Tumor copy-number alterations predict response to immune-checkpoint-blockade in gastrointestinal cancer

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ABSTRACT

Background Despite the great achievements made in immune-checkpoint-blockade (ICB) in cancer therapy, there are no effective predictive biomarkers in gastrointestinal (GI) cancer.

Methods This study included 93 metastatic GI patients treated with ICBs. The first cohort comprising 73 GI cancer patients were randomly assigned into discovery (n=44) and validation (n=29) cohorts. Comprehensive genomic profiling was performed on all samples to determine tumor mutational burden (TMB) and copy-number alterations (CNAs). A subset of samples was collected for RNA immune oncology (IO) panel sequencing, microsatellite instability (MSI)/mismatch repair and program death ligand 1 (PD-L1) expression evaluation. In addition, 20 gastric cancer (GC) patients were recruited as the second validation cohort.

Results In the first cohort of 73 GI cancer patients, a lower burden of CNA was observed in patients with durable clinical benefit (DCB). In both the discovery (n=44) and validation (n=29) subsets, lower burden of CNA was associated with an improved clinical benefit and better overall survival (OS). Efficacy also correlated with a higher TMB. Of note, a combinatorial biomarker of TMB and CNA may better stratify DCB patients from ICB treatment, which was further confirmed in the second validation cohort of 20 GC patients. Finally, patients with lower burden of CNA revealed increased immune signatures in our cohort and The Cancer Genome Atlas data sets as well.

Conclusions Our results suggest that the burden of CNA may have superior predictive value compared with other signatures, including PD-L1, MSI and TMB. The joint biomarker of CNA burden and TMB may better stratify DCB patients, thereby providing a rational choice for GI patients treated with ICBs.

INTRODUCTION

Encouraged by the great achievements in other cancer types, many ongoing clinical trials are assessing immune-checkpoint-blockade (ICB) therapy in gastrointestinal (GI) cancers. However, the efficacy is still not satisfactory.1 Recently, mounting evidence has tried to identify molecular features relevant to immune responses.2 However, data are limited regarding predictive biomarkers in GI cancer patients undergoing ICB treatments.

In particular, microsatellite instability high (MSI-H)/mismatch repair deficient (dMMR) identifies only a small subset (0–5%) of patients with GI cancers.3 4 Efficacy analysis to determine the correlation between tumor mutational burden (TMB) and response to ICBs in GI cancers is still underway due to the high variance in TMB within tumor types.5 6 Most recently, evidence has pointed to the predictive role of copy-number alteration (CNA) and immune-related gene expression profile in cancer patients treated with ICB.8–10 In specific, a lower burden of copy-number loss (CNloss) was observed in responders to ICB treatment in melanoma.8 A combined somatic CNA (SCNA) and TMB score was a candidate biomarker for clinical benefit in patients with melanoma treated with anticytotoxic T-lymphocyte-associated protein 4 (CTLA-4).11 Similarly, SCNA was lower in patients with a partial response (PR) than those with progressive disease (PD)/stable disease (SD) in a cohort of advanced nonsmall cell lung cancer (NSCLC) patients receiving ICB treatment.12 Recently, a negative correlation between CNA and immune parameters including immune cell infiltration and program death ligand 1 (PD-L1) expression was identified in the gastric cancer (GC) and colorectal cancer (CRC) in the The Cancer Genome Atlas (TCGA) data sets.13 However, additional insights are still needed for an in-depth view of the burden of CNAs, including copy-number gain (CNgain) and/or CNloss, in GI cancer patients who received ICB treatments.

Here, we examined tumor specimens from 93 GI cancer patients who were treated with ICBs, including anti-programmed cell death 1 (PD-1), anti-PD-L1, and anti-CTLA-4 antibodies. The predictive and prognostic
significance of tumor CNA burden was extensively investigated, as well as other biomarkers, including TMB, MSI, and PD-L1.

METHODS

Patients and study design

In the first cohort, data of 73 GI cancer patients were retrospectively collected from all GI cancer patients treated with ICBs in the Department of GI Oncology, Peking University Cancer Hospital and Institute from August 1, 2015, to June 8, 2018. To explore genomic correlates of therapeutic efficacy, the 73 GI cancer patients were randomly assigned into the discovery (n=44) and validation (n=29) GI cohorts. Briefly, formalin-fixed paraffin-embedded (FFPE) samples from the 73 patients were subjected to whole-exome sequencing (WES) analysis. To characterize the different immunological features between CNA low and high GI patients, FFPE specimens from a subset of these GI patients (n=65) were subjected to an RNA IO panel sequencing (online supplementary materials).

In addition, 20 patients with advanced GC who were treated with ICBs between January 31, 2018 and May 24, 2019 in the Department of GI Oncology, Peking University Cancer Hospital and Institute were included in this study as an independent GC validation cohort (online supplementary table S1). Aiming to validate the predictive value of CNA burden, we performed WES analysis on these 20 GC samples.

Tumor burden was measured by imaging studies or physical examinations according to the Response Evaluation Criteria in Solid Tumors (RECIST) V.1.1 and iRECIST. Patients were stratified by clinical response. Briefly, the durable clinical benefit (DCB) group was defined as complete response, PR, and SD lasting for ≥24 weeks. No durable benefit (NDB) included patients with PD or SD that lasted <24 weeks.

TMB evaluation

For the determination of TMB value, the number of somatic non-synonymous single nucleotide variants (SNVs) detected using next-generation sequencing was quantified, and the values were extrapolated to the WES analysis using a validated algorithm. TMB was measured in mutations per Mb.

Copy-number analysis

For copy-number analysis, blood cell samples from patients were used as paired controls, and the CONTRA assay was used to call copy-number variations from the FFPE tumor samples for each patient. CNA burden analysis included measurements of the total CNA burden, CNgain, and CNloss. CNgain/CNloss was defined as the total number of genes with CNgain/CNloss per sample, as previously described. The CNA burden was calculated as the total number of genes with CNgains/CNlosses.

RNA IO panel sequencing

RNA IO panel sequencing was conducted as previously described. Briefly, 10 ng of RNA that was extracted from the FFPE sample was reverse transcribed into cDNA, amplified with a primer pool, and ligated to unique barcode adapters. After purification, the libraries were quantified, pooled, and sequenced on the Ion S5 530 chip (Thermo Fisher Scientific, Waltham, MA). The detailed process of gene expression normalization and measurement of normalized reads per million (nRPM) values are described in the online supplementary materials.

TCGA data sets

CNA and gene expression data of stomach adenocarcinoma (STAD) and colon adenocarcinoma (COAD) types analyzed in TCGA project were obtained from the cBioPortal (http://www.cbioportal.org). The CNA burden was calculated as the total number of genes with CNgains or CNlosses. Gene expression levels of the IFN-γ signature and expanded immune signature (gene list was provided in the online supplementary materials) were calculated by using the RNA-Seq data. To quantify the relative infiltration of immune cell types into the tumor microenvironment (TME), single-sample gene set enrichment analysis (ssGSEA) was employed as previously reported.

Statistical analyses

Statistical analyses were performed using SPSS V.22.0 software. Statistical tests included Fisher’s exact tests, Mann-Whitney U tests, and Student’s t-test. All statistical tests were two-sided.

RESULTS

Patient characteristics

This study included 93 metastatic GI patients treated with ICBs between August 1, 2015 and June 8, 2018 (n=73, comprising discovery and first validation GI cohorts, median follow-up time, 11.97 months, IQR: 6.03–17.3) and between January 31, 2018 and May 24, 2019 (n=20, second validation GC cohort, median follow-up time, 4.87 months, IQR: 2.83–10.47). To comprehensively explore the genomic correlates with immunotherapy efficacy, 73 GI patients were randomly assigned into discovery (n=44) and validation (n=29) cohorts. The baseline and treatment characteristics of all patients are shown in online supplementary table S1. Briefly, of 73 included GI patients, the cancer types were GC (50.7%), CRC (27.4%), and others types, including pancreatic neuroendocrine tumors, GI-NETs, and cholangiocellular carcinoma. In GI combined cohort, 47 patients (64.4%) were treated with anti-PD-1 therapy, 15 cases (20.5%) were treated with anti-PD-L1 therapy, and 11 cases (15.1%) received ICB combinational therapy. In the second validation GC cohort (n=20), the cancer types were GC (100%). Among the GC cases, 15 (75%) patients were treated with anti-PD-1 therapy, 3 cases (15%) were treated with anti-PD-L1 therapy, and 2 cases (10%) were treated with ICB combinational therapy (online supplementary table S1).
Figure 1 The burden of tumor copy-number alteration and clinical benefit in the GI cancer cohort. (A) The correlation of burden of copy-number changes (CNA, CNgain and CNloss) and clinical benefit in the discovery (n=44), validation (n=29), and combined GI cancer cohort (n=73). (B) ROC curves for prediction of DCB by CNA burden in the discovery, validation, and combined GI cancer cohort. DCB, durable clinical benefit; GI, gastrointestinal; NDB, no durable benefit; ROC, receiver operating characteristic curve; AUC, area under curve.

Burden of CNA predicts response to ICB in the GI cancer cohort

Tumor somatic copy-number changes have recently been associated with the clinical response to ICB in melanoma. We therefore investigated the predictive value of the three indices: CNA burden, CNgain, and CNloss in the GI cancer cohort (n=73), which was randomly assigned into the discovery and validation cohorts. First, we observed no remarkable differences in the number of copy-number changes among different GI cancer types (online supplementary figure S1 A-C).

Next, we identified that the CNA burden index revealed significantly decreased levels in DCB versus NDB patients receiving immunotherapy in the GI cohort, with AUCs of 0.712 (discovery, p=0.021), 0.864 (validation, p=0.001), and 0.775 (the combined GI cancer patients, p<0.001) (figure 1A,B). Accordingly, a trend toward a lower burden of CNgain or CNloss was also observed in DCB group compared with NDB group with an AUC value of 0.728 (all GI cancer patients, p=0.001) and 0.757 (all GI cancer patients, p=0.0003) for CNgain and CNloss, respectively (online supplementary figure S2 A-B). Notably, the change in CNgain between DCB and NDB patients did not attain statistical significance in the discovery cohort (online supplementary figure S2 A).

Moreover, efficacy was further compared between groups with high (CNA>10) and low (CNA≤10) copy-number variation. Specifically, a drastically improved DCB rate was observed in CNA-low (69%) GI cancers compared with CNA-high (15.9%) cases (online supplementary table S2, p<0.001). A similar trend of responsiveness was identified in the CNgain-low (57.1%) and CNloss-low (57.5%) groups when compared with the CNgain-high (18.4%) and CNloss-high (12.1%) cases, respectively (p<0.01 for all comparisons, online supplementary table S2). Taken together, our findings strongly indicate that features for tumor genome alterations, including total CNA, CNgain, and CNloss, are potential predictive biomarkers for GI cancers with ICB treatment. Notably, the CNA burden demonstrated the highest AUC value, and therefore, the utility of the CNA burden was further evaluated.
**Tumor mutational burden (TMB) and clinical benefit.** (A) The correlation of TMB (number of non-synonymous mutations per Mb) and clinical benefit in the discovery, validation, and combined GI cancer cohort. (B) ROC curves for prediction of DCB by TMB in the discovery, validation, and combined GI cancer cohort. GI, gastrointestinal.

The evaluation of predictive value of MSI/MMR and PD-L1 in GI cancers

Since MSI/MMR and PD-L1 expression have emerged as potential predictive biomarkers for PD-1/PD-L1 blockade, we examined the correlation between MSI/MMR or PD-L1 and the clinical benefit in GI cancer patients. Our data show that MSI-H/dMMR patients experienced a significantly higher DCB rate (59.1%) than MSI-L/MSS/pMMR patients (28.6%, p=0.022), while no association was found between PD-L1 positivity and the efficacy of ICB therapy in our cohort (online supplementary table S2).

Analysis of the predictive value of TMB in GI cancers

As shown in online supplementary figure S1D, higher TMB was identified in CRC (median 38.7 mutations/Mb) than other cancer types (median 3.5 mutations/Mb). The median number of TMB in DCB patients was significantly higher than that of NDB patients, with AUCs of 0.696 (p=0.032) and 0.702 (p=0.004) in the discovery and combined cohorts, respectively (figure 2A,B). On the other hand, we did not observe a statistically significant AUC value (0.697) of TMB in the validation cohort (p=0.080) (figure 2B). In all GI cancer patients, 60% (18/30) of the patients with a higher mutation burden (designated as above 5, including intermediate and high TMB levels) experienced a DCB compared with 20.9% (9/43) of those with a lower TMB (online supplementary table S2, p=0.001).

Clinical outcomes stratified by baseline CNA, TMB, MSI/MMR and PD-L1 status in GI cancers

To assess whether the above-tested signatures, including CNA, TMB, MSI/MMR and PD-L1 status, influence the OS and progression-free survival (PFS), we used Kaplan-Meier analysis for the GI cancer cohort (figures 3 and 4 and online supplementary figure S3-4). As expected, CNA-low patients demonstrated a longer median OS (unreached for all three cohorts) than the CNA-high group (discovery, 7.63 months; validation, 5.6 months; and combined cohorts, 7.43 months) (figure 3A, log-rank test, p<0.05 for all comparisons). A similar trend was identified between CNgain/CNloss and OS in GI cancer patients, although statistical significance was not attained for CNgain in the validation cohort (online supplementary figure S3). In addition, patients with higher TMB levels revealed longer OS than patients with lower TMB in the validation and combined cohorts (figure 3B). In our cohort, neither MSI/MMR nor PD-L1 status was able to predict the OS outcome after ICB (figure 3C,D).
Consistently, longer PFS was identified in the lower burden groups for CNA, CNgain or CNloss than in the higher burden groups (figure 4A and online supplementary figure S4A–B, log-rank test, p<0.05 for all comparisons, except for CNgain in the validation cohort). In contrast, higher TMB levels were able to stratify patients with favorable PFS in all three cohorts (figure 4B). Moreover, the median PFS time of MSI-H/dMMR patients (7.24 months) was significantly longer than that of MSI-L/MSS/pMMR patients (2.67 months, p=0.0173) (figure 4C). However, no significant association was identified between PD-L1 status and PFS in the combined cohort (figure 4D).

**Joint utility of burden of CNA with TMB**

Due to the predictive and prognostic value of CNA and TMB, we sought to explore the possibility of combining the two parameters in identifying DCB patients receiving immunotherapy. As revealed in figure 5A, no significant correlation was observed between CNA burden and TMB,
Figure 4  Association between tumor CNA burden, TMB level, PD-L1, and MMR/MSI status with the progression-free survival (PFS) of patients with GI cancer receiving ICB treatment. (A) Kaplan-Meier estimates of PFS according to tumor CNA burden in the discovery cohort, the validation cohort, and the combined GI cancer cohort. (B) Kaplan-Meier estimates of PFS according to TMB levels in the discovery cohort, the validation cohort, and the combined GI cancer cohort. (C) Kaplan-Meier estimates of PFS according to MSI/MMR status in the combined GI cancer cohort. (D) Kaplan-Meier estimates of PFS according to PD-L1 expression in the combined GI cancer cohort. CNA, copy-number alterations; dMMR, mismatch repair deficient; GI, gastrointestinal; ICB, immune-checkpoint-blockade; MMR, mismatch repair; MSI-H, microsatellite instability high; PD-L1, program death ligand 1; TMB, tumor mutational burden.
The association of a joint tumor CNA burden/TMB with clinical outcome of patients with GI cancer receiving ICB treatment. (A) Correlation between the burden of CNA and TMB in the GI cancer cohort. (B) Relationships of both CNA burden and TMB signatures with clinical benefit (DCB, blue dot; NDB, black dot). (C) Proportions of patients with DCB calculated within each of the four indicated subgroups. (D) Overall survival (OS, Kaplan-Meier curves) of patients within each of the four indicated subgroups in combined GI cancer cohort. (E) Waterfall plot of tumor response to ICB according to the CNA burden (CNA-high, red bar; CNA-low, green bar) and TMB (asterisk). The Y-axis represents the percentage of maximum tumor reduction assessed according to the RECIST V.1.1 criteria. CNA, copy-number alterations; DCB, durable clinical benefit; GI, gastrointestinal; ICB, immune-checkpoint-blockade; NDB, no durable benefit; RECIST, Response Evaluation Criteria in Solid Tumors; TMB, tumor mutational burden.

Intriguingly, a high proportion of DCB rate was also identified in the TMB-high/CNA-low subgroup of patients (five of six), as presented in online supplementary figure S5C-D. A larger sample size may help improve the performance of the predictive model.

Lower CNA burden correlates with activated immune responses

Recent analyses have linked cancer genomic features, including TMB and CNA, with antitumor immunity. In particular, studies have proposed that high mutational load and low aneuploidy may correlate with increased T-cell
The correlation of the CNA burden and the immune-related RNA signatures. (A) Immune RNA signature levels in CNA-low and CNA-high subgroups in our combined GI cancer cohort. (B) Upregulated gene clusters in CNA-low vs CNA-high subgroups in our combined GI cancer cohort. (C) Immune RNA signature levels in CNA-low and CNA-high subgroups in the TCGA STAD and COAD cohorts. CNA-low, tumors with a CNA burden level within the 25th percentile; CNA-high, tumors with a CNA burden level higher than the 75th percentile. CNA, copy-number alterations; COAD, colon adenocarcinoma; GI, gastrointestinal; STAD, stomach adenocarcinoma; TCGA, The Cancer Genome Atlas.

We therefore sought to determine the association between CNA burden and the inflammatory characteristics of the TME by using an RNA IO sequencing platform. IFN-γ-related (6-gene set) and expanded immune signature (18-gene set) mRNA profiles have recently been developed as pan-cancer biomarkers to predict the immune response to ICB in melanoma, head and neck squamous cell carcinoma and GC cohorts. Herein, we measured the IFN-γ and expanded immune signatures in 65 patient samples with an available RNA data set. Briefly, the CNA burden was partitioned into high or low at the upper quartile or lower quartile value for the 65 GI patients. As figure 6A demonstrates, the CNA-low subgroup revealed higher IFN-γ and expanded immune signatures than the CNA-high subgroup (p<0.05). Moreover, we observed 26 upregulated genes in the CNA-low samples versus the CNA-high group (figure 6B). These upregulated immune-related genes belong to the following pathways: lymphocyte markers, interferon signaling, lymphocyte regulation, and checkpoint pathways.

To confirm the correlation between tumor CNA burden and immune signature more broadly, we examined the IFN-γ and expanded immune signatures in the TCGA, STAD, and COAD cohorts. For each cohort, the burden of CNA was measured and then partitioned into high or low at the upper quartile or lower quartile value. Consistently, CNA-low samples showed higher immune signatures in both cohorts (figure 6C). Further ssGSEA analysis revealed that CNA-low GC and CRC samples were infiltrated with diverse immune cell types, including activated CD8+ T cells, activated CD4+ T cells, natural killer (NK) cells, and NK T cells (online supplementary figure S6A-B). Collectively, our
data indicate that a lower CNA burden may correlate with an activated inflammatory response in the TME.

**DISCUSSION**

In this study, we used WES analysis to characterize genomic determinants of the clinical benefit from ICB in GI cancers. Here, we show that the CNA burden may have superior predictive value compared with other signatures, including PD-L1, MSI, and TMB.

PD-L1 and MSI/MMR status have been suggested to be independent predictive biomarkers for ICB, and determining MSI/MMR and PD-L1 status of patients based on immunohistochemistry becomes a cost-effective screening tool.25,26 In our study, MSI-H/dMMR GI patients showed a significantly increased DCB rate (59.1%) versus patients with MSS/MSI-/pMMR status (28.6%) (online supplementary table S2, p=0.022). A larger sample size may increase the predictive value of MSI-H status; however, the overall incidence (0–5%)3–5 was still too low, and even half of MSI-H/dMMR GI cancers are non-responsive to ICB treatment.27,28 On the other hand, we observed that PD-L1 positivity showed little value for stratification of DCB patients with GI cancers (online supplementary table S2). Similarly, in the combination arm of KEYNOTE-062 study, KEYTRUDA plus chemotherapy was not found to be superior for OS (CPS≥1 or CPS≥10) or PFS (CPS≥1) compared with chemotherapy alone.29 The result confirmed our findings that PD-L1 expression is not a predictive biomarker in advanced gastric or gastroesophageal junction adenocarcinoma. Therefore, we recruited the second validation cohort including 20 GC patients for further investigation.

TMB was recently elevated as a powerful predictive marker for immunotherapy in various cancer types.17,23,30,31 Here, we identified that a relatively higher level of TMB (>5 mutations/Mb) in GI cancers correlated with improved DCB (60%) compared with lower TMB (≤3 mutations/Mb, 20.9%) (online supplementary table S2). We also examined the landscape of TMB across different GI cancer types, revealing that CRC had significantly higher TMB than other malignancies (online supplementary figure S1D). This may result from the high percentage of MSI-H/dMMR CRC patients enrolled in the ICB treatment, and MSI-H/dMMR tumors usually exhibit high TMB, especially in CRC.17,32 We can reasonably hypothesize that the predictive power of TMB can partially be attributed to MSI-H/dMMR CRC. Consequently, the best cut-off values for TMB across different GI cancer types still need to be thoroughly explored in larger cohorts in the future. While growing evidence revealed a significant association between TMB and therapeutic efficacy in multiple cancers receiving ICIs, the utility of TMB still remains controversial.4,24 For instance, TMB may function as an effective predictor of response in anti-PD-1 monotherapy in PD-L1-positive lung cancer.25 In contrast, in metastatic NSCLC patients receiving pembrolizumab plus platinum-based chemotherapy, TMB did not show robust predictive value.34

Most importantly, WES analysis demonstrated a strong correlation between CNA burden and DCB from ICB (figure 1 and online supplementary figure S5A) and that a lower CNA burden was also associated with favorable OS and PFS (figures 3A and 4A). A similar trend was identified in the other two indices, CNgain and CNloss, although statistical significance was not attained in some cohorts (online supplementary figure S2-4). This finding is in accordance with a previous investigation which reported that a higher burden of CNloss is associated with persistent resistance in melanoma patients with sequential CTLA-4 and PD-1 blockades. However, the study did not report significant differences in the CNA burden, CNgain or CNloss in the context of each individual agent response.8 Thus, the association between copy-number variations and immune response may vary among different cancer types. Our data strongly indicate that the CNA burden may function as a potential predictive and prognostic biomarker in GI cancers.

Recently, CNA burden has also been correlated with clinical outcome in multiple cancer types.35,36 In addition, great efforts have been made to understand how copy-number changes affect the characteristics of cancer and/or the microenvironment.37–40 For instance, in hepatocellular carcinoma, a subgroup of patients defined as immune class with inflammatory response showed a lower burden of CNgains and CNlosses;41 similarly, pancreatic ductal adenocarcinoma (PDA) with high cytolytic activity exhibited fewer copy-number changes at loci important in PDA, including NOTCH2, MYC, and FGFR1.42 A comprehensive analysis of genetic and immunologic characteristics based on the TCGA data set demonstrated a negative correlation between CNA load and immune parameters in COAD and STAD.15 This finding was in accordance with our observations that CNA-low samples exhibited higher IFNγ and expanded immune signatures than the CNA-high subgroup (figure 6). Additionally, ssGSEA analysis further proved that CNA-low GC and CRC samples were infiltrated with activated CD8+ T cells, activated CD4+ T cells, and NK cells (online supplementary figure S6). Taken together, these results indicate somatic CNA levels might be a strong biomarker for cytotoxic immune cell infiltration and hence predict ICB efficacy.

Finally, we also evaluated the joint utility of CNA burden and TMB (figure 5, and online supplementary figure S5). Intriguingly, a limited DCB (1/28) and OS (6.23 months) from ICB was observed in the TMB-low/CNA-high subgroup, whereas the highest DCB rate (12/14) and longest OS (unreached) occurred in the TMB-high/CNA-low subgroup. It has been proposed that higher TMB levels may increase the chances of generating immunogenic neoantigens.43 On the other hand, SCNA load, which has been identified as a significant survival predictor independent of TMB in several cancer types,8,44 revealed a significant correlation with immunological features. Specifically, lower levels of CNAs showed

our study has limitations. First, since this cohort was relatively heterogeneous, the proposed signature may vary across GI cancer types or checkpoint inhibitor regimens. Second, this is a retrospective single-center study. Further prospective study within specific cancer type is warranted in the future. In recent years, cost-effective sequencing panels have been developed to accurately measure TMB levels. However, data on the parameters influencing panel-based CNA burden quantification are limited. Finding a refined set of panel tools that can accurately estimate both CNA and mutational burden will become promising for clinical application.

Our data indicate that lower CNA burden correlates with better clinical outcome. Moreover, a joint biomarker of CNA burden and TMB may better stratify DCB patients with GI cancers receiving ICB treatment. Further prospective studies are needed to validate this observation across multiple GI cancer types.

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