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Alcohol consumption, DNA methylation and colorectal cancer risk: Results from pooled cohort studies and Mendelian randomization analysis

Xuan Zhou ¹	Lijuan Wang ¹ Jiarui Xiao ¹ Jing Sun ¹ Lili Yu ¹
Han Zhang ²	Xiangrui Meng ³ Shuai Yuan ⁴ Maria Timofeeva ^{5,6}
Philip J. Law ⁷ 📵	Richard S. Houlston ⁷ Kefeng Ding ⁸ Malcolm G. Dunlop ⁶
Evropi Theodora	atou ^{6,9} Xue Li ¹

Correspondence

Xue Li, Department of Big Data in Health Science School of Public Health, and Center of Clinical Big Data and Analytics of The Second Affiliated Hospital, Zhejiang University School of Medicine, Hangzhou, Zhejiang, China. Email: xueli157@zju.edu.cn

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Abstract

Alcohol consumption is thought to be one of the modifiable risk factors for colorectal cancer (CRC). However, the causality and mechanisms by which alcohol exerts its carcinogenic effect are unclear. We evaluated the association between alcohol consumption and CRC risk by analyzing data from 32 cohort studies and conducted two-sample Mendelian randomization (MR) analysis to examine for casual relationship. To explore the effect of alcohol related DNA methylation on CRC risk, we performed an epigenetic MR analysis with data from an epigenome-wide association study (EWAS). We additionally performed gene-alcohol interaction analysis nested in the UK Biobank to assess effect modification between alcohol consumption and susceptibility genes. We discovered distinct effects of alcohol on CRC incidence and mortality from the meta-analyses, and

Abbreviations: ALSPAC, Avon Longitudinal Study of Parents and Children; ARIES, Accessible Resource for Integrated Epigenomic Studies; AUD, alcohol use disorder; CI, confidence interval; CRC, colorectal cancer; EWAS, epigenome-wide association study; GSCAN, GWAS and Sequencing Consortium of Alcohol and Nicotine use; GWAS, genome-wide association study; HR, hazard ratio; IVs, genetic instruments; IVW, inverse variance weighted; mQTLs, methylation quantitative trait loci; MR, Mendelian randomization; MR-PRESSO, MR pleiotropy residual sum and outlier; OR, odds ratio; PAU, problematic alcohol use; SNPs, single nucleotide polymorphisms.

Xuan Zhou, Lijuan Wang and Jiarui Xiao should be considered joint first authors.

Evropi Theodoratou and Xue Li should be considered joint last authors.

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¹Department of Big Data in Health Science School of Public Health, and Center of Clinical Big Data and Analytics of The Second Affiliated Hospital, Zhejiang University School of Medicine, Hangzhou, Zhejiang, China

²College of Public Health, Zhengzhou University, Zhengzhou, Henan, China

³Division of Psychiatry, University College of London, London, UK

⁴Unit of Cardiovascular and Nutritional Epidemiology, Institute of Environmental Medicine, Karolinska Institutet, Stockholm, Sweden

⁵Danish Institute for Advanced Study (DIAS), Epidemiology, Biostatistics and Biodemography Research Unit, Institute of Public Health, University of Southern Denmark, Odense, Denmark

⁶Cancer Research UK Edinburgh Centre, Medical Research Council Institute of Genetics and Cancer, University of Edinburgh, Edinburgh, UK

⁷Division of Genetics and Epidemiology, The Institute of Cancer Research, London, UK

⁸Department of Colorectal Surgery and Oncology, Key Laboratory of Cancer Prevention and Intervention, Ministry of Education, The Second Affiliated Hospital, Zhejiang University, Hangzhou, Zhejiang, China

⁹Centre for Global Health, Usher Institute, University of Edinburgh, Edinburgh, UK

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genetic predisposition to alcohol drinking was causally associated with an increased CRC risk (OR = 1.79, 95% Cl: 1.23-2.61) using two-sample MR approaches. In epigenetic MR analysis, two alcohol-related CpG sites (cg05593667 and cg10045354 mapped to COLCA1/COLCA2 gene) were identified causally associated with an increased CRC risk ($P < 8.20 \times 10^{-4}$). Gene-alcohol interaction analysis revealed that carriage of the risk allele of the eQTL (rs3087967) and mQTL (rs11213823) polymorphism of COLCA1/COLCA2 would interact with alcohol consumption to increase CRC risk ($P_{Interaction} = .027$ and $P_{Interaction} = .016$). Our study provides comprehensive evidence to elucidate the role of alcohol in CRC and highlights that the pathogenic effect of alcohol on CRC could be partly attributed to DNA methylation by regulating the expression of COLCA1/COLCA2 gene.

KEYWORDS

alcohol, colorectal cancer, DNA methylation, Mendelian randomization

What's new?

While alcohol consumption is suspected of being a modifiable risk factor for colorectal cancer (CRC), causal relationships and carcinogenic mechanisms linked to alcohol consumption remain unclear. In this study, investigation of interactions between alcohol consumption and susceptibility genes revealed associations between genetic predisposition to alcohol drinking and increased CRC risk. Epigenetic analyses showed that the pathogenic effect of alcohol can be attributed in part to DNA methylation via regulation of the expression of the COLCA1/COLCA2 gene. Polymorphisms in risk alleles in COLCA1/COLCA2 interacted with alcohol consumption to increase CRC risk, providing insight into how alcohol modulates CRC tumorigenesis.

1 | INTRODUCTION

Colorectal cancer (CRC) is the third most commonly diagnosed cancer worldwide, and the second leading cause of cancer-related death worldwide, with more than 1.9 million new cases and 935 000 deaths estimated to occur in 2020.1 Alcohol drinking is thought to be one of the modifiable risk factors of CRC.² The World Cancer Research Fund and the American Institute for Cancer Research concluded that alcohol intake more than 30 g of ethanol per day (about two drinks a day) is associated with increased risk of CRC, however, the association between light and moderate levels of alcohol consumption and CRC risk are unclear. A pooled analysis identified a J-shaped association between alcohol drinking and CRC risk, where moderate alcohol consumption showed a protective effect comparing to nondrinkers,3 while other studies reported a positive dose-response relationship or nonsignificant positive association at moderate levels of alcohol consumption.^{4,5} Additionally, it is uncertain whether the effect of alcohol consumption on cancer risk is different between colon and rectum. One study reported a stronger alcohol-cancer risk association for the colon, whereas others discovered a stronger or similar association for the rectum.⁷⁻⁹ Overall, the observational association between alcohol and CRC risk is inconsistent and these discrepancies could be due to confounding, reverse causation, and other biases inherited from conventional epidemiological studies.

Mendelian randomization (MR) is a method commonly applied in epidemiology to estimate the causal relationship between a modifiable

environmental exposure and a medical relevant trait or disease by utilizing genetic variants as instrumental variables (IVs).¹⁰ Recent meta-analyses of genome-wide association studies (GWAS) identified a number of single nucleotide polymorphisms (SNPs) associated with general alcohol consumption (drinks per week),¹¹ and the sub-phenotypes of more severe drinking behaviors and physiological alcohol dependence (alcohol use disorder and problematic alcohol use),¹² which can be selected as IVs to proxy the genetic predisposition of different alcohol drinking behaviors in MR analyses.

Beyond causality, the underlying mechanisms are not fully understood. Epigenome-wide association studies (EWAS) showed that alcohol consumption can affect DNA methylation both in blood and tissues, ^{13,14} and aberrant patterns of DNA methylation, an important epigenetic mechanism of transcriptional control, can also lead to CRC development. ¹⁵ It has been reported that alcohol drinking could also interact with the genetic polymorphism of several susceptibility genes in the tumorigenesis of CRC. ^{16,17} Therefore, we hypothesize that alcohol related DNA methylation and gene-alcohol interaction might be involved in the pathological effect of alcohol drinking on CRC risk.

In our study, we comprehensively evaluated the relationship between alcohol consumption and CRC risk by performing an updated meta-analysis of prospective cohort studies and conducting two-sample Mendelian randomization (MR) analyses for causal inference. To shed light on the underlying biology, we explored the causal effects of alcohol related DNA methylation on the risk of CRC and performed gene-alcohol interaction analysis nested in the UK Biobank cohort.

2 MATERIALS AND METHODS

2.1 Meta-analysis of prospective cohort studies

2.1.1 Literature search

We performed a comprehensive literature search for cohort studies in MEDLINE, EMBASE and BIOSIS citation index, using "alcohol", "CRC" and "cohort" as the keywords. The detailed search strategy can be seen in Supporting Information Methods. Reference lists of retrieved reviews and meta-analyses were manually searched to identify additional studies of interest.

Studies were included if they met the following criteria: (i) examined the relationship between alcohol consumption and CRC incidence or mortality; (ii) reported odds ratio (OR), relative risk (RR) or hazard ratio (HR) (or providing sufficient data to compute them) for at least two levels of alcohol consumption vs nondrinkers and/or occasional drinkers (<1 drink/week): (iii) reported standard errors or confidence intervals (CIs) of the risk estimates (or providing sufficient data to calculate them). Studies were excluded if they met any of the following criteria: (i) not prospective cohort studies; (ii) included individuals with prior CRC history; (iii) did not report the independent effect of alcohol on CRC risk; (iv) reported the risk estimates of a specific type of alcoholic beverage only; (v) when more than one study was published for the same cohort, only the most recent and comprehensive one was included and the others were excluded.

2.2 Data extraction and quality assessment

The retrieved articles were independently reviewed by two authors to determine the eligibility for inclusion in the meta-analysis. Doubts and disagreements were resolved by consensus among all the investigators. For each included study, two authors independently extracted the following items: first author, publication year, country, study population, the number (or person-years) of cases and subjects at risk, age, gender, anatomical subsite, exposure levels and corresponding risk estimates with 95% CIs, outcome, adjustment variables, and duration of follow-up. As the measurement scale of alcohol was different among included studies, we set grams per day as a standard, considering one drink as 12.5 g, 1 mL as 0.8 g, and 1 oz as 28 g ethanol, unless otherwise specified in original studies.4 If alcohol consumption was reported by a range, the midpoint was taken as the exposure dose; and for an open-ended upper category, the dose was assigned as the lower bound plus three-quarters of the length of the previous category. 4,18 We divided alcohol consumption into three categories, considering ≤12.5, ≤50 and >50 g/day as light, moderate and heavy drinking, respectively. When a study reported two or more risk estimates for a single dose category, we used the method developed by Hamling et al to combine them into one single estimate. 19 The Newcastle-Ottawa Scale (NOS) was used to evaluate the methodological quality of included studies.²⁰ The scale has a score system (ranging from 0 to 9 points) to assess three domains: selection,

comparability, and outcome. In the comparability domain, we set age, sex, and BMI as the most important covariates that should be adjusted, and in the outcome section, we regarded more than 5 years follow-up as enough for outcomes to occur and follow-up rate larger than 80% as adequacy. We considered the overall score 0-5, 6-7, and 8-9 as low, moderate, and high quality, respectively.

2.3 Statistical analysis

We computed a pooled RR for each exposure category compared to nondrinkers and/or occasional drinkers using the inverse variance weighted random-effects model. Statistical heterogeneity across included studies was evaluated by the I2 index. We conducted metaanalysis for CRC incidence (the risk to develop CRC) and CRC-specific mortality (the risk of death due to CRC) separately; subgroup analyses for different sex, anatomical site of tumor and study population were also performed. For the dose-response analysis, we assigned the transformed exposure levels as the doses and their corresponding RR or HR estimates as responses, using cubic-spline model with 0, 12.5, and 50 g per day alcohol drinking as knots to estimate the doseresponse trend, and the adjusted R-square to quantify the model fit.

Genetic instruments for predisposition to 2.4 alcohol drinking

We derived genetic instruments for predisposition to general alcohol drinking (drinks per week) from a GWAS conducted by the GWAS and Sequencing Consortium of Alcohol and Nicotine use (GSCAN) in 1.2 million individuals of European ancestry. 11 Data of drinks per week of all participants were collected using a questionnaire, left-anchored at 1 and log-transformed (natural log). 11 For each SNP that passed quality control, linear regression was applied to calculate the effect sizes of the exposure. 11 A total of 99 SNPs of genome-wide significance $(P < 5 \times 10^{-8})$ were identified to be associated with the phenotype, and 84 SNPs remained after removing those in linkage disequilibrium $(r^2 > .01)$. To complement with more severe drinking behaviors and alcohol dependence, genetic instruments for alcohol use disorder (AUD) and problematic alcohol use (PAU) were also used. A total of 30 SNPs and 42 SNPs at genome-wide significance (P < 5×10^{-8}) were identified for AUD and PAU respectively from a GWAS metaanalysis containing 435 563 individuals of European-ancestry (Million Veteran Program, Psychiatric Genomics Consortium, and UK Biobank),¹² and 19 and 26 SNPs remained after clumping those in linkage disequilibrium ($r^2 > .01$).

Methylation quantitative trait loci (mQTL) for alcohol-related DNA methylation

We obtained the DNA methylation pattern related to alcohol consumption from an epigenome-wide association study with meta-analysis of 13 population-based cohorts using whole blood DNA.²¹ Illumina Infinium HumanMethylation450 (HM450) BeadChip was used to measure the level of methylation and the association between alcohol consumption and methylation level of epigenome-wide CpG sites was adjusted for multiple covariates, such as age, sex, BMI, batch effects and white cell blood counts (ie, CD4 cells, CD8 cells, natural killer cells, B cells and monocytes).²¹ A total of 363 CpG sites were identified to be associated with alcohol consumption at epigenome-wide significance level (P<1 \times 10⁻⁷). Then, we derived genetic variants robustly associated with those CpG sites from an mQTL database based on the Accessible Resource for Integrated Epigenomic Studies (ARIES) project.²² The ARIES was launched using the Illumina Infinium HumanMethylation450 (HM450) BeadChip to acquire epigenetic data (CpG sites) and the Illumina Infinium Human Hap550 and 660-w quad genome-wide SNP genotyping platform to acquire genetic data (SNPs) on cord blood and peripheral blood samples from 1018 mother-offspring pairs in the Avon Longitudinal Study of Parents and Children (ALSPAC) cohort.²³⁻²⁵ The Matrix eQTL software was used for the preliminary association analysis of SNPs with CpG sites.²² SNPs at epigenome-wide association significance ($P < 1 \times 10^{-7}$) were further analyzed using exact linear regression including covariates in PLINK 1.07, and genome-wide complex trait analysis (GCTA) was performed to obtain the most representative independent loci in relation to each CpG site.²² For each of those CpG sites associated with alcohol consumption, we identified mQTL-DNA methylation effect estimates at middle-age time point for the methylation MR analysis.

2.6 | GWAS summary-level data of CRC

We investigated the relationship between the selected genetic IVs and CRC risk using summary data from 12 previously reported CRC GWASs. Briefly, these GWASs included individuals of European ancestry from the following studies: CCRR1, CCFR2, COIN, CORSA, Croatia, DACHS, FIN, NSCCG-OncoArray, SCOT, UK1, VQ58, and Scottish case-control series, comprising 20 049 cases and 22 661 controls. Comprehensive details of the cases and controls are available in our previously published work. After standard quality control procedures, a total of 16 871 cases and 26 328 controls were included in the meta-GWAS analysis.

2.7 | Two-sample MR

To test the causality between genetic predisposition to alcohol consumption and CRC risk, drinks per week, AUD, and PAU were considered as the alcohol exposures, and 84, 19, and 26 SNPs were used as the IVs. To investigate the causal effect of alcohol related DNA methylation on the risk of CRC, methylation at each CpG site induced by alcohol was regarded as the exposure and its proxy mQTLs were used as the IVs. For each genetic instrument, $\beta_{\rm GP}$ is the estimate of the genetic association with the exposure (ie, alcohol consumption or DNA methylation) and $\beta_{\rm GD}$ is the estimate of the genetic association

with the outcome (ie, CRC). We calculated the effect estimates in CRC risk per SD (SD) increase in alcohol consumption (or DNA methylation) using the formula β_{GD}/β_{GP} (Wald ratio) and combined in a fixed-effect meta-analysis after weighing each ratio estimate using the inverse variance weighted (IVW) approach. We additionally undertook multiple analyses to assess the risk of horizontal pleiotropy (a violation of the second MR assumption). First, we used the MR Egger method, which allows for an additional intercept (alpha) term and provides an estimate of directional horizontal pleiotropy.²⁷ Then, we used three additional MR methods known to be more robust to the presence of horizontal pleiotropy (but at the expense of statistical power): simple mode, weighted median, and weighted mode.²⁸ Given possible instability in MR estimates, we applied the global test, outlier test, and distortion test using the MR pleiotropy residual sum and outlier (MR-PRESSO) R package as an additional control for pleiotropy. 29 To control for potential pleiotropic effect on confounders, we queried PhenoScanner³⁰ to identify all reported association for the IVs and performed sensitivity analyses by excluding those associated with obesity related traits (ie, BMI, waist circumference, hip circumference, and weight), physical activity, smoking, education. Crohn's disease and inflammatory bowel disease at the threshold of 5×10^{-8} in European ancestry samples. MR analyses were performed using the TwoSampleMR R package.31

While evaluating the causal effect of DNA methylation at alcohol-drinking related CpG sites, the effect allele of mQTL was unified to be in the same direction with the effect of alcohol intake on DNA methylation. If there was only one mQTL for a CpG site, only the Wald ratio with its corresponding SE was calculated, but if it was associated with multiple independent mQTLs, the aforementioned IVW approach was applied. Bonferroni correction was applied to account for multiple testing.

2.8 | Gene-alcohol interaction analysis nested in the UK Biobank

UK Biobank is a large cohort study that recruited more than half a million people aged 40 to 69 years across the UK during 2006 and 2010. The details of baseline information and biological samples collection can be found elsewhere.³² To validate the MR estimates and test whether the effect of alcohol-related methylation on CRC was distinct by the expression and methylation levels of COLCA1/COLCA2 gene, we obtained the genotyping data of the eQTL SNP rs3087967³³ and the mQTL SNP rs11213823 of the COLCA1/COLCA2 gene along with alcohol drinking information and performed gene-alcohol interaction analysis among incident CRC cases and population-based controls nested in the UK Biobank. We excluded former drinkers and finally included 1892 incident CRC cases and 9386 matched controls for the analysis after quality control. Various types of alcoholic beverages were transformed into gram per day of ethanol and the amount of consumption was divided into three categories (light, moderate and heavy) as mentioned before. Then we investigated the independent estimates of alcohol drinking (per 10 g/d) using the R function "glm", the independent estimates of rs3087967 and rs11213823 and the

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interaction estimates using R function "snp.logistic" in CGEN R package³⁴ with two regression models under the framework of a nested case-control study. We basically adjusted for age and sex of the participants, and additionally adjusted for area deprivation index, red meat consumption, processed meat consumption, aspirin intake, BMI and smoking for the multivariable model. Based on the genotypes of rs3087967 and rs11213823, we further conducted stratification analyses to assess the dose-response effects of alcohol drinking (per 10 g/d). Time-to-event analysis was performed in CRC incident cases to test if the time to CRC occurrence was different among nondrinkers, light, moderate and heavy drinkers stratified by the genotype of rs3087967 and rs11213823. The study design is shown in Figure 1. All analyses were conducted using R version 4.1.0.

RESULTS 3

3.1 Meta-analysis of observational studies

A total of 3407 articles were retrieved from MEDLINE, EMBASE and BIOSIS citation index databases. After screening of title, abstract and full text, 32 unique prospective cohort studies were eligible to be included in the meta-analysis. More details regarding the process of parallel screening can be found in Figure S1. Of them, 25 (78.1%) reported incidence and seven (21.9%) reported mortality as the outcome of interest; 11 (34.4%) were conducted in North America, nine (28.1%) in Europe, 10 (31.3%) in Asia, two (6.3%) in mixed or other areas; five (15.6%) reported estimates for men, two (6.3%) for women, 15 (46.8%) for both men and women in subgroups and 10 (31.2%) for both men and women together: three (9.4%) reported the risk of colon cancer only, 11 (34.4%) reported the risk of colon cancer and rectum cancer separately, and 18 (56.3%) reported the incorporative risk of colorectal cancer; 22 (68.8%) included abstainers whereas

eight (25.0%) included occasional drinkers in the reference category (Table S1). The details of all included studies can be seen in Table S2. Quality assessment based on the NOS scale suggested 28 studies (87.5%) as high quality and four (12.5%) as moderate quality (Table S3).

Figure S2 presents the pooled RR estimates for light, moderate and heavy drinking as compared to nondrinkers and occasional drinkers in the risk of CRC incidence and mortality as well as several subgroups stratified by sex, histologic subtypes and main population. No association were observed between light drinking and CRC risk. Significant associations were identified for CRC incidence (RR = 1.24, 95% CI: 1.17-1.32, $P = 5.27 \times 10^{-12}$ for moderate drinking, RR = 1.54, 95% CI: 1.15-2.06. P = .004 for heavy drinking) but not for CRC mortality (RR = 0.98, 95% CI: 0.86-1.12, P = .789 for moderate drinking, RR = 1.16, 95% CI: 0.85-1.57, P = .359 for heavy drinking). The association between alcohol drinking and CRC was identified in men (RR = 1.24, 95% CI: 1.15-1.33, $P = 7.26 \times 10^{-9}$ for moderate drinking, and RR = 1.45, 95% CI: 1.16-1.80, P = .001 for heavy drinking), while no significant detrimental effect of drinking on CRC risk was observed in women drinkers. Considering colon cancer and rectum cancer separately, moderate and heavy drinking were associated with higher risk (RR = 1.25, 95% CI: 1.14-1.38, P = 5.45×10^{-6} for moderate drinking and RR = 1.59, 95% CI: 1.06-2.38, P = .026 for heavy drinking on colon cancer, RR = 1.38, 95% CI: 1.28-1.48, $P = 1.65 \times 10^{-16}$ for moderate drinking and RR = 1.92, 95% CI: 1.47-2.51, $P = 2.17 \times 10^{-6}$ for heavy drinking on rectum cancer). For different anatomical sites of colon, the risk of distal colon cancer increased both in moderate and heavy drinkers (RR = 1.24, 95% CI: 1.15-1.35, $P = 1.25 \times 10^{-7}$ and RR = 1.66, 95% CI: 1.17-2.36, P = .005, respectively) while no significant associations were observed in proximal colon cancer. Focusing on the study population, the risk of CRC increased among moderate and heavy drinkers in European populations as well as Asian populations, while the latter

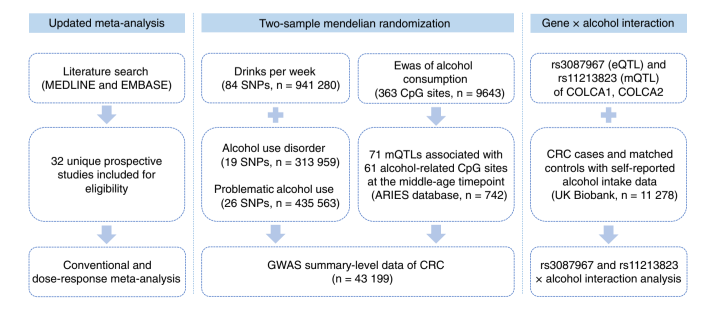


FIGURE 1 Study design

had larger magnitude and wider confidence intervals (RR = 1.20, 95% CI: 1.10-1.31, $P=4.39\times10^{-5}$ for moderate drinking and RR = 1.41, 95% CI: 1.16-1.71, $P=4.24\times10^{-4}$ for heavy drinking in European populations, RR = 1.29, 95% CI: 1.09-1.54, P=.003 for moderate drinking and RR = 1.53, 95% CI: 1.02-2.29, P=.038 for heavy drinking in Asian populations). In addition, a significant nonlinear doseresponse relationship between alcohol consumption and CRC risk was

identified, in which the dose of alcohol was treated as a continuous variable. Results showed that the risk of CRC incidence increased distinctly with increasing dose of alcohol intake between 0 and 50 g per day and increased modestly afterwards ($P=1.32\times10^{-7}$, adjusted R-square = .18). However, the risk of CRC mortality tended to reveal a J-shaped association with increasing dose of alcohol intake and the turning point was approximately 25 g per day (P=.014, adjusted

TABLE 1 Two-sample Mendelian randomization estimates for the relationship between alcohol consumption and colorectal cancer risk

Exposure	SNPs	MR method	OR (95% CI)	P _{Effect}	P _{Heterogeneity}	P _{Intercep}
Main analysis						
Drinks per week	84	IVW	1.79 (1.23, 2.61)	.003	1.39×10^{-4}	
		MR Egger	2.03 (0.93, 4.43)	.080.	$\textbf{1.10}\times\textbf{10}^{-4}$.717
		Weighted median	1.49 (0.91, 2.43)	.112		
		Simple mode	1.55 (0.44, 5.50)	.498		
		Weighted mode	1.19 (0.59, 2.41)	.635		
		MR-PRESSO	1.93 (1.36, 2.73)	3.90×10^{-4}		
Alcohol use disorder	18	IVW	1.33 (0.95, 1.85)	.093	.110	
		MR Egger	0.90 (0.49, 1.68)	.753	.155	.174
		Weighted median	1.13 (0.76, 1.69)	.541		
		Simple mode	1.62 (0.72, 3.65)	.257		
		Weighted mode	1.04 (0.65, 1.65)	.878		
		MR-PRESSO	1.33 (0.95, 1.85)	.111		
Problematic alcohol use	24	IVW	1.53 (1.02, 2.29)	.040	.002	
		MR Egger	0.91 (0.45, 1.83)	.787	.006	.093
		Weighted median	1.07 (0.69, 1.65)	.759		
		Simple mode	0.86 (0.35, 2.09)	.736		
		Weighted mode	0.95 (0.60, 1.50)	.819		
		MR-PRESSO	1.40 (0.99, 1.98)	.071		
Sensitivity analysis						
Drinks per week	63	IVW	1.77 (1.17, 2.69)	.007	.005	
		MR Egger	1.66 (0.75, 3.66)	.216	.004	.846
		Weighted median	1.53 (0.90, 2.60)	.114		
		Simple mode	2.25 (0.58, 8.69)	.243		
		Weighted mode	1.30 (0.63, 2.68)	.472		
		MR-PRESSO	1.96 (1.36, 2.83)	6.38×10^{-4}		
Alcohol use disorder	13	IVW	1.31 (0.86, 1.98)	.206	.063	
		MR Egger	0.95 (0.46, 1.98)	.901	.072	.325
		Weighted median	1.15 (0.74, 1.78)	.546		
		Simple mode	1.43 (0.57, 3.56)	.461		
		Weighted mode	1.06 (0.64, 1.76)	.827		
		MR-PRESSO	1.31 (0.86, 1.98)	.230		
Problematic alcohol use	17	IVW	1.27 (0.88, 1.81)	.199	.271	
		MR Egger	0.99 (0.55, 1.79)	.972	.279	.321
		Weighted median	1.08 (0.69, 1.69)	.745		
		Simple mode	0.87 (0.33, 2.30)	.776		
		Weighted mode	1.01 (0.63, 1.64)	.955		
		MR-PRESSO	1.27 (0.88, 1.81)	.218		



R-square = .14). Stratification by sex, cancer site and study population showed similar trends as incidence, except for colon cancer that showed an almost linear association ($P = 6.47 \times 10^{-11}$, adjusted *R*-square = .45) (Figure S3).

3.2 | Causal effect of genetic predisposition to alcohol drinking on CRC risk

We firstly calculated the F-statistic of each instrument and no weak instruments were identified (F-statistic > 10, Table S4). Among the instrumental variables for alcohol consumption (drinks per week, AUD and PAU), 84 (100%), 18 (94.7%) and 24 (92.3%) were matched with GWAS summary-level data of CRC, respectively. Using the IVW approach, we observed a significant causal association between genetically predicted drinks per week and increased risk of CRC: for per one SD increase of log-transformed alcoholic drinks per week, the risk of CRC would increase 79% (OR = 1.79, 95% CI: 1.23-2.61. P = .003). Genetic predisposition to PAU was nominally associated with increased risk of CRC (OR = 1.53, 95% CI: 1.02-2.29, P = .040), while no significant causal effect was observed between genetic predisposition to AUD and CRC risk (OR = 1.33, 95% CI: 0.95-1.85. P = .093) (Table 1), which might be due to the limited statistical power. There was substantial heterogeneity in all MR analyses except for that of AUD, but no apparent horizontal pleiotropy was observed in MR-Egger regression ($P_{\rm Intercept}=.717$ for drinks per week, $P_{\rm Intercept}=.174$ for AUD, $P_{\rm Intercept}=.093$ for PAU). Using MR-PRESSO approach, we discovered one outlier among the IVs for drinks per week, one outlier among the IVs for PAU but no outlier among the IVs for AUD (Table S5); the evidence of causal association remained for drinks per week (OR = 1.93, 95% CI: 1.36-2.73, $P=3.90\times10^{-4}$), but the evidence against the null hypothesis of no association was limited for PAU (OR = 1.40, 95% CI: 0.99-1.98, P=.071) after removing the outlier. As sensitivity analyses, we excluded IVs associated with potential confounders (Table S6) and observed evidence for the causal effect of drinks per week on CRC risk (OR = 1.77, 95% CI: 1.17-2.29, P=.007), but limited evidence for the effect of AUD and PAU on CRC risk due to the inadequate study power (OR = 1.31, 95% CI: 0.86-1.98, P=.206 for AUD and OR = 1.27, 95% CI: 0.88-1.81, P=.199 for PAU).

3.3 | Causal effect of alcohol-related DNA methylation on CRC risk

We identified 363 CpG sites correlated to alcohol consumption of European ancestry from the EWAS conducted by Liu et al.²¹ A total of 71 mQTLs were identified to be robustly associated with 61 CpG sites from ARIES database at the timepoint of middle age using GCTA results and included in the methylation MR analysis, among which

TABLE 2 Two-sample Mendelian randomization estimates for alcohol-related CpG sites and colorectal cancer risk

CpG sites	Chr	Position	Nearest gene (s)	Method	SNP	OR (95% CI)	P value
cg10045354	11	111 169 427	COLCA2, COLCA1	Wald ratio	1	1.67 (1.47, 1.89)	$\textbf{3.54}\times\textbf{10}^{-\textbf{16}}$
cg05593667	6	35 490 744	-	Wald ratio	1	0.89 (0.84, 0.95)	$\textbf{1.25}\times\textbf{10}^{-\textbf{4}}$
cg03575969	10	82 172 508	STARD10, ARAP1	Wald ratio	1	1.17 (1.05, 1.31)	5.91×10^{-3}
cg22871253	6	159 238 744	EZR	Wald ratio	1	1.12 (1.02, 1.24)	2.08×10^{-2}
cg12662084	6	17 809 126	KIF13A	Wald ratio	1	0.91 (0.84, 0.99)	2.31×10^{-2}
cg26312998	6	43 337 775	ZNF318	Wald ratio	1	1.05 (1.01, 1.10)	2.80×10^{-2}
cg17390562	6	159 238 463	EZR	Wald ratio	1	1.10 (1.01, 1.19)	3.48×10^{-2}
cg10456541	2	8 721 512	-	Wald ratio	1	1.14 (1.01, 1.30)	3.94×10^{-2}

Note: The bold ones were those that survived multiple-testing correction (Bonferroni P < .05). Abbreviation: Chr. chromosome.

TABLE 3 Gene-alcohol interaction estimates for the risk of CRC nested in the UK Biobank

	Basic model		Multivariable model		
Variables	OR (95% CI)	P value	OR (95% CI)	P value	
Alcohol (10 g/day)	1.08 (1.05, 1.10)	4.10×10^{-12}	1.06 (1.04, 1.09)	3.86×10^{-8}	
rs3087967	1.08 (1.01, 1.15)	.022	1.08 (1.01, 1.16)	.018	
rs11213823	1.07 (1.00, 1.14)	.043	1.07 (1.00, 1.15)	.035	
$rs3087967 \times Alcohol \text{(10 g/day)}$	1.04 (1.01, 1.06)	.001	1.03 (1.00, 1.05)	.027	
$rs11213823 \times Alcohol \text{(10 g/day)}$	1.04 (1.02, 1.06)	.001	1.03 (1.00, 1.05)	.016	

Note: Basic model: adjusted for age and sex; multivariable model: additionally adjusted for area deprivation index, red meat consumption, processed meat consumption, aspirin intake, BMI and smoking.

	Basic model		Multivariable mode	el
Alcohol (10 g/day)	OR (95% CI)	P value	OR (95% CI)	P value
rs3087967 CC	1.03 (0.97, 1.10)	.392	1.01 (0.95, 1.09)	.710
rs3087967 CT	1.07 (1.04, 1.11)	6.99×10^{-6}	1.07 (1.04, 1.11)	$\textbf{1.54}\times\textbf{10}^{-5}$
rs3087967 TT	1.09 (1.06, 1.13)	7.09×10^{-9}	1.08 (1.05, 1.12)	4.43×10^{-7}
rs11213823 CC	1.03 (0.96, 1.09)	.401	1.01 (0.94, 1.08)	.822
rs11213823 CT	1.07 (1.04, 1.11)	5.66×10^{-6}	1.06 (1.03, 1.10)	$\textbf{1.72}\times\textbf{10}^{-4}$
rs11213823 TT	1.09 (1.06, 1.12)	5.88×10^{-8}	1.07 (1.04, 1.11)	1.58×10^{-5}

TABLE 4 Dose-response effect of alcohol drinking on the risk of CRC stratified by the genotype of rs3087967 in the UK Biobank

Note: Basic model: adjusted for age and sex; multivariable model: additionally adjusted for area deprivation index, red meat consumption, processed meat consumption, aspirin intake, BMI and smoking.

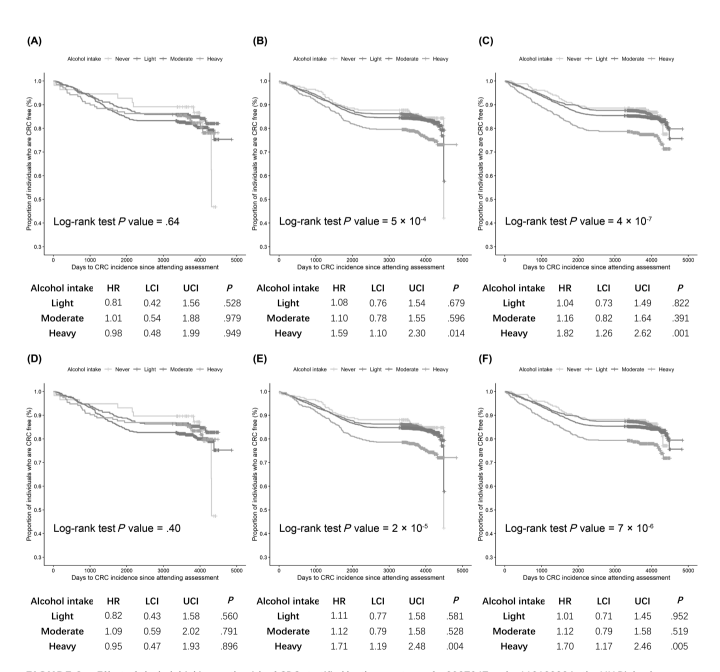


FIGURE 2 Effect of alcohol drinking on the risk of CRC stratified by the genotype of rs3087967 and rs11213823 in the UK Biobank: (A) rs3087967 CC genotype; (B) rs3087967 CT genotype; (C) rs3087967 TT genotype; (D) rs11213823 CC genotype; (E) rs11213823 CT genotype; (F) rs11213823 TT genotype

70 (98.6%) mQTLs were cis mQTLs (Table S7). To assess the causal effect of DNA methylation at alcohol consumption-related CpG sites on CRC risk, we conducted two-sample MR analyses using the identified mQTLs in the CRC GWAS summary data (Table S8). Eight CpG sites (cg03575969, cg05593667, cg10045354, cg10456541, cg12662084, cg17390562, cg22871253 and cg26312998) were identified to be causally associated with CRC risk at nominal significance level (P < .05) and two of them (cg05593667, cg10045354) survived Bonferroni correction (P < $.05/61 = 8.20 \times 10^{-4}$) (Table 2). CpG site cg10045354 was significantly associated with an increased risk of CRC (OR = 1.67, 95% CI: 1.47-1.89, $P = 3.54 \times 10^{-16}$), and located on the CpG island mapping to gene COLCA2 and COLCA1, which are key molecules involved in immunity and defense and have been identified as CRC susceptibility genes (Figure S4). To investigate whether the mQTL rs11213823 of cg10045354 would influence the expression of gene COLCA2 and COLCA1 in colon tissue, we queried on GTEx Portal and found it was a strong eQTL of gene COLCA2 (Figures S5 and S6). For the other identified CpG site cg05593667 which showed a casual association with CRC (OR = 0.89, 95% CI: 0.84-0.95, $P = 1.25 \times 10^{-4}$), we did not find any mapped gene(s) regulated by this CpG site and little is known about its nearest gene.

3.4 Gene-alcohol interaction effect on the risk of CRC

The baseline characteristics of the incident CRC cases and controls included for gene-alcohol interaction analysis were summarized in Table S9. Table 3 shows the effect estimates of alcohol drinking (per 10 g/day), rs3087967, rs11213823 and their interaction effects on the risk of CRC. When considering alcohol drinking and the two genetic variants independently, we observed that 10 g alcohol intake per day was significantly associated with 6% higher risk of CRC (OR = 1.06, 95% CI: 1.04-1.09, $P = 3.86 \times 10^{-8}$), rs3087967 was correlated with 8% higher risk of CRC (OR = 1.08, 95% CI: 1.01-1.16, P = .018) and rs11213823 was correlated with 7% higher risk of CRC (OR = 1.08, 95% CI: 1.00-1.15, P = .035) after adjustment for a wide range of covariates. When examining the gene-alcohol interaction effect, we found evidence for rs3087967-alcohol and rs11213823-alcohol interaction on CRC risk (P_{Interaction} = .027 and P_{Interaction} = .016, respectively). When assessing the dose-response effect of alcohol drinking (per 10 g/ d) stratified by the genotypes of rs3087967 and rs11213823, no significant association was observed among those who carried no risk allele of rs3087967 (OR = 1.01, 95% CI: 0.95-1.09, P = .710) and rs11213823 (OR = 1.01, 95% CI: 0.94-1.08, P = .822). However, alcohol intake was significantly linked with 7% and 8% higher risk of CRC among those with rs3087967 CT and TT genotypes (OR = 1.07, 95% CI: 1.04-1.11, $P = 1.54 \times 10^{-5}$ and OR = 1.08, 95% CI: 1.05-1.12, $P = 4.43 \times 10^{-7}$), and 6% and 7% higher risk of CRC among those with rs11213823 CT and TT genotypes (OR = 1.06, 95% CI: 1.03-1.10, $P = 1.72 \times 10^{-4}$ and OR = 1.07, 95% CI: 1.04-1.11, $P = 1.58 \times 10^{-5}$) for 10 g alcohol intake per day (Table 4). In the time-to-event multivariable Cox regression models, we observed significant higher risk of CRC development among

heavy drinkers, with the HR of 1.59 (95% CI: 1.10-2.30, P = .014) and 1.82 (95% CI: 1.26-2.62, P = .001) for carriers of rs3087967 CT and TT genotypes, and the HR of 1.71 (95% CI: 1.19-2.48, P = .004) and 1.70 (95% CI: 1.17-2.46, P = .005) for carriers of rs11213823 CT and TT genotypes comparing to never drinkers (Figure 2).

DISCUSSION

In our study, we conducted an updated meta-analysis of prospective cohort studies to explore the observational association between alcohol consumption and CRC risk, and then evaluated the causal impact of alcohol drinking on CRC risk using two-sample MR approaches. We additionally performed epigenetic MR analysis from the perspective of DNA methylation and gene-alcohol interaction analysis nested in the UK Biobank to understand how alcohol consumption modulates CRC risk.

The updated meta-analysis revealed that moderate and heavy drinking were observationally associated with an increased risk of CRC incidence but not for mortality. Dose-response analysis also illustrated distinct dose-risk manners: the risk of CRC incidence increased distinctly when alcohol consumption ranged from 0 to 50 g per day and modestly afterwards, nonetheless, the risk of CRC morality showed a J-shaped association, with the turning point at approximately 25 g per day. Taken these together, it is possible that the pathogenic effect of alcohol differs in terms of CRC incidence and mortality, but due to the limited adjusted R-square, further studies are warranted to proof our findings. Although we confined the metaanalysis to include only prospective cohort studies, these results should be interpreted with caution due to potential reverse causation. for that only eight studies set a lag time between the assessment of alcohol consumption and CRC occurrence. Using genetic instruments from two large-scale GWASs in European ancestry as proxies, 11,12 we appraised the causal relationship between genetic predisposition to alcohol consumption and the risk of CRC through two-sample MR analyses, in which the phenotype of "drinks per week" represents the general exposure of alcohol consumption, and other two phenotypes "AUD" and "PAU" reflects pathological drinking behaviors. Convincing evidence in support of a significant causal effect of drinks per week and suggestive evidence for a nominally significant causal effect of PAU with CRC were observed. These findings are similar to a previously published meta-analysis based on a MR approach by Wang et al who used the aldehyde dehydrogenase 2 (ALDH2) variant as a surrogate of alcohol exposure and reported an odds ratio of 1.31 (95% CI: 1.01-1.70) supporting the role of alcohol in colorectal carcinogenesis.³⁵

Along with observational association, the mechanisms by which alcohol consumption exerts its carcinogenic effect are not fully understood. One possible hypothesis is that DNA methylation may mediate the effects of alcohol on CRC risk. Alcohol may influence DNA methylation patterns by inhibiting the activity and expression of enzymes involved in DNA methylation, such as DNA methyltransferases.³⁶⁻³⁹ To appraise the causal effect of DNA methylation induced by alcohol consumption on CRC risk, we conducted a two-sample MR using

mQTLs of alcohol drinking related CpG sites as proxies. We found two CpG sites (cg05593667, cg10045354) significantly associated with CRC risk after multiple correction. The CpG site cg10045354 was annotated to genes *COLCA1* and *COLCA2*, which have been identified as genes whose expression levels were significantly associated with CRC susceptibility as reported by previous GWAS.⁴⁰

A number of studies have examined the interactions between alcohol intake and CRC susceptibility genes to investigate any effect modification. For instance, Gong et al identified interactions between alcohol consumption and variants in the 9q22.32/HIATL1 region, and found that light to moderate drinkers with rs9409565 CT and TT genotype were associated with a lower risk of CRC compared to nondrinkers or occasional drinkers. 16 Our previous study reported a weak evidence suggesting that heavy drinkers with G allele of MTR A2756G variant conferred higher risk of CRC.41 To validate our MR findings and test whether the effect of alcohol-related methylation on CRC was influenced by the expression level of COLCA1/COLCA2 gene, we performed alcohol-gene interaction analysis using the eQTL rs3087967 and mQTL rs11213823. Our study identified novel alcohol-gene interactions at both the eQTL and mQTL of COLCA1/ COLCA2 in terms of CRC risk. Dose-response analysis stratified by the genotypes provided supporting evidence for the dose dependent carcinogenesis effect of alcohol drinking on CRC risk among risk allele carriers of rs3087967 and rs11213823, and time-to-event analysis revealed that the effect was more prominent among heavy drinkers when comparing with light and moderate drinkers. Our findings vielded insight to understand how alcohol interacts with the methylation and expression level of COLCA1/COLCA2 gene to modulate the tumorigenesis of CRC.

COLCA1 is a protein-coding gene and expressed in many cells that are frequently involved in immunity and defense, such as eosinophils and mast cells.⁴² Eosinophils can induce apoptosis and kill tumor cells via releasing ECP, EDN, TNF and granzyme,⁴³ while mast cells may be damaging to tumor cells through cytokines and proteolytic enzyme secretion,⁴⁴ indicating that COLCA1 may exert its tumoricidal functions through immunity. COLCA2 is found to be coregulated with COLCA1 and is present in many epithelial cells. The expression of COLCA2 is increased in immune and other cells of the microenvironment and reduced in tumor cells,⁴⁵ which suggests a potential protective role of COLCA2 in antitumor immunity. Alcohol intake may exert the carcinogenic effect on CRC by downregulating the expression of COLCA1/COLCA2 gene through epigenetic modification.

Our study has several strengths, which include the comprehensive evaluations of the association between alcohol consumption with CRC outcomes through updated meta-analysis and two-sample MR analyses. Furthermore, by integrating several large-scale datasets, causal effects of alcohol related CpG sites on CRC risk and positive interaction effect between susceptibility gene and alcohol drinking were detected to possibly explain potential pathogenic mechanisms underlying significant observations. However, our study also has limitations. Our MR findings were concluded within the European descents, while a strong genetic determinant of alcohol consumption on aldehyde dehydrogenase 2 (ALDH2 rs671) which is East Asian-

specific genetic polymorphism⁴⁶ was not investigated in our study. The mQTL effects used in the methylation MR analysis were obtained from the ARIES middle-age time point, which only contained females at this time point. Without access to the full EWAS summary-level data, we were not able to perform mediation analysis to estimate the magnitude of carcinogenesis effect of alcohol drinking on CRC mediated by DNA methylation. To better control the potential covariates, we performed the gene-alcohol interaction under the framework of a nested case-control study in UK biobank, further study is needed to validate the findings in a perspective way.

In conclusion, our study found that genetic predisposition to alcohol drinking was causally associated with increased CRC risk and the pathogenic effect of alcohol could be partly attributed to DNA methylation via regulating the expression of *COLCA1/COLCA2* gene; the eQTL rs3087967 and mQTL rs11213823 polymorphism of *COLCA1/COLCA2* gene would interact with alcohol consumption to increase the risk of CRC. Further studies with larger-scale and tissue-specific dataset are warranted to confirm the findings by providing molecular evidence.

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CONFLICT OF INTEREST

The authors declare no potential conflict of interests.

DATA AVAILABILITY STATEMENT

The results of our study are included in this published article and its Supporting Information files. The UK Biobank is an open access resource and bona fide researchers can apply to use the UK Biobank dataset by registering and applying at http://ukbiobank.ac.uk/registerapply/. Further information is available from the corresponding author upon request.

ETHICS STATEMENT

All studies were approved by the respective institutional review boards and conducted with appropriate ethical criteria in each country and in accordance with the Declaration of Helsinki.

ORCID

Maria Timofeeva https://orcid.org/0000-0002-2503-4253 Philip J. Law https://orcid.org/0000-0001-9663-4611

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Kefeng Ding https://orcid.org/0000-0002-2380-3717 Xue Li https://orcid.org/0000-0001-6880-2577

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SUPPORTING INFORMATION

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