

ORIGINAL RESEARCH

ABSTRACT

DNA damage signalling as an anti-cancer barrier in gastric intestinal metaplasia

Vaidehi Krishnan (D), ^{1,2} Debbie Xiu En Lim (D), ¹ Phuong Mai Hoang (D), ¹ Supriya Srivastava (D), ^{3,4} Junichi Matsuo (D), ¹ Kie Kyon Huang (D), ² Feng Zhu (D), ⁴ Khek Yu Ho (D), ^{4,5} Jimmy Bok Yan So (D), ^{6,7} Christopher Khor (D), ⁸ Stephen Tsao (D), ⁹ Ming Teh (D), ³ Kwong Ming Fock (D), ¹⁰ Tiing Leong Ang (D), ¹⁰ Anand D Jeyasekharan (D), ¹ Patrick Tan (D), ^{2,7} Khay-Guan Yeoh (D), ^{4,5,7} Yoshiaki Ito (D), ^{1,7}

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For numbered affiliations see end of article.

Correspondence to

Professor Yoshiaki Ito, Cancer Science Institute of Singapore, National University of Singapore, Singapore 117599, Singapore; csiitoy@nus.edu.sg and Professor Khay-Guan Yeoh, Department of Gastroenterology and Hepatology, National University Health System, Singapore 119074, Singapore; mdcykg@nus.edu.sg

DXEL, PMH and SS contributed equally.

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Objective Intestinal metaplasia (IM) is a premalignant stage that poses a greater risk for subsequent gastric cancer (GC). However, factors regulating IM to GC progression remain unclear. Previously, activated DNA damage response (DDR) signalling factors were shown to engage tumour-suppressive networks in premalignant lesions. Here, we interrogate the relationship of DDR signalling to mutational accumulation in IM lesions. **Design** IM biopsies were procured from the gastric cancer epidemiology programme, an endoscopic surveillance programme where biopsies have been subjected to (epi) genomic characterisation. IM samples were classified as genome-stable or genome-unstable based on their mutational burden/somatic copy-number alteration (CNA) profiles. Samples were probed for DDR signalling and cell proliferation, using the markers yH2AX and MCM2, respectively. The expression of the gastric stem cell marker, CD44v9, was also assessed. Tissue microarrays representing the GC progression spectrum were included.

Results MCM2-positivity increased during GC progression, while γ H2AX-positivity showed modest increase from normal to gastritis and IM stages, with further increase in GC. γ H2AX levels correlated with the extent of chronic inflammation. Interestingly, genomestable IM lesions had higher γ H2AX levels underscoring a protective anti-cancer role for DDR signalling. In contrast, genome-unstable IM lesions with higher mutational burden/CNAs had lower γ H2AX levels, elevated CD44v9 expression and modest promoter hypermethylation of DNA repair genes *WRN*, *MLH1* and *RAD52*.

Conclusions Our data suggest that IM lesions with active DDR will likely experience a longer latency at the premalignant state until additional hits that override DDR signalling clonally expand and promote progression. These observations provide insights on the factors governing IM progression.

INTRODUCTION

Gastric cancer (GC) is the third leading cause of cancer-related deaths with high incidence in high-risk Asian regions.¹ The most common form of GC, the intestinal-type, is initiated as chronic inflammation which progresses sequentially as gland loss

Significance of this study

What is already known on this subject?

- Activated forms of DNA damage response (DDR) signalling factors accumulate in premalignant lesions of the lung, colon and breast and engage in downstream tumoursuppressive networks.
- Thus, a protective role for DDR signalling against cancer progression has been proposed. However, due to the difficulty in obtaining sufficient premalignant biopsy material for parallel immunohistochemistry and genomic studies, the relationship between DDR signalling to mutational burden/copy-number alterations (CNAs) remains unclear.
- Moreover, the applicability of the DDR paradigm to gastric cancer (GC) is unclear.

What are the new findings?

- MCM2 levels, indicative of cell proliferation gradually increased with GC progression, while γH2AX accumulation, suggestive of DDR signalling correlated with chronic inflammation and increased with GC progression.
- Importantly, an inverse correlation between DDR activation and mutational/CNAs was observed in precancerous intestinal metaplasia lesions.
- The gastric stem cell marker CD44v9 which is known to provide higher adaptation to inflammation-dependent reactive oxygen species accumulated in genome-unstable IM lesions.
- Modest promoter hypermethylation of the DNA repair factors WRN, MLH1 and RAD52 was also observed in genome-unstable samples.

or atrophic gastritis, intestinal metaplasia (IM), dysplasia and adenocarcinoma.² The principal initiator of intestinal-type GC (IGC) is recognised as *Helicobacter pylori* infection which fosters an environment of chronic inflammation and triggers (epi) genomic downstream cascades.^{3 4}



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Significance of this study

How might it impact on clinical practice in the foreseeable future?

- These data are among the first to directly correlate higher DDR signalling with lower propensity to accumulate genomic instability at the premalignant IM stage.
- We propose that the γH2AX/MCM2/CD44v9 profiling of larger IM cohorts with accompanying genomic characterisation will facilitate design of better risk stratification strategies for IM lesions.

IGC is proposed to be preceded by a series of precancerous events in a model of multistep carcinogenesis.² The first recognisable histological change is chronic inflammation which can be either accompanied by no gastric gland loss (chronic nonatrophic gastritis), or with gland loss (chronic atrophic gastritis). Atrophic gastritis is followed by IM in which normal gastric mucosa is replaced by glands having an intestinal morphology.²³ In recent studies, IM has been shown to arise by parietal cell loss and consequent metaplastic trans-differentiation of the chief cells into intestinal-like glandular structures.⁶ IM is classified as the small intestine/complete-type or colonic/incomplete type with the incomplete-type more likely to progress into adenocarcinoma.⁷ In particular, much attention has been given to the precancerous IM stage because it is considered a point-of-noreturn in H. pylori-infected patients even after bacterial eradication.⁸⁹ IM prevalence is greater in patients with relatives having GC, in smokers, increases with age and poses an estimated annual risk of 0.13% to 0.25% for GC per year.^{10 11}

Although IM is detected in nearly one out of four patients undergoing endoscopy, there is no clear consensus on how IM lesions should be stratified for future routine endoscopic surveillance. According to the international guideline MAPS (management of precancerous conditions and lesions of the stomach), patients with moderate-to-severe atrophic gastritis and marked IM in both the corpus and antrum together with dysplasia are recommended to undergo endoscopic surveillance.¹² ¹³ Serum pepsinogen levels, and classifications of gastritis known as the operative link on gastritis assessment (OLGA) and operative link on gastritis intestinal metaplasia (OLGIM) integrating both the severity and location of atrophy/IM, are proposed to assist risk stratification.¹² ¹⁴ As such, more studies are needed to elucidate the underlying biology of IM to GC transformation.

One potential mechanism that can regulate IM to GC progression is the DNA damage response (DDR) pathway. Cells have evolved sophisticated mechanisms to precisely detect, signal and repair DNA damage, and mount a tumour-suppressive response collectively called as the DDR.¹⁵ DDR includes a network of proteins that detect DNA lesions and activate downstream responses like DNA repair, cell cycle checkpoints, senescence or apoptosis and can be quantified experimentally by the levels of phosphorylated H2AX, ATM, CHK2 and p53. In landmark studies, DDR was found activated in precancerous lesions of the lung, colon, breast, skin and urinary bladder by preventing further propagation of such damaged cells inducing them to undergo senescence.¹⁶⁻¹⁹ However, the original studies were based on immunohistochemical examination of precancerous biopsies and contemporaneous data on the mutational status was unavailable. Thus, the role of DDR signalling in premalignant lesions especially pertaining to the accumulation of mutational/ chromosomal aberrations remains mostly unknown.

We used a resource of biopsy samples from patients enrolled in gastric cancer epidemiology programme (GCEP), an endoscopic surveillance programme with minimum clinical follow-up of 5 years to conduct IHC analysis. Our earlier comprehensive (epi)genomic characterisation revealed that IM samples had a low mutational burden (mutation rate - 2.6 and 6.9 mutations per Mb in IM and GC, respectively) and only <5% of all IM samples harboured known cancer driver mutations.²⁰ Similarly, somatic copy-number alterations (CNAs) were found in ~12.5% of IMs as against none in the normal samples. Notably, samples with greater CNAs had a higher propensity for progression in a multivariate analysis.²⁰

Here, we used H2AX phosphorylation at serine 139 (yH2AX) by the apical DDR kinases, ATM, ATR and DNAPK-cs, as a surrogate marker for DNA damage signalling.²¹ We developed quantitative immunohistochemistry IHC (qIHC) methods to reliably quantify DNA damage and cell proliferation through automated cell recognition and image analysis. We also integrated our immunohistochemical findings with previously reported (epi)genomic features for the same samples. Our results suggest that IM lesions with activated DNA damage signalling have lower accumulation of mutation/CNAs and thus may exhibit a different trajectory towards tumour progression as compared with lesions that have not activated their DDR pathway. Our studies also imply a potential role for CD44v9 in promoting greater reactive oxygen species (ROS) defence²² and in attenuating tumour-suppresive DDR signalling in genome-unstable IM lesions. These observations, in turn raise valuable insights for developing biomarkers for IM risk stratification.

METHODS

Clinical samples

- 1. Commercial tissue microarrays (TMAs) Formalin-fixed, paraffin-embedded (FFPE) stomach tissue from anonymised patient samples, spotted as TMAs (US Biomax, Rockville, Maryland) were used. The TMA, #ST1001 (US Biomax), spotted as cores with 1 mm diameter and 5 μ M thickness was used to compare MCM2 and γ H2AX across stomach tissues graded as normal, inflammation or atrophic gastritis with IM. The TMA #IC00011B (US Biomax), spotted as cores with 1.5 mm diameter and 5 μ M thickness was used to compare MCM2 and γ H2AX across stomach tissues grated as normal, inflammation or atrophic gastritis with IM. The TMA #IC00011B (US Biomax), spotted as cores with 1.5 mm diameter and 5 μ M thickness was used to compare MCM2 and γ H2AX across stomach tissues graded as gastritis with IM or intestinal-type gastric cancer (IGC). For both ST1001 and IC00011B TMAs, the cores which were insufficient were excluded from the final analysis.
- 2. Archival samples of gastritis To compare the relationship between chronic inflammation and DDR signalling, nine non-atrophic gastritis (NAG) biopsies from *H. pylori*negative gastritis subjects were retrieved from the archives of the Department of Pathology, National University Hospital, Singapore, following our Institutional Review Board regulations. Chronic inflammation of the gastritis tissue was categorised as negative and positive by certified pathologists (SS and MT).²³
- 3. Tissue biopsies from the GCEP cohort GCEP is a multicentre longitudinal study comprising of 2980 subjects who were of Chinese ethnicity, above the age of 50 and were suffering from dyspepsia, heartburn or had a family history of gastric cancer. These patients were followed-up for at least 5 years. The study was approved by the Domain Specific Review Board, a part of the National Healthcare group and Centralised Institutional Board of Singapore Health Services. Written informed consents were obtained from

all participants. Upper gastrointestinal endoscopy was performed by credentialed endoscopists and six biopsies were taken for histological examination and were classified as mild, moderate or marked for the degree of chronic gastritis, H. pylori infection, gastric atrophy and IM according to the updated Sydney system for classification and gastritis gradation.²³ IM diagnosis and extent were confirmed by pathological examination of Alcian Blue-stained serial sections. From this GCEP cohort, we selected two groups of patients; the first group comprised of normal, marked gastritis and marked IM (n=19) and a second group comprised of samples with IM and had been subjected to prior genomic characterisation (n=15). Normal gastric biopsies were those who were H. pylori serology-negative, free of early gastric neoplasm and any other cancers, without gastric atrophy, without IM, free of dysplasia or polyps throughout the period of surveillance endoscopies.

Immunofluorescence staining of FFPE sections, multispectral acquisition and image analysis

Detailed methods for qIHC staining, imaging and data analysis are provided in supplementary methods, online supplementary figure S1 and online supplementary table 1. Briefly, FFPE sections were deparaffinised, rehydrated and subjected to heat-based antigen retrieval. Following blocking, slides were stained using the Opal 4-colour Multiplex Kit (Perkin Elmer, # NEL810001KT) based on the tyramide signal amplification method. Immunofluorescence images were captured on the Vectra multiplex pathology system (Perkin Elmer) using the 20x objective lens. The inForm 2.2 (Perkin Elmer) image analysis software with built-in capability for tissue segmentation, cell segmentation and precise fluorescence quantification was used.

For tissue scoring, fluorescent intensity cut-off values were used to calculate positivity. The detailed method used for scoring tissues is described in supplementary methods. The panel labelled as 'Digital H&E' in the Figures is a simulated bright field image obtained by using Hoechst 33342 and tissue autofluorescence as masks. Markers of interest such as MCM2, γ H2AX or CD44v9 were pseudo-coloured brown and overlaid with the digital H&E image to enable the visualisation of positive cells in the context of the gastric gland architecture. The panel labelled as 'Scoring map' refers to inForm software-generated digital images where cells positive for the marker of interest were pseudo-coloured and overlaid on digital H&E images.

Antibodies used in the study

γH2AX (clone JBW301, #05–636, Merck Millipore), MCM2 (clone D7G11, #3619S, Cell Signalling technology), EpCAM (9C4, #324202, BioLegend), pan-Cytokeratin (C-11, #Ab7753, Abcam), CDX2 (#EPR2464Y, Ventana), Ki67 for mouse IHC (clone TEC-3, #M7249, Dako, Agilent), Ki67 for human IHC (clone MIB-1, #7240, Dako, Agilent), CD44v9 (clone RV3, #LKG-M001, Cosmo Bio USA), anti-Rat Alexa Fluor 488 (#A21202, Thermo Fisher Scientific), Click-iT EdU Alexa Fluor 488 Imaging kit (#C10337, Thermo Fisher Scientific).

Statistical analysis

All qIHC data were subjected to pair-wise comparisons and tested for statistical differences using the non-parametric Mann-Whitney U test. Differences in DNA methylation levels between groups were compared using the Student's t-test. Pearson's correlation coefficient was computed to study the correlation between two markers of interest. Comparisons with p < 0.05

were considered statistically significant. *denotes p<0.05, **denotes p<0.01, ***denotes p<0.001.

RESULTS

Optimisation of epithelial segmentation, cell proliferation and DDR signalling markers for qIHC analysis of gastric tissues

We describe below optimised qIHC methods for quantifying cell proliferation and DDR signalling in the stomach epithelial compartment.

Epithelial segmentation - Since the inForm image analysis software conducts automated epithelial segmentation, antibodies against the epithelial markers EpCAM or Cytokeratins (pan-Cytokeratin) were assessed for their ability to stain epithelial cells. However, a comparison of the mean fluorescence intensities (M.F.I) for EpCAM or pan-Cytokeratin individually revealed their lower expression in normal gastric tissues (online supplementary figure S2A,S2B). Hence, to enable robust algorithmbased tissue segmentation, tissues were probed for both EpCAM and pan-Cytokeratin. Tissue segmentation using

EpCAM and pan-Cytokeratin staining patterns revealed excellent identification of epithelial cells (online supplementary figure S2C). Thus, combined EpCAM/pan-Cytokeratin staining was conducted throughout this study for optimal tissue segmentation.

MCM2 as a cell proliferation marker - Next, to identify a reliable marker for gastric cell proliferation, we conducted in vivo experiments using mouse models. Wild-type mice were injected the synthetic thymidine analogue, EdU (5-Ethynyl-2'-deoxyuridine), which is incorporated into newly synthesised DNA and thus marks proliferating cells with the gastric gland. Immunofluorescence analysis of EdU using Click-iT chemistry showed EdU positivity within the isthmus region of corpus and at the base of the glands in the antrum (online supplementary figure S2D,S2E).

To further identify a sensitive marker of gastric epithelial proliferation in an IHC setting, two commonly used proliferation markers, Ki67 or MCM2 were compared with EdU. Of these, Ki67 is a nuclear antigen that marks dividing cells with widespread clinical use.²⁴ Mini-chromosome maintenance (MCM) proteins are DNA helicases that form the pre-replication complex and initiate DNA replication.²⁵ MCMs are expressed abundantly across all phases of the cell cycle but are degraded in cells that have exited the cell cycle such as differentiated, quiescent and senescent cells, enabling them to be used as biomarkers of cell cycle proliferation.²⁶

Similar to EdU incorporation, MCM2 and Ki67 antibodies also stained cells within the isthmus region of the corpus and at the base of the glands of the antrum. Closer examination revealed a broader zone of positivity for MCM2 with all EdU-positive cells being positive for MCM2. On the other hand, several EdU-positive cells were negative for Ki67 (online supplementary figure S2D,S2E). Validation studies in a K-Ras expressing precancerous GC model (eR1-CreERT2; Kras^{G12D} mice)²⁷ further revealed robust MCM2-positivity within the abnormal glands (online supplementary figure S2F). Lastly, human normal stomach, gastritis and IM tissues (n=3) were subjected to IHC staining of MCM2 and Ki67 to compare the two markers in human gastric tissues. Percent Ki67 or MCM2 positive nuclei over the nuclei of representative gastric/metaplastic cells were quantified in representative areas of the entire endoscopic biopsy. Ki67 and MCM2 staining closely correlated in normal and gastritis tissues, whereas the percent MCM2 positivity was slightly higher than Ki67 in some areas of IM (online supplementary figure S2G, online supplementary table 2). These

Stomach

data demonstrate the value of MCM2 as a sensitive gastric cell proliferation marker.

 γ H2AX as a marker of DDR signalling - The phosphorylation of histone H2AX at serine 139, also called as γ H2AX, is one of the most widely used markers of DNA damage signalling.²¹ FFPE cell blocks of HGC-27 cells were left untreated or exposed to irradiation (2 Gy) and stained with the γ H2AX antibody. As expected, radiation treatment induced γ H2AX signals (online supplementary figure S3A). As further validation of the γ H2AX antibody, M.F.I values for γ H2AX were significantly lower in HGC27 cells subjected to H2AX-knockdown as compared with control-knockdown cells on exposure to etoposide (online supplementary figure S3B,S3C).

Increased MCM2 positivity during gastric cancer progression

Using the methods established above, we profiled tissues spanning the stomach cancer spectrum. The TMA slide (ST1001) (online supplementary table 3) was stained with EpCAM/pan-Cytokeratin and MCM2 antibodies. MCM2 positivity per core was then averaged across two technical replicates and computed as $2.3\%\pm1.2\%$, $12.1\%\pm2.1\%$ and $25.9\%\pm3.1\%$, in the normal, gastritis and IM tissues, respectively (normal vs NAG, p=0.0043; normal vs IM, p=0.0022) (figure 1A and C).

To compare the IM stage with IGC, we used another TMA slide (IC00011B) with greater number of IM samples. Of the 40 cores analysed, 26 cores were atrophic gastritis with IM and 14 were IGC (online supplementary table 4). The percent positivity of MCM2 in the IM and adenocarcinomas was $25\% \pm 3.3$ and $37.81\% \pm 3.1\%$, respectively (IM vs IGC, p=0.0063) (figure 1B and D). Thus, positivity for MCM2 increased in the order of normal, NAG, IM and gastric adenocarcinomas, ascertaining increased proliferation through the carcinogenesis cascade.

Moderate increase in γH2AX positivity in gastric precancerous lesions

The TMA slide ST1001 (online supplementary table 5) were subjected to multiplex IHC staining with EpCAM/pan-Cytokeratin and yH2AX antibodies. The percent yH2AX positivity per core was then averaged across the two technical replicates and computed in the normal, gastritis and IM as 9.5%±3.9%, 19.75%±9.3% and 30.28%±9.2%, respectively (normal vs NAG, p=0.55; normal vs IM, p=0.0359; NAG vs IM, p=0.2810) (figure 2A and C). To compare DDR signalling levels between IM and IGC, TMA slide IC00011B was stained with EpCAM/pan-Cytokeratin and anti-yH2AX antibodies (online supplementary table 4). IHC analysis revealed significantly greater yH2AX-positivity at the IGC stage (IM vs IGC, 12.79%±2.10% and 33.9%±6.3%, respectively, p=0.0025) (figure 2B and D). These results indicate an increase in DDR signalling from normal tissue to IM, with further elevation in gastric adenocarcinomas.

Since the TMAs used above lacked clinical information or follow-up, we used samples from the GCEP cohort (n=19), comprising of normal, NAG and IM tissues for further validation (online supplementary table 6). γ H2AX-positivity was computed in the normal, gastritis and IM tissue as $6.4\% \pm 1.9\%$, $26.89\% \pm 9.34\%$ and $21.53\% \pm 6.7\%$, respectively (figure 3A and B) (normal vs NAG, p=0.1807; normal vs IM, p=0.0411). An overlay of γ H2AX-fluorescence the digital H&E image clearly showed γ H2AX-positive cells within glands harbouring the intestinal phenotype. Consistently, when serial sections of IM were stained either with antibodies against γ H2AX or CDX2 a known marker for IM,²⁸ many CDX2-positive cells were also positive

for γ H2AX, confirming that gastric cells that have undergone intestinalisation exhibit activated DDR signalling (online supplementary figure S4).

Taken together, the above results revealed elevated DDR signalling at the IM stage with further increase as lesions progress to IGC. Although γ H2AX-positive cells also accumulated at the NAG stage, inter-sample heterogeneity was observed rendering the data for this group statistically not significant when compared with other groups.

γH2AX positivity in gastric precancerous lesions strongly correlated to extent of chronic inflammation

Next, we then asked if the extent of chronic inflammatory cell infiltration determines DDR signalling. Chronic inflammation is known to create DNA damage by increasing the levels of ROS within cells.²⁹ ROS can damage DNA through the formation of oxidative adducts that collide with the replication fork or by activating DDR signalling directly through ATM activation.^{30 31} To investigate the relationship between yH2AX-positivity and chronic inflammation, we chose nine H. pylori negative gastritis tissues in order to avoid the confounding effects of H. pylori infection-dependent direct DNA damage.³² Since H. pylorinegative gastritis is associated with much lower levels of chronic inflammation as compared with H. pylori-positive gastritis,³³ it was possible to obtain a few regions on pathological examination that were negative for chronic inflammation (n=3) and compare DDR signalling within these with regions to those that were positive for chronic inflammation (n=6) (online supplementary table 7).

Tissues were subjected to IHC staining with EpCAM/pan-Cytokeratin, MCM2 and γ H2AX. The extent of cell proliferation was equivalent between the chronic inflammation negative and positive groups and computed as 27.35%±7.645 and 26.92±6.022, respectively (p=0.7143) (figure 4A and B). In contrast, γ H2AX-positivity was substantially higher in the chronic-inflammation positive (35.14%±9.76) as compared with the negative samples (3.074%±0.5863), (p=0.0238) (figure 4A and C). Thus, chronic inflammation seems to be a major activator of DDR signalling in precancerous gastric tissues.

IM lesions with heightened DNA damage signalling have lower mutational accumulation

Lastly, we interrogated the relationship of DDR signalling to mutational accumulation and CNAs at the IM stage. In a recent study, we characterised 146 IM lesions within the GCEP cohort for mutational counts, CNAs and average telomere length.²⁰ Notably, IM samples had a low mutational burden and CNAs as compared with gastric adenocarcinomas. Mutations frequently found in GC like *TP53*, *ARID1A* and *FBXW7* were infrequent in IM. Importantly, these observations implied that there exists a window of premalignant cell latency where IM lesions retain their genomic integrity before any further progression.

To interrogate the relationship of DDR signalling to mutational accumulation and CNAs, we extracted 15 samples of IM from the same cohort of 146 IM samples used in our previous published study. Of the 146 IM samples with genomic characterisation, IM samples without any mutational burden and/or CNAs were classified as the genome-stable category (n=8, samples IDs: 1 to 8) (online supplementary table 8, figure 5A and B). GCEP samples harbouring the highest mutational burden and CNAs were classified them as the genome-unstable category (n=7, samples IDs: 9 to 15). Notably, all samples were wild-type for p53.

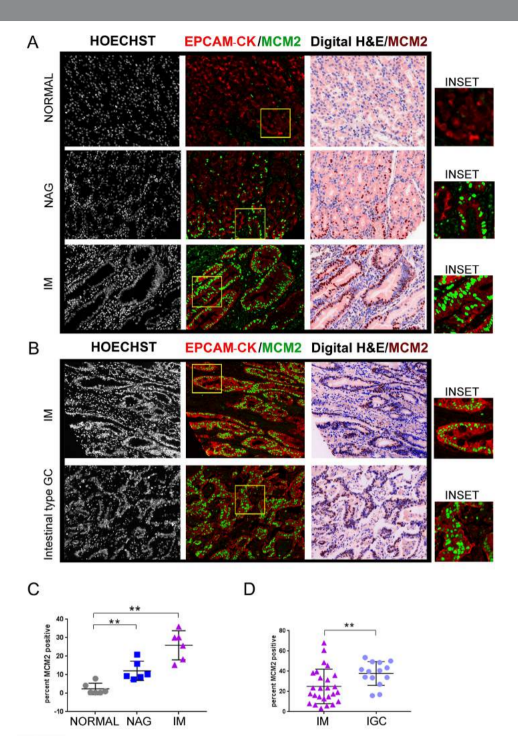


Figure 1 MCM2 positivity increases during GC progression. (A) TMA slide (ST1001) was stained with antibodies against EpCAM/pan-Cytokeratin (CK) and MCM2. Six cases each of normal gastric tissues, NAG and IM were analysed. (B) TMA slide (IC00011B) was stained with antibodies against EpCAM/pan-Cytokeratin (CK) and MCM2. Twenty-six cases of IM and 14 cases of IGCs were analysed. The panel labelled as digital H&E is a simulated bright field image obtained by using Hoechst 33342 and tissue autofluorescence as masks. The MCM2 staining pseudo-coloured as brown, has been overlaid on the the digital H&E image. Areas marked by the yellow rectangle within the fluorescent images have been zoomed by two-fold under the images marked as 'inset'. (C) Plot shows percent MCM2 positivity across the three stages, normal, NAG and IM. (D) Plot shows percent MCM2 positivity across IM and adenocarcinoma. Line within the vertical scatter plot shows the mean value. Error bar shows ±SD. Statistical significance was analysed by the Mann-Whitney non-parametric test. **p<0.01. GC,gastric cancer; IM, intestinal metaplasia; NAG, non-atrophic gastritis; TMA,tissue microarrays.

To test the hypothesis that DDR acts as an anti-cancer barrier, we asked if genome-stable samples had higher levels of DDR signalling as compared to the genome-unstable samples. FFPE-tissues obtained from the same biopsy site subjected previously to genomic characterisation were probed for EpCAM/pan-Cytokeratin, MCM2 and γ H2AX. Cell proliferation comparisons

through MCM2 percent positivity revealed slightly higher cell proliferation within the genome-stable group, although the difference did not reach statistical significance (genome-stable vs genome-unstable, mean percent positivity - 33.14 ± 9.69 vs 22.15 ± 9.5 , respectively, p=0.115) (figure 5C and E). A comparison of telomere length between the genome-stable and

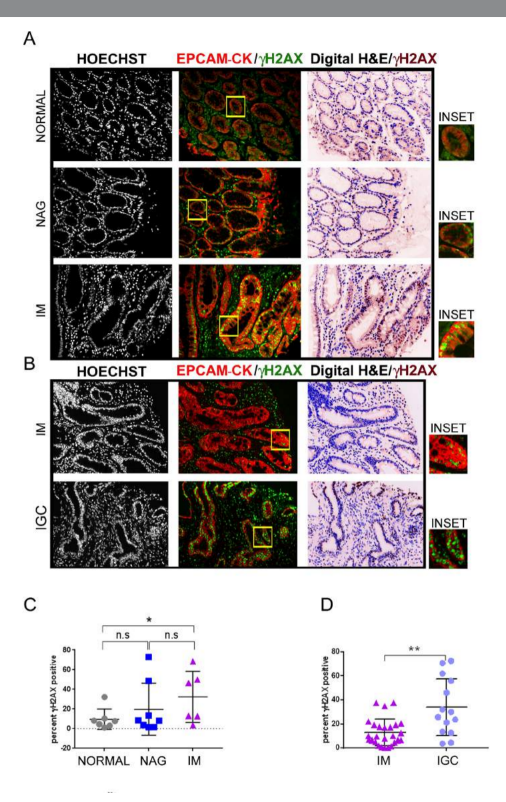


Figure 2 Modest increase in γ H2AX positivity during GC progression. (A) TMA slide (ST1001) was stained with antibodies against EpCAM/pan-Cytokeratin and γ H2AX. Seven cases of normal, eight cases of NAG and six cases of IM were analysed. (B) TMA slide (IC00011B) was stained with antibodies against EpCAM/pan-Cytokeratin (CK) and γ H2AX. Twenty-six cases of IM and 14 cases of IGCs were analysed. The γ H2AX staining shown with pseudo-brown colour has been overlaid on the digital H&E image. Areas marked by the yellow rectangle within the fluorescent images have been zoomed by two-fold under the images marked as 'inset'. (C) Plot shows percent γ H2AX positivity across the three stages, normal, NAG and IM. (D) Plot shows percent γ H2AX positivity across IM and IGCs. Line within the vertical scatter plot shows the mean value. Error bar shows ±SD. Statistical significance was analysed by the Mann-Whitney non-parametric test. * p<0.05, **p<0.01. GC,gastric cancer; IGC, intestinal-type gastric cancer; IM, intestinal metaplasia;NAG, non-atrophic gastritis; TMA, tissue microarrays.

genome-unstable groups also revealed no difference (data not shown). A comparison of IM-type (complete vs incomplete) did not reveal any significant differences between the two groups.

In contrast to the above parameters, DDR signalling was significantly higher in the genome-stable group ($\%\gamma$ H2AX positivity - 20.45±7.5,) as compared with the genome-unstable

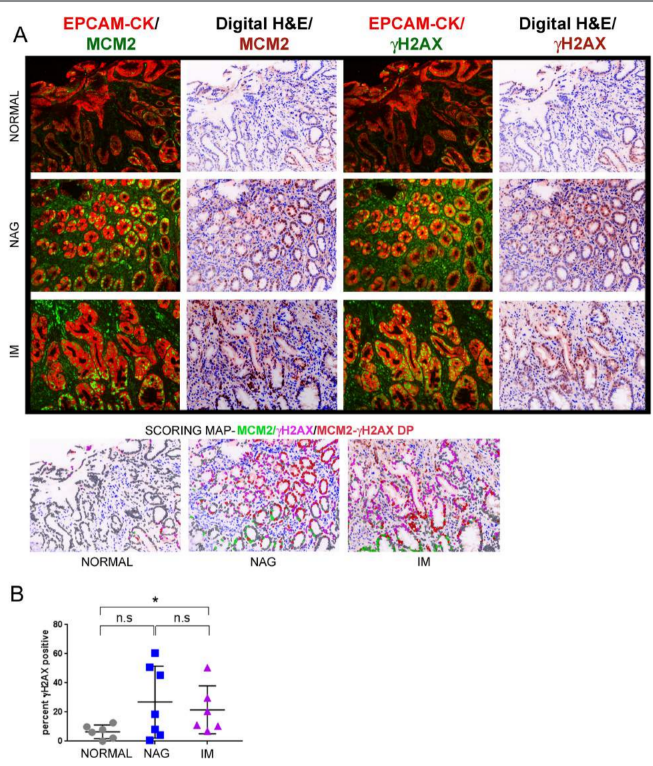


Figure 3 γ H2AX positivity increase in IM lesions as compared to normal controls in the GCEP cohort. (A) Samples from the GCEP cohort comprising of six cases of normal gastric tissue, seven cases of NAG and six cases of IM were subjected to multiplex IHC staining with antibodies against EpCAM/ pan-Cytokeratin (CK), MCM2 and γ H2AX. Immunofluorescence images of γ H2AX or MCM2 with EpCAM/pan-Cytokeratin (CK) are shown. Scoring map depicts epithelial cells positive for γ H2AX (magenta), MCM2 (green), γ H2AX/MCM2 double-positive (DP) (red) and γ H2AX/MCM2 double-negative (grey). (B) Plot shows percent γ H2AX positivity across the three stages, normal, NAG and IM. Line within the vertical scatter plot shows the mean value. Error bar shows ±SD. Statistical significance was analysed by the Mann-Whitney non-parametric test. * p<0.05, n.s-not significant. GCEP, gastric cancer epidemiology programme; IHC, immunohistochemistry; IM, intestinal metaplasia; NAG, non-atrophic gastritis.

group (% γ H2AX positivity - 6.3±4.5) in most of the samples (except one case) (p=0.04) (figure 5C,D and E). On manual histological examination of the scoring maps, glands harbouring the intestinal phenotype harboured most of the DDR activation.

Since DDR signalling can either arrest cells in the cell cycle or render them senescent, we asked if γ H2AX signals and MCM2 expression are correlated. Nuclear fluorescence values generated by the image analysis software was used to accurately

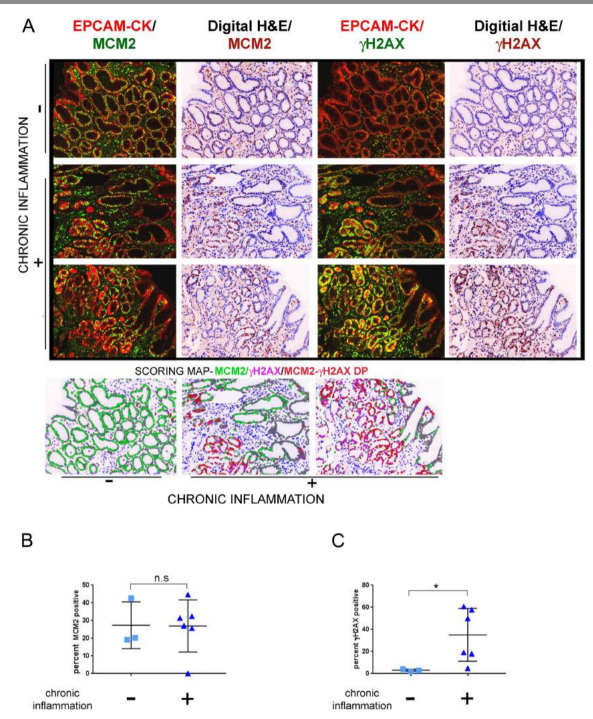


Figure 4 γ H2AX positivity correlates with extent of chronic inflammation. (A) Tissues were classified according to extent of chronic inflammatory cell infiltration as positive (n=6) or negative (n=3) and subjected to multiplex immunohistochemistry staining with antibodies against EpCAM/pan-Cytokeratin (CK), MCM2 and γ H2AX. Immunofluorescence images of MCM2 or γ H2AX with EpCAM/pan-Cytokeratin (CK) are shown. Scoring map depicts epithelial cells positive for γ H2AX (magenta), MCM2 (green), γ H2AX /MCM2 double-positive (DP) (red) and γ H2AX/MCM2 double-negative (grey). (B) Plot shows percent MCM2 positivity in the chronic inflammation positive and negative groups. (C) Plot shows percent γ H2AX positivity in the chronic inflammation positive and negative groups. Line within the vertical scatter plot shows the mean value. Error bar shows ±SD. Statistical significance was analysed by the Mann-Whitney non-parametric test. * p<0.05, n.s-not significant.

compute the Pearson's correlation coefficient (*r*) between the two markers. Intriguingly, MCM2 and γ H2AX signals had a varied relationship across difference samples and were positively correlated in two samples (sample ID #6; *r*=+0.66, p<0.00001 and sample ID #7; *r*=+0.68, p<0.00001), and had no clear correlation in others. Manual examination further confirmed the heterogeneous relationship between MCM2 and γ H2AX

signals and imply that despite activated DDR signalling many IM lesions may remain MCM2 positive and do not necessarily exit the cell cycle. The potential implication of these observations to IM progression is discussed below ('Discussion').

The intriguing observation of reduced DDR signalling in genome-unstable lesions led us to hypothesise that these IM lesions may have elevated expression of CD44v9, a variant

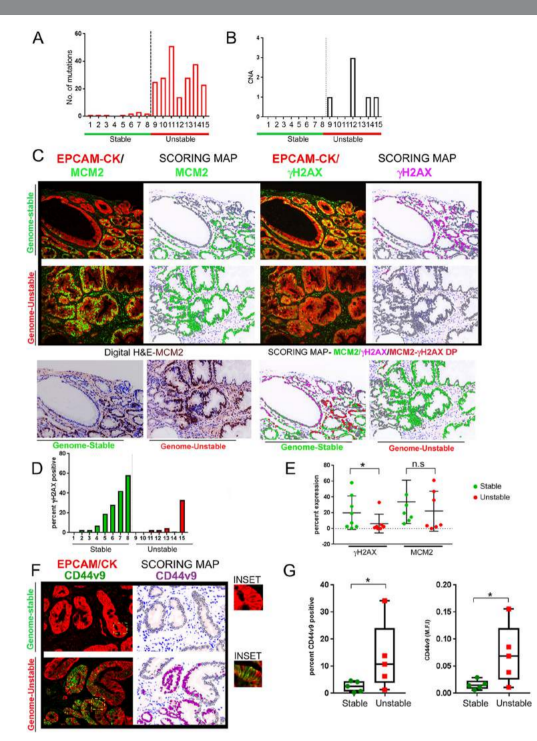


Figure 5 γ H2AX positivity inversely correlates with mutational accumulation in IM samples. (A) IM samples were chosen from the GCEP cohort (n=15) and categorised either as genome-stable or genome-unstable, based on their mutational counts and CNAs. Mutational count of the 15 samples is shown. Genome-stable (n=8), genome-unstable (n=7). (B) CNAs for the 15 samples are shown. Genome-stable (n=8), genome-unstable (n=7). (C) IM tissues were subjected to multiplex IHC staining with antibodies against EpCAM/pan-Cytokeratin (CK), MCM2 and γ H2AX. Regions with low epithelial content (<30%) or low IM coverage (<70%) were removed. (upper) Immunofluorescence images of MCM2 or γ H2AX with EpCAM/ pan-Cytokeratin (CK) are shown. (lower left) Digit H&E to show IM tissue architecture. (lower right) Scoring map depicts epithelial cells positive for γ H2AX (magenta), MCM2 (green), γ H2AX/MCM2 double-positive (red) and γ H2AX/MCM2 double-negative (grey). (D) Graph shows percent γ H2AX positivity across the IM samples. Bars in green and red represent the genome-unstable and genome-unstable categories, respectively. (E) Plot shows percent γ H2AX positivity and MCM2 in genome-stable and genome-unstable categories. (F) Immunofluorescence images of CD44v9 co-staining with EpCAM/pan-Cytokeratin (CK) are shown (left). Scoring map shows epithelial cells positive for CD44v9 (magenta coloured) or negative for CD44v9 (grey) (right). CD44v9 signal swere either 'dim' or 'bright' within IM regions, and both types of signals were included for CD44v9 signal quantification. Areas marked by the yellow rectangle within the fluorescent images have been zoomed by two-fold under the images marked as 'inset'. (G) Plots show CD44v9 percent positivity (left) and M.F.I (right) in the genome-stable and genome-unstable groups. Line within the vertical scatter plot shows the mean value. Error bar shows ±SD. Statistical significance was analysed by the Mann-Whitney non-parametric test. CNA, copy-number alteration; GCEP, gastric cancer epidemiology

form of CD44. CD44 is complex gene comprising of multiple variant forms of CD44 (CD44v), of which CD44v9, an exon 9 containing variant isoform has been shown to be a functional gastric stem cell marker and predictive of recurrence in early gastric cancers.^{34–36} CD44v9 has been shown to detoxify chronic inflammation-mediated ROS and accelerate cancer progression by increasing ROS defence.²² We hypothesised that higher expression of CD44v9 in the genome-unstable IM lesions may defend these cells from inflammation-generated ROS and facilitate their escape from tumour-suppressive consequences of DDR signalling.

We tested the hypothesis that genome-unstable samples with attenuated DDR signalling have higher CD44v9 expression. The expression of CD44v9 and its relationship to yH2AX levels was studied in IM lesions. IM samples were subjected to qIHC analyses with the antibodies against EpCAM/pan-Cytokeratin, CD44v9 and yH2AX. Due to insufficient patient material, five samples each from the genome-stable and genome-unstable groups were used. Clear positive regions of membrane-associated CD44v9 staining were seen within the IM regions (figure 5F). Following image quantification, percent positivity of CD44v9 was higher in the genome-unstable group (%CD44v9 in genome-stable vs genome-unstable, 2.57±1.94 and 13.3±12.59, respectively, p=0.0278) (figure 5G, left). Similarly, fluorescence intensity of CD44v9 was also higher in the genome-unstable group (M.F.I CD44v9 in genome-stable vs genome-unstable, 0.015±0.008 and 0.072 ± 0.054 , respectively, p=0.0476) (figure 5G, right).

Next, since CD44v9 expression promotes ROS defence, we asked whether CD44v9 staining and γ H2AX are negatively correlated and performed correlation analysis between the two markers. Interestingly, γ H2AX and CD44v9 intensities were inversely correlated across all the samples and the values reached statistical significance (*r* values ranged from -0.476 to -0.70, p<0.001, online supplementary table 9).

Then, towards our effort to integrate IHC-based analysis with (epi)genomic characterisation we examined whether DDR genes are mutated or if their promoters undergo hypermethylation in IM samples. A list of genes found mutated in the IM lesions used for this study were compiled (online supplementary appendix 1). An overlap of the mutated genes from genome unstable group with manually curated DNA repair gene-set (online supplementary appendix 2) revealed that *ATM*, *FANCA*, *HLTF*, *POLE* and *REV3L* were mutated in two of the samples from the genome-unstable group (*ATM* in sample ID #10 and *FANCA*, *HLTF*, *POLE* and group).

Lastly, we also asked if genome-unstable IM samples displayed epigenetic inactivation of DDR genes through promoter hypermethylation. In our earlier study, DNA methylation levels of IM lesions was compared with normal samples, and DNA methylation levels were categorised as 'low', 'intermediate' or 'high' relative to normal gastric tissue.²⁰ Around 2537 regions were hypermethylated in the methylation-high IMs relative to normal antral tissues. Of note, methylation-high antral IMs showed more mutational accumulation and CNAs leading us to test here whether IM-associated promoter hypermethylation of DDR genes is higher in the genome-unstable group.

The DNA repair genes WRN, MLH1 and RAD52 showed modest but consistent increase in promoter hypermethylation in the genome-unstable IM samples when compared with 39 normal control gastric tissues (delta β values for WRN, MLH1 and RAD52 following comparison between genome unstable vs normal samples was 0.15 ± 0.14 , 0.079 ± 0.042 and 0.073 ± 0.044). On the other hand, the aforementioned DNA

repair genes has minimal hypermethylation in the genome-stable samples as compared with normal controls (delta β values *WRN*, *MLH1* and *RAD52* following comparison between genome stable vs normal samples was 0.009 ± 0.019 , 0.005 ± 0.012 and 0.005 ± 0.008). The promoter methylation differences between genome stable and unstable samples was statistically significant (p<0.05) (online supplementary appendix 3). Taken together, reduced DNA damage signalling possibly through the increased expression of CD44v9, along with mutational/epigenetic alterations in DNA repair genes may be factors contributing towards the greater genome instability of some IM lesions. An integrated view of the major alterations observed in genome-unstable IM lesions based on genomic analysis and qIHC findings is compiled in online supplementary table 10.

DISCUSSION

Here, we interrogate the relationship of DDR signalling to mutational accumulation and gain insights on the tumour-suppression mechanisms operational in IM. In earlier studies, activated forms of DDR signalling factors were shown to accumulate and engage tumour suppressive networks in premalignant lesions. It was proposed that the DDR creates an inducible barrier against cancer progression by preventing the evolution of malignant mutants from DNA damage-afflicted premalignant cells.^{37 38} However, due to the difficulty in procuring sufficient precancerous tissue for concurrent genomic and IHC study, it has been difficult to test if DDR signalling indeed prevents mutational accumulation. By conducting qIHC analysis of DDR signalling and cell proliferation, our studies reveal that IM lesions with higher levels of DDR signalling have a lower mutational accumulation. By linking yH2AX-positivity indicative of DDR activation to the degree of mutational accumulation, this work provides evidence for DDR as an anti-cancer barrier in preneoplastic lesions.

Our studies with the genome-stable IM lesions reinforce previous studies on DDR as anti-cancer barrier in lung, colon, breast, urinary bladder and prostate cancers.^{39–42} In these lesions, oncogene-induced DNA replication stress and the ensuing double strand breaks were seen as the main stimuli for DDR signalling activation. However, gastric IMs are unique in that they have a very low mutational burden (less than ${\sim}5\%$ of samples) and classic oncogene activation is seldom seen.²⁰ Thus, chronic inflammation-elicited ROS, instead of oncogene-induced replication stress, is likely to be the major activator of DDR signalling in precancerous gastric lesions. Supportive of this assumption, yH2AX-positivity and chronic inflammation were strongly correlated, suggesting an important role for chronic inflammation as the driver of gastric DDR signalling (figure 4). Intriguingly, many of the cells positive for γ H2AX in IM lesions were also MCM2-positive (figure 5), indicating that despite exhibiting activated DDR signalling, such cells do not exit the cell cycle. Instead, we speculate that cells with an activated DDR signalling are arrested in the cell cycle and may turn into a future reservoir of malignant clones once their DDR signalling is turned off due to mutations or epigenetic alterations in DDR/ cell cycle checkpoint components. Examining the status of cell cycle checkpoint proteins such as pCHK1 or pCHK2 and senescent markers in IM lesions, especially in longitudinal studies can provide greater clarification.

Our studies with the genome-unstable IM lesions suggest a role for CD44v9 in attenuating DDR signalling arising from chronic inflammation. This is because CD44v has been shown to detoxify chronic inflammation-mediated ROS, by interaction with xCT (glutamate-cystine transporter) and increasing intracellular glutathione levels.^{22 43} Interestingly, CD44v9 expression was found to be an important marker of recurrence in primary early gastric cancer.³⁵ Indeed, CD44v9 expression was found to be higher in the genome-unstable lesions and showed negative correlation with YH2AX staining intensities (online supplementary table 9). Thus, greater ROS adaptability and the concomitant attenuation in tumour-suppressive DDR signalling could be a major risk factor for IM lesions. Additionally, DNA repair gene mutational inactivation (2/5 samples) coupled with epigenetic inactivation of DDR genes (4/5 samples) could potentially collaborate with elevated ROS defence and facilitate IM progression (Model, online supplementary figure S5B and S5C). It is to be noted that despite being statistically significant (p < 0.05), the degree of promoter hypermethylation is modest in genomeunstable IM because our study has used very early precancerous lesions and compared their promoter methylation with normal samples. Also, since IM lesions have relatively lower levels of genomic instability, our sample numbers had to be restricted. More studies on larger cohorts are needed to further clarify the role of the above-identified risk factors in IM to GC progression.

Lastly, the epithelial segmentation protocol and algorithms described here for quantitative IHC are broadly applicable to any epithelial cancer. The value of MCM2 as an early marker of cancer proliferation has been demonstrated for several precancerous lesions tissues such as bronchial dysplasia, premalignant lung lesions, human papillomavirus-infected squamous lesions of the cervix, epithelial dysplasia of the tongue.^{44–47} MCM2 was identified as a sensitive proliferation marker of gastric cardia cancer in a previous study and possibly with potential prognostic value.⁴⁸ Our study re-enforces the view that MCM2 and Ki67 staining should be combined to obtain a better measure of cancer cell proliferation.

The origins of intestinal-type gastric cancer reflect the interplay between bacterial infection, chronic inflammation, tumour-associated epigenetic/epigenomic changes and host factors.^{49 50} Of these, chronic inflammation is a well-known perpetrator of DNA damage and genomic instability. Our observations imply that the competence with which IM cells respond to inflammation-dependent ROS through DDR signalling likely determines the mutational status and future fate of premalignant lesions. An integrated approach that combines multiplex γ H2AX/MCM2/CD44v9 profiling with genomic data on larger cohorts will shed more light on the biological determinants of IM to GC evolution.

Author affiliations

- ¹Cancer Science Institute of Singapore, National University of Singapore, Singapore 117599, Singapore
- ²Program in Cancer and Stem Cell Biology, Duke-NUS Graduate Medical School, Singapore 169857, Singapore
- ³Department of Pathology, National University of Singapore, Singapore 119228, Singapore
- ⁴Department of Medicine, Yong Loo Lin School of Medicine, National University of Singapore, Singapore 119228, Singapore
- ⁵Department of Gastroenterology and Hepatology, National University Health System, Singapore,119074, Singapore
- ⁶Department of Surgery, National University of Singapore, Singapore 119228, Singapore
- ⁷Singapore Gastric Cancer Consortium, Singapore 119074, Singapore
- ⁸Department of Gastroenterology & Hepatology, Singapore General Hospital, Singapore 169854, Singapore
- ⁹Department of Gastroenterology and Hepatology, Tan Tock Seng Hospital, Singapore
 ¹⁰Department of Gastroenterology, Changi General Hospital, Singapore 529889,
- ¹⁰Department of Gastroenterology, Changi General Hospital, Singapore 529889 Singapore

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Contributors IHC data acquisition and analysis, DL, PMH, VK and SS; Pathological interpretations and grading of biopsy tissues, SS and MT; Imaging platform and staining optimisation, PMH, VK and ADJ; Mouse data acquisition, JM; Genomic analysis, KKH and PT; Clinical data collection, ZF, JS, CK, ST, KMF and TLA; Data interpretation, VK, SS, ADJ, MT and TI; Drafting and writing manuscript, VK and YI; Revising work, VK, PMH, SS and MT; Conceptualisation and design, YI, YKG and VK; Final approval for publishing work, YKG and YI.

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Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information.

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ORCID iDs

Vaidehi Krishnan https://orcid.org/0000-0002-5028-6567 Debbie Xiu En Lim https://orcid.org/0000-0002-2772-0580 Phuong Mai Hoang https://orcid.org/0000-0002-2772-0580 Supriya Srivastava https://orcid.org/0000-0002-6727-3076 Junichi Matsuo https://orcid.org/0000-0002-2378-2370 Kie Kyon Huang https://orcid.org/0000-0002-6727-3076 Feng Zhu https://orcid.org/0000-0001-6030-9251 Khek Yu Ho http://orcid.org/0000-0002-2932-1962 Jimmy Bok Yan So https://orcid.org/0000-0002-9772-7905 Christopher Khor https://orcid.org/0000-0002-1409-5691 Stephen Tsao https://orcid.org/0000-0002-9432-3382 Ming Teh https://orcid.org/0000-0001-6879-8678 Kwong Ming Fock https://orcid.org/0000-0002-2254-2260 Tiing Leong Ang https://orcid.org/0000-0001-9993-8549 Anand D Jeyasekharan https://orcid.org/0000-0001-9816-6137 Patrick Tan https://orcid.org/0000-0002-0179-8048 Khay-Guan Yeoh https://orcid.org/0000-0002-7802-4606 Yoshiaki Ito http://orcid.org/0000-0002-9037-1184

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Figure S1

Details on experimental conditions during multiplex qIHC staining. A sequential representation of the major steps during qIHC staining are (a) tissue staining with fluorescent antibodies (b) image acquisition (c) fluorophore unmixing (d) tissue segmentation (e) cell segmentation (f) scoring for marker positivity.

Figure S2

Optimization of protocols for gastric epithelial cell segmentation and cell proliferation. (A) TMA slide (ST1001) comprising of normal gastric tissues (n=8), non-atrophic gastritis (n=9), IM (n=6) and IGC (n=13) were stained with antibodies against (A) EpCAM (B) pan-Cytokeratin. Plot shows M.F.I for the indicated proteins across cancer stages. (C) Normal stomach tissue was stained by mixing EpCAM and pan-Cytokeratin antibodies. (Left) Gastric architecture was revealed by a digitally generated H&E image using autofluorescence and Hoechst 33342 staining as a mask. (Right) Tissue segmentation was performed using combined EpCAM and pan-Cytokeratin (CK) staining. (D and E) Wild type mice were injected twice with EdU within 24 hours. Tissues harvested after 24 hours were subjected to IHC staining with antibodies against Ki67 and MCM2. EdU signals were detected using click-IT chemistry. D-antrum, E-corpus. (F) Stomach tissue from transgenic *eR1*-CreERT2;LSL-*Kras*^{G12D} mice were subjected to IHC staining with antibodies against MCM2, E-cadherin and Ki67. (G) Ki67 and MCM2 nuclear expression in representative cases of normal gastric glands, marked gastritis and marked IM in GCEP samples. Arrows show nuclear expression (200X).

Figure S3 Antibody optimization for DNA damage detection in gastric precancerous lesions. (A) HGC-27 cells were exposed to irradiation (2Gy) and cells were harvested after 2 hours. Plot depicts percent the mean fluorescence intensity (M.F.I) of γ H2AX. (B) A549 cells were subjected to H2AX-siRNA for three days. Q-PCR analysis demonstrates efficient H2AX-knockdown. (C) Control or H2AX-knockdown cells were exposed to Etoposide (10 μ M) for 6 hours, followed which cells were harvested and processed for quantification of signals using the operetta high-throughput system. γ H2AX nuclear fluorescence is shown in the plot. ***p<0.001. Error bars represent S.D.

Figure S4 Immunohistochemical analysis of CDX2 and γ H2AX in serial sections of Intestinal metaplasia. (Left) A IM biopsy from the GCEP cohort was subjected to multiplex immunofluorescence staining with EpCAM/pan-Cytokeratin, MCM2 and γ H2AX. Scoring map shows the MCM2-positive cells in green. γ H2AX-positive cells in magenta, MCM2/ γ H2AX double-positive (DP) cells in red and MCM2/ γ H2AX double-negative cells in grey. (Right) A serial section from the same tissue was subjected to conventional DAB-mediated IHC staining. Note: since the serial sections were subjected to different protocols (Multiplex qIHC vs conventional IHC with DAB-based detection), the area of the image and orientation of glands are slightly different between the right and left images. Two representative images labelled as image 1 and image 2 are shown.

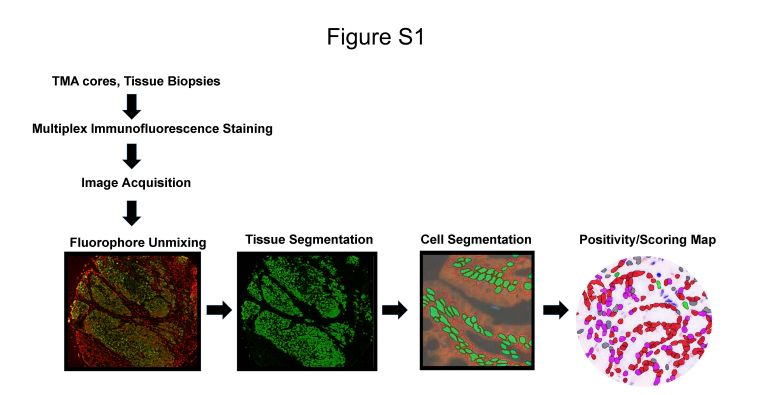
Figure S5 Status of DNA damage signaling and DNA hypermethylation in IM lesions and an integrated model summarizing the role of DNA damage signaling in premalignant gastric lesions. (A) Heatmap shows a list of genes that are differentially hypermethylated in genome unstable IM samples. Briefly, DNA methylation data from Huang et al (1) was used compute delta β values for the promoter regions of DNA repair genes in IM samples against 39 normal gastric samples. The student's T-test was used to compare the average delta β value of genome-stable and genome-unstable samples for statistical significance. * p<0.05 (B) A schematic representing the pattern of MCM2 and γ H2AX staining spanning the premalignant stages. (C) Model depicting the relationship between DDR signalling in

premalignant lesions and GC progression. Left- *Genome-Stable IM*: When the genomes of IM cells encounter chronic inflammation-mediated DNA damage, the activation of the DDR pathway (high γ H2AX levels and wild-type p53 status) suppresses mutational accumulation via cell cycle arrest. As long as such IM lesions do not accumulate additional hits that incapacitate their DDR signaling, IM samples with higher DNA damage are predicted to have a lower chance for progression. Right-*Genome-Unstable IM*: When the genomes of IM cells with high CD44v9 expression encounter chronic inflammation-mediated DNA damage, DDR signalling is suppressed in the IM lesions. Such IM lesions have higher ROS defence thus manage to avoid tumor-suppressive networks like the DDR, enabling cells to divide and accumulate mutational/chromosomal aberrations. Epigenetic alterations in DNA repair gene may further fuel genomic instability is such IM lesions. Thus, greater ROS defence in combination with (epi)genomic aberrations in DNA repair pathways can render some IM lesions more permissive to the accumulation of genomic alterations and pose a higher risk for progression.

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Gut



Supplementary Table 1

Details on experimental conditions for multiplex qIHC staining. Table shows the antibodies used in the current study, antigen retrieval conditions and details on dilution of primary and secondary antibodies. NA-not applicable

Supplementary Table 2

Scoring of Ki67 and MCM2 staining in human gastric samples. Immunohistochemical staining for Ki67 and MCM2 were performed and cells positive for nuclear signals are shown as a percentage of all nuclei in representative areas. Values shown are mean positivity per sample.

Supplementary Table 3

Patient details for TMA used for MCM2 staining. For cores stained with anti-MCM2 and data shown in figure 1A and figure 1C, patient clinical information is provided.

Supplementary Table 4

Patient details for TMA with IM and IGC cores. For cores stained with anti- MCM2/γH2AX and data shown in figure 1B and figure 1D, patient clinical information is provided.

Supplementary Table 5

Patient details for TMA used γ **H2AX staining**. For cores stained with anti- γ H2AX and data shown in figure 2A and figure 2C, patient clinical information is provided.

Supplementary Table 6

Patient details for GCEP samples used for \gammaH2AX staining. For GCEP tissues stained with anti-MCM2/ γ H2AX and data shown in figure 3, patient clinical information is provided.

Supplementary Table 7

Patient details for MCM2 and γ H2AX staining of gastric tissues with different extents of inflammation. For gastric tissues stained with anti-MCM2/ γ H2AX and data shown in figure 4, patient clinical information is provided.

Supplementary Table 8

Patient details for MCM2 and γ H2AX staining of IM samples from GCEP with different extents of genomic instability. For IM samples stained with anti-MCM2/ γ H2AX and data shown in figure 5, patient clinical information is provided.

Supplementary Table 9

Correlation between CD44v9 and γ **H2AX in IM samples**. Pearson's correlation analysis was performed to compare the relationship between CD44v9 and γ H2AX staining. Cells positive for γ H2AX, CD44v9 or both were subjected to Pearson's correlation analysis and tested for statistical significance. *r* (Pearson's correlation coefficient) was computed for highest biopsy region expressing CD44v9. *Sample ID numbers are matched to patient IDs from figure 5A-D.

Supplementary Table 10

An integrated view of alterations observed in the IM lesions subjected to genomic and IHC analysis. For the samples used for DDR signalling and cell proliferation analysis (Figure 5A-E), % CD44v9 expression, mutational accumulation, DNA repair gene mutations, CNAs, promoter hypermethylation and differential hypermethylation of DNA repair genes is shown. CNAs were identified using the GATK4 workflow for allelic copy number variation (ACNV; https://software.broadinstitute.org/gatk/). *Sample ID numbers are matched to patient IDs from figure 5A-D. **NA- Data not available.

Supplemental Appendix

Supplemental Appendix 1

A list of genes found mutated in IM samples. Sample ID numbers are matched to patient IDs from figure 5A-D.

Supplemental Appendix 2

Manually curated list of DNA repair and DNA damage signaling genes

Supplemental Appendix 3

DNA methylation analysis of IM samples. The methylation data from Huang et al (1) was used measure the average β values for the promoter regions of DNA repair genes. The column labelled "Normal" is the average methylation β values of 39 normal gastric samples. Sample ID numbers are matched to patient IDs from figure 5A-D.

References

1. Huang KK, Ramnarayanan K, Zhu F, Srivastava S, Xu C, Tan ALK, et al. Genomic and Epigenomic Profiling of High-Risk Intestinal Metaplasia Reveals Molecular Determinants of Progression to Gastric Cancer. Cancer Cell. 2018;33(1):137-50 e5.

| Antibody | Antigen Retrieval | Primary Antibody | Secondary Antibody)/ dilution | Hoechst 33342 | TSA/ dilution |
|---|---|---|--|------------------|-----------------------|
| Anti-MCM2 | Steaming at pH.6.0, 20 mins | 1:1000 | Anti-Rabbit, 1:1000 | 1:100 | Opal 520, 1:100 |
| Anti-γH2AX | Microwaving at pH 9.0, 16 mins | 1:1000 | Anti-Mouse, 1:1000 | 1:100 | Opal 690, 1:100 |
| Anti- EpCAM, Anti-pan- Cytokeratin | Steaming at pH.6.0, 20 mins and microwaving at pH 6.0, 16 mins | 1:250 (EpCAM), 1:1000 (pan- Cytokeratin) | Anti-Mouse, 1:1000 | 1:100 | Opal 570, 1:100 |
| Anti-CD44v9 | Steaming at pH 9.0, 20 mins | 1:100 | Anti-Rat (Alexa Fluor 488), 1:200 | 1:100 | NA |

Supplementary Table 1. Details on experimental conditions during multiplex qIHC staining

Supplementary Table 2. Scoring of Ki67 and MCM2 staining in human gastric samples.

| Sample | Ki67/MCM2-Tissue (A) | Ki67/MCM2-Tissue (B) | Ki67/MCM2-Tissue (C) |
|-----------|----------------------|----------------------|----------------------|
| | | | |
| Normal | <10%/<10% | <10%,/<10% | <1%/<1% |
| Gastritis | 40%/40% | 50%/50% | 30%/20% |
| Gastritis | 40 78/40 78 | 50 787 50 78 | 3078/2078 |
| IM | 30%/70% | 40%/60% | 15%/5% |

Supplementary Table 3. Patient details for TMA used for MCM2 staining

| Clinicopathological features | Normal | Inflammation | Intestinal Metaplasia |
|---------------------------------|--------|--------------|-----------------------|
| | N=6 | N=6 | N=6 |
| Age (Mean) | 40.5 | 57.7 | 55.5 |
| Age range | 16-56 | 53-69 | 40-74 |
| Sex | | | |
| Male | 5 | 3 | 5 |
| Female | 1 | 3 | 1 |

Supplementary Table 4. Patient details for TMA with IM and IGC cores

| Clinicopathological features | Intestinal Metaplasia | Adenocarcinoma | |
|------------------------------|-----------------------|----------------|--|
| | N=26 | N=14 | |
| Age (Mean) | 58.9 | 58.1 | |
| Age range | 32-74 | 43-68 | |
| Sex | | | |
| Male | 23 | 11 | |
| Female | 3 | 3 | |

Supplementary Table 5. Patient details for TMA used γ H2AX staining

| Clinicopathological features | Normal | Inflammation | Intestinal Metaplasia |
|------------------------------|--------|--------------|-----------------------|
| | N=7 | N=8 | N=6 |
| Age (Mean) | 37.4 | 58.6 | 55.5 |
| Age range | 16-50 | 52-71 | 40-74 |
| Sex | | | |
| Male | 6 | 4 | 5 |
| Female | 1 | 4 | 1 |

Supplementary Table 6. Patient details for GCEP samples used for γ H2AX staining

| Clinicopathological features | Normal | Gastritis | Intestinal Metaplasia |
|---------------------------------|--------|-----------|-----------------------|
| | N=6 | N=7 | N=6 |
| Acute Inflammation | | | |
| Negative | 6 | 3 | 3 |
| Mild/Moderate | - | 3 | 3 |
| Marked | - | 1 | - |
| Chronic Inflammation | | | |
| Negative | 6 | - | - |
| Moderate | - | - | 3 |
| Marked | - | 7 | 3 |
| H. pylori status | | | |
| Negative | 6 | 2 | 5 |
| Mild/Moderate | - | 3 | 1 |
| Marked | - | 2 | |
| Atrophy | | | |
| Negative | 6 | 7 | - |
| Mild/Moderate | - | - | 3 |
| Marked | - | - | 1 |
| Unknown | - | - | 2 |
| Smoking | | | |
| Y | - | 1 | 3 |
| Ν | 6 | 6 | 3 |
| Alchohol | | | |
| Y | - | 1 | 1 |
| Ν | 6 | 6 | 5 |

Supplementary Table 7. Patient details for MCM2 and γ H2AX staining of gastric tissues with different extents of inflammation

| Clinicopathological features | |
|------------------------------|-----|
| | N=9 |
| Inflammation Score | |
| Negative | 3 |
| Mild/Moderate | 6 |
| Marked | - |
| H. pylori status | |
| Negative | 9 |
| Positive | - |

Supplementary Table 8. Patient details for MCM2 and γ H2AX staining of IM samples from GCEP with different extents of genomic instability

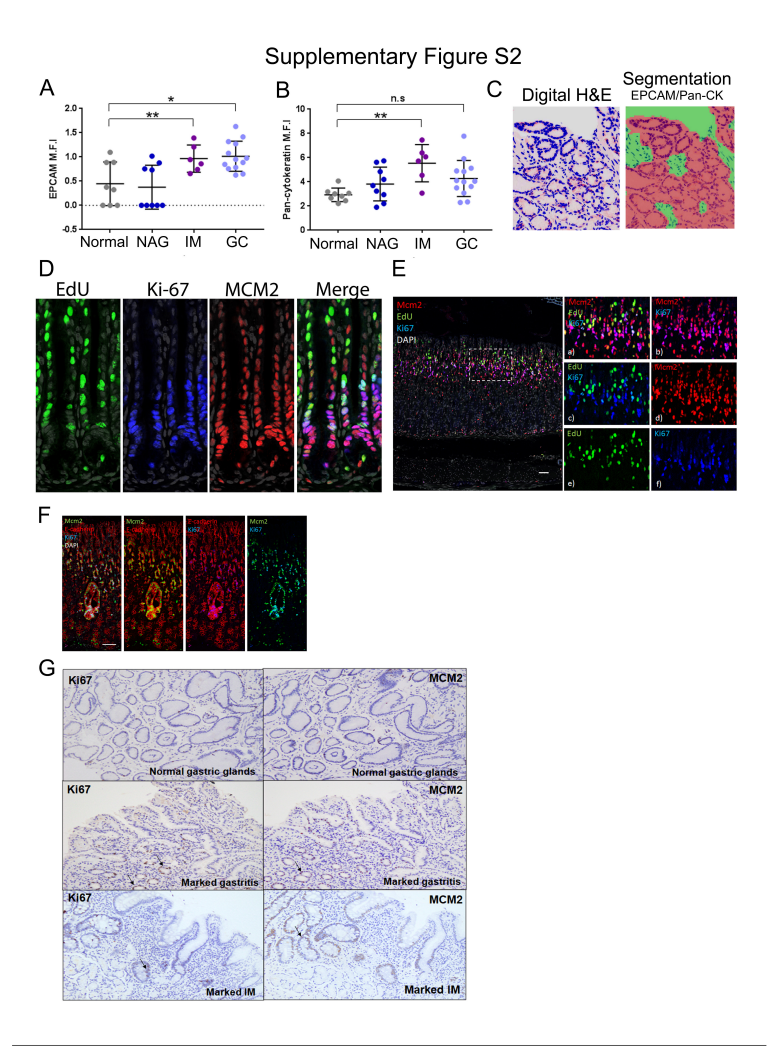
| Clinicopathological features | Genome stable | Genome unstable |
|------------------------------|---------------|-----------------|
| | N=8 | N=7 |
| Chronic Inflammation | | |
| Negative | 1 | - |
| Mild/Moderate | 7 | 7 |
| Marked | - | - |
| Acute Inflammation | | |
| Negative | 7 | 6 |
| Mild/Moderate | 1 | 1 |
| Marked | - | - |
| Atrophy | | |
| Negative | 2 | 1 |
| Mild/Moderate | 3 | 2 |
| Marked | - | - |
| Unknown | 3 | 4 |
| <i>H. pylori</i> status | | |
| Negative | 7 | 7 |
| Positive (mild) | 1 | - |
| IM status | | |
| Mild/Moderate | - | 1 |
| Marked | 8 | 6 |

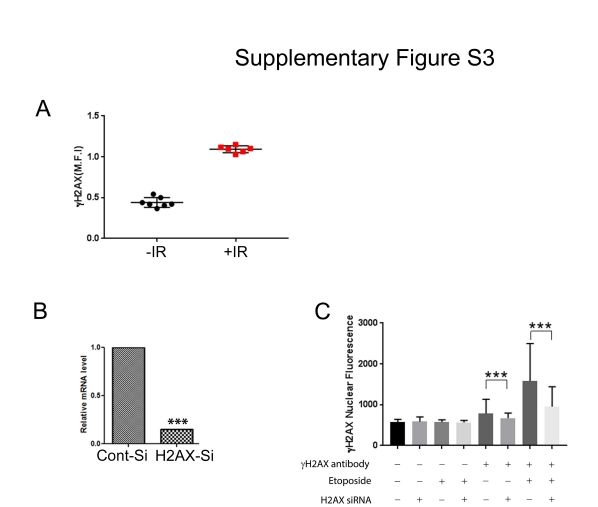
| | Sample ID* | Category | r | p-value |
|----|------------|-----------------|--------|-----------|
| 1 | #1 | Genome-stable | -0.629 | < 0.00001 |
| 2 | #3 | Genome-stable | -0.476 | < 0.00001 |
| 3 | #6 | Genome-stable | -0.576 | < 0.00001 |
| 4 | #7 | Genome-stable | -0.7 | < 0.00001 |
| 5 | #8 | Genome-stable | -0.658 | < 0.00001 |
| 6 | #9 | Genome-unstable | -0.628 | < 0.00001 |
| | | | | |
| 7 | #12 | Genome-unstable | -0.547 | < 0.00001 |
| 8 | #13 | Genome-unstable | -0.653 | < 0.00001 |
| 9 | #14 | Genome-unstable | -0.416 | < 0.00001 |
| 10 | #15 | Genome-unstable | -0.401 | < 0.00001 |

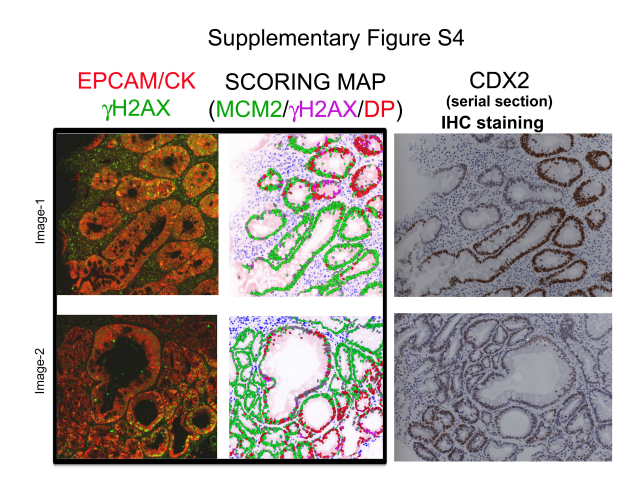
Supplementary Table 9. Correlation analysis between CD44v9 and $\gamma H2AX$ in IM samples

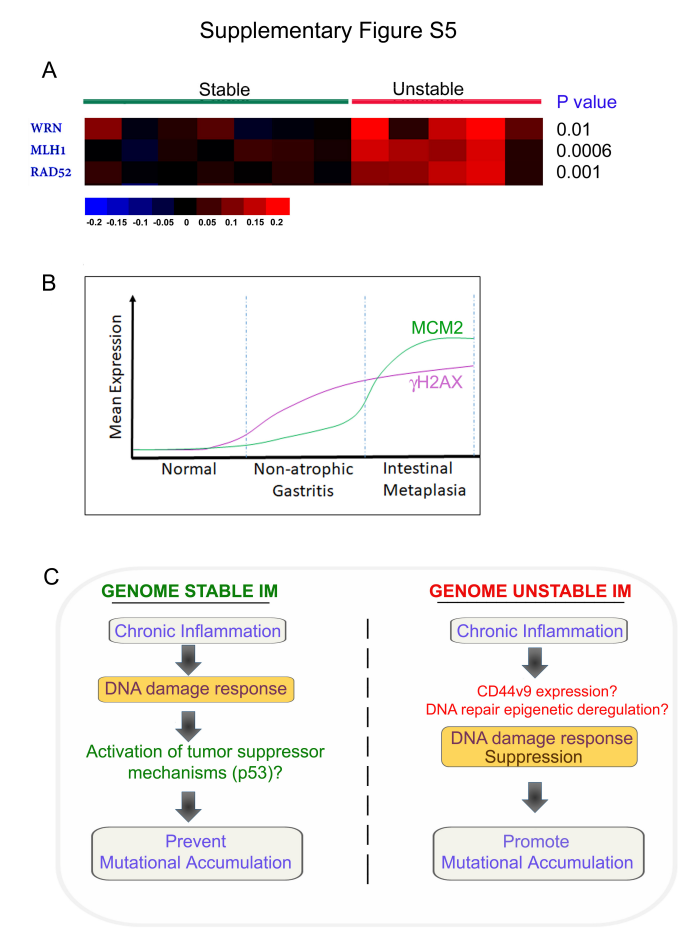
Supplementary Table 10. An integrated view of alterations observed in the genome-unstable IM lesions subjected to genomic and IHC analysis.

| | Sample ID* | % CD44V9+ | Mutation No. | Mutation | CNA segments | Methylation cluster | Hypermethylation |
|---|---------------|--------------|-----------------|------------------------------------|-----------------|---------------------|--|
| | | | | Mutation in DNA repair genes | | | Hypermethylation of DNA repair genes |
| 1 | #9 | 10.79 | 25 | No | 1 | High | Yes |
| 2 | #10 | NA** | 28 | ATM | 0 | High | Yes |
| 3 | #11 | NA** | 51 | FANCA,HLTF, REV3L, POLE | 0 | Intermediate | No |
| 4 | #12 | 13.9 | 14 | No | 3 | High | Yes |
| 5 | #13 | 1.31 | 28 | No | 0 | NA** | NA** |
| 6 | #14 | 34.17 | 38 | No | 1 | High | Yes |
| 7 | #15 | 6.33 | 23 | No | 1 | NA** | NA** |









Gut