 2014; 226: 214–221.

87 Shinmura K, Kasai H, Sasaki A, Sugimura H, Yokota J. 8-hydroxyguanine (7,8-dihydro-8-oxoguanine) DNA glycosylase and AP lyase activities of hOGG1 protein and their substrate specificity. Mutat Res 1997; 385: 75–82.

88 Arai K, Morishita K, Shinmura K, Kohno T, Kim SR, Nohmi T et al. Cloning of a human homolog of the yeast OGG1 gene that is involved in the repair of oxidative DNA damage. Oncogene 1997; 14: 2857–2861.

Supplementary Information accompanies this paper on the Oncogenesis website (http://www.nature.com/oncogenesis)