Research Article

Genome-Wide Association and Transcriptome-Wide Association Studies Identify Novel Susceptibility Genes Contributing to Colorectal Cancer

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Received 15 April 2022; Accepted 31 May 2022; Published 1 July 2022

Academic Editor: Fu Wang

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Background. Colorectal cancer (CRC) is among the most common cancers diagnosed worldwide. Although genome-wide association studies have effectively identified the genetic basis of CRC, there is still unexplained variability in genetic risk. Transcriptome-wide association studies (TWAS) integrate summary statistics from CRC genome-wide association studies (GWAS) with gene expression data to prioritize these GWAS findings and uncover additional gene-trait correlations.

Methods. First, we carried out a post-GWAS analysis using summary statistics from a large-scale GWAS of CRC (n = 4,562 cases, n = 382,756 controls). Second, combined with the expression weight sets from GTEx (v7), susceptibility genes were identified with the FUSION software. Colocalization, conditional and fine-mapping analyses, phenome-wide association study (pheWAS), and Mendelian randomization were employed to further characterize the observed correlations.

Results. In the post-GWAS analyses, we first identified new genome-wide significant associations: three genomic risk loci were identified at 8q24.21 (rs6983267, \( P = 6.98 \times 10^{-12} \)), 15q13.3 (rs58658771, \( P = 1.40 \times 10^{-10} \)), and 18q21.1 (rs6507874, \( P = 1.91 \times 10^{-14} \)). In addition, the TWAS also identified four loci statistically significantly associated with CRC risk, largely explained by expression regulation, including six candidate genes (DUSP10, POUSF1B, C11orf53, COLCA1, COLCA2, and GREM1-AS1). We further discovered evidence that low expression of COLCA2 is correlated with CRC risk with Mendelian randomization.

Conclusions. We discovered novel CRC risk loci and candidate functional genes by merging gene expression and GWAS summary data, offering new insight into the molecular processes underlying CRC development. This makes it easier to prioritize potential genes for follow-up functional research in CRC.

1. Introduction

Colorectal cancer (CRC) is one of the most common cancers identified globally [1], accounting for around 10% of all cancers and cancer-related deaths identified each year, with over 1.2 million people detected with CRC each year and millions of deaths from CRC each year [2]. Notably, in some high-income countries, such as Australia, Canada, the United States, and some European countries, 5-year survival rates have reached nearly 65%, but in low-income countries, 5-year relative survival rates are still less than 50%, and survival rates decline with age [3–5].

Unlike monogenic diseases, CRC is a polygenic disease produced by genetic inheritance and environmental factors (such as obesity, physical inactivity, poor diets, alcohol consumption, and smoking) which play a major part in the
etiology of both familial and sporadic CRC [6–8]. Twin family studies have shown heritability of about 33%–48% for CRC, implying that a major genetic component is generating the phenotypic variation [9]. Patients with CRC who have a positive family history represent approximately 10–20% of the total. The risk of disease varies with the quantity of relatives of patients with CRC, the severity of the disease, and the age at which CRC is diagnosed [10, 11].

Several GWAs on CRC have discovered around 60 correlation signals at more than 50 loci over the past decades, while an increasing amount of single nucleotide polymorphisms (SNPs) have shown statistical significance but often only a little fraction of the risk for CRC risk [12–14]. Even though GWAS has been quite successful in identifying elements that contribute to the genetic architecture of CRC, the loci detected are generally difficult to characterize biologically. In contrast, a transcriptome-wide association study (TWAS) provides more interpretable biologically relevant findings owing to the usage of disease-relevant cell types and tissues, and also databases detailing tissue-specific expression [15]. TWAS could detect genes whose gene-regulated expression may be related to the risk of diseases by merging expression quantitative loci (eQTL) results with GWAS summary data [15].

To identify genetically regulated risk loci associated with CRC, we performed gene-based and gene-set tests utilizing the CRC GWAS summary statistics from the publicly available UK Biobank with the FUMA online tool, an available online website at http://fuma.ctglab.nl [16, 17]. Then, we also leverage the currently available CRC GWAS summary statistical to conduct a TWAS; the cohort includes 4,562 CRC cases and 382,756 controls from Europe. Possibly relevant tissue-derived transcriptomic expression weights were employed, comprising the whole blood and 2 CRC-relevant tissue (colon transverse and colon sigmoid) panels from GTEx (v7). Subsequently, we conducted the conditional analysis of all significant TWAS correlations to identify the jointly significant TWAS genes (i.e., the driven genes at each risk locus). Follow-up analyses, including summary data-based Mendelian randomization (SMR), colocalization analysis, fine-mapping analysis, and phenome-wide association study (pheWAS), were performed to broadly characterize the associations identified. Altogether, our findings demonstrate TWAS’s ability to identify CRC risk genes with small effect sizes and present a testable target for future functional validation of CRC.

2. Method

2.1. Study Cohort. Included in this analysis were the following data: (i) genome-wide summary data from the GWAS of CRC by Zhou et al. [17], (ii) 3 SNP weight sets from GTEx (v7) transcriptomic reference samples, and (iii) the 1000 Genomes Project reference for linkage disequilibrium (LD) estimation.

First, we utilized the CRC GWAS summary data from UK Biobank analysis results (https://www.leelabsg.org/resources), in which individuals (n = 4,562 cases, n = 382,756 controls) had European genetic heritage [17]; second, SNP weight sets from the relevant tissues were used. SNP weight sets indicate the correlation of the SNP with its annotated gene expression [15]; SNP weight sets from colon transverse, colon sigmoid, and whole blood were obtained from the FUSION website (http://gusevlab.org/projects/fusion/); third, the LD reference of the 1000 Genomes Phase 3 European (N = 489) was also obtained from the FUSION website (http://gusevlab.org/projects/fusion/).

2.2. Identification of Independent SNPs and Lead SNPs with FUMA. Utilizing information from multiple biological resources, FUMA has established a range of SNP functional annotation methods on the website platform, providing functional annotations of GWAS association signals, gene prioritization [16]. The identification of lead SNPs and candidate SNPs is based on the following criteria: (i) independently significant SNPs were identified by $P < 5 \times 10^{-8}$ and independent from each other at $r^2 < 0.6$; (ii) independent lead SNPs were identified as independent significant SNPs and independent from each other at $r^2 < 0.1$; (iii) genomic risk loci was identified by merging lead SNPs within a 250 kb window and all known SNPs in LD of $r^2 \geq 0.6$ with one of the independent significant SNPs; and (iv) the 1000 Genomes Phase 3 European was defined as the reference panel population [16].

2.3. MAGMA for Gene-Based and Gene-Set Tests. Gene-based tests/gene-set analyses are methods capable of summarizing SNP associations at the gene level and associating gene sets with biological pathways. To determine prospective candidate genes and biological processes enriched for markers with low but not necessarily genome-wide significant $P$ values in GWAS, gene-based tests/gene-set analyses were conducted by MAGMA, which was implemented in FUMA [16].

The CRC GWAS summary statistics were utilized in MAGMA’s gene-based analysis to aggregate the association of SNPs within gene regions without accounting for SNP effects on gene expression [18]. The SNP-based $P$ values from the CRC GWAS summary statistics were utilized as the input file for the gene-based analysis. For gene-based studies in MAGMA, we employed all 19,427 protein-coding genes from the NCBI 37.3 gene definition as the foundation (https://ctg.cnclr.nl/software/magma). After SNP annotation, 19,252 genes had at least one SNP covering. Note that the LD relationship between SNPs was taken into account when performing the gene association test. We employed a strict Bonferroni correction to account for numerous testing, and the genome-wide threshold for significance was set at $P = 2.60 \times 10^{-6}$ (0.05/19252). Gene-set tests were carried out in MAGMA utilizing competitive genomic analysis, which integrated the test statistics for all genes in a given genome to provide a joint association statistic, which was then implemented in FUMA [16]. This statistic was compared to that of all other genes not included in that set, while taking into consideration the quantity of SNPs within each gene, gene density, and differential sample size (unequal sample size contributing to each
gene). 10,678 gene sets (curated gene sets: 4761, GO terms: 5917) from MsigDB v6.2, as well as a default competitive test model, were employed in the MAGMA gene-set analysis. Bonferroni correction was performed by employing a false discovery rate threshold of 5%.

2.4. Transcriptome-Wide Association Study. We used TWAS analyses with FUSION (http://gusevlab.org/projects/fusion) with default settings to identify genes whose gene-regulated expression may be related to the risk of CRC [15]. TWAS was carried out using reference panels obtained from tissue-specific gene expression and the CRC GWAS summary data [15], taking into account the LD structure between SNPs. To interpret the LD structure, we used the 1000 Genomes Phase 3 (European, N = 489) data as the LD reference panel. FUSION software was used to generate SNP weight sets from BLUP, BSLMM, LASSO, Elastic Net, and top SNPs utilizing genotype and expression data unless BLUP/BSLMM was eliminated owing to sample size or convergence issues [15]. A strict Bonferroni-corrected threshold was utilized: $P < 1.11 \times 10^{-6}$ (0.05/44,953) (considering the relations between features within and across SNP weight sets).

2.5. Bayesian Colocalization. To determine whether GWAS SNPs colocalized with eQTLs, Bayesian colocalization was examined using the COLOC package (https://cran.r-project.org/web/packages/coloc/, version 5.1.0) in R implemented by FUSION for all associations with $P_{\text{TWAS}} < 0.05$ within a 1 megabase (Mb) window [19]. This Bayesian colocalization technique revealed the posterior probability (PP) that relationships within a locus for two outcomes (GWAS and eQTL signals) were caused by a common causal variable or variants in strong LD. COLOC evaluated five hypotheses: PP0, no eQTL and GWAS association; PP1, association with eQTL, but no GWAS; PP2, association with GWAS, but no eQTL; PP3, eQTL and GWAS association, but independent signals; and PP4, shared eQTL and GWAS associations. The main objective is to determine whether the GWAS and eQTL signals are consistent with common causal variants (i.e., PP4). In reality, a high PP (PP4 > 80%) implies that the GWAS and eQTL signals colocalize [19].

2.6. Joint/Conditional Analysis and Permutation. To determine multiple correlated features within a locus (or the same feature from multiple tissues), we performed a conditional analysis and hopefully identified which were conditionally independent. Moreover, we also wondered how much the GWAS signal remained after the correlation of the function was excluded [20, 21]. This process identified which features indicate independent associations (called jointly significant) and which features were not significant when the predicted expression of the jointly significant genes in the region was ignored (called marginally significant) [20]. We also computed the extent to which GWAS correlations within each locus can be described by the functional connections detected in this TWAS. Additionally, the conditional analysis enables us to evaluate the extent to which the GWAS association signal within each locus may be described by the functional connections identified in this TWAS. To account for correlations between features within and across SNP weight sets, we randomized the eQTL weights and recalculated empirical association statistics conditional on GWAS effects by FUSION. In this study, 1,000 permutation tests were performed for each TWAS gene, setting the significance of the permutation test at $P < 0.05$ [15, 20].

2.7. TWAS Fine Mapping. Fine-mapping of CaUsal gene Sets (FOCUS) is software for fine-mapping transcriptome-wide correlation study statistics to genomic risk areas discovered by FUSION and producing a set of reasonable features interpreting the observed genomic risk. The software accepts GWAS summary data and eQTL weights as input, as well as FUSION findings and weights. FOCUS calculates the posterior inclusion probability (PIP) for each feature in the region of interest and decides whether TWAS-significant genes are included in the default 90% credible set, which is the set of features most likely to include causal features [22]. PIP values greater than 0.05 for each feature in the related region suggest that the feature is more likely to be causal than any other feature in the associated region [21]. Consistent with the TWAS analysis performed by the FUSION software, the FOCUS software used the same TWAS reference panel from FUSION.

2.8. Summary Data-Based Mendelian Randomization of CRC. We applied summary data-based Mendelian randomization (SMR) based on pooled data (https://cnsgenomics.com/software/smr/) to determine genetic signals correlated with phenotypic and gene expression variation, utilizing pooling from independent GWASs and eQTL weight data (colon sigmoid, colon transverse, whole blood from GTEx (v7), and Westra, CAGE eQTL summary data) that can be employed to evaluate whether the effects of genetic variation are mediated by gene expression levels [23]. This method employs the concept of Mendelian randomization (MR), a methodology for detecting causal effects [24]. The first step in the SMR approach is MR analysis, in which genetic variants (such as SNPs) were defined as instrumental variables, gene expression levels were defined as exposures, and traits were defined as outcomes [25]. To control the genome-wide type I error rate, $P_{\text{SMR}}$ values were Bonferroni corrected for the number of genes tested, meaning that genes less than 1 Mb away from the GWAS lead SNPs were considered significant using the Bonferroni-corrected SMR significance threshold [26]. Following the SMR test, the heterogeneity independent instrument (HEIDI) test was used to determine whether the connection was attributed to a common causal variant rather than widespread LD across the genome [25]. Considering that this analysis is conservative for gene analysis and maintains fewer genes than when correcting for multiple testing, we did not correct for multiple testing and a $P_{\text{HEIDI}}$ threshold of 0.05 for the HEIDI test was identified [23].

2.9. Phenome-Wide Association Studies. To determine phenotypes associated with the best eQTL in the locus identified via TWAS, a pheWAS was performed using publicly
available data from the GWAS Atlas (https://atlas.ctglab.nl). Only the top phenotypes were recorded (excluding CRC). Accounting for the current GWAS Atlas website which contains a total of 3,302 unique traits, a Bonferroni-corrected cut-off of $1.68 \times 10^{-5}$ (0.05/the number of unique traits) was used.

3. Result

3.1. Study Overview. First, we used the FUMA online website and performed the post-GWAS analysis with CRC GWAS summary statistics as input files to identify independent significant SNPs and lead SNPs. Second, gene-based and gene-set tests were conducted using MAGMA and implemented in FUMA, combining the test statistics for all genes in a particular genome to obtain a joint association statistic [16]. After that, we used FUSION software to perform TWAS analysis to determine genes whose gene regulatory expression may be related to the risk of CRC with default settings [15]. Afterward, we conducted a Bayesian colocalization analysis for all associations with $P_{TWAS} < 0.05$ utilizing the COLOC package in R (https://cran.r-project.org/web/packages/cocoloc) implemented in FUSION to predict the PP association that an SNP contributed to the association signal in the GWAS and the eQTL [19]. Joint analysis was employed in regions with multiple significant associations to determine conditionally independent associations [22]. Beyond that, most genes remained significant following the permutation, indicating that their signal was genuine, not accidental. Finally, we used SMR, pheWAS, and FOCUS software to analyze and verify the above results [23] (Figure 1).

3.2. FUMA’s Functional Annotation Analysis Results. To determine genetic loci that contribute to CRC and outcome, SAIGE, scalable and accurate software for generalized mixed-model association testing, was utilized to efficiently analyze a CRC case-control cohort, including a sample of 387,318 Caucasians of British European ancestry, with an imbalanced control case-control ratios and sample correlations [17]. A Manhattan plot of the GWAS results was drawn with the 28,146,008 SNPs that satisfied quality controls (Figure S1a). The quantile-quantile plot revealed an excellent match between the observed $P$ value distributions and the predicted $P$ value distribution by chance ($\lambda = 1.035$ ; Figure S1b), indicating that the genome-wide statistical results were not overinflated.

In the post-GWAS process, we first utilized the FUMA online website to conduct the functional annotation with the CRC GWAS summary statistics including 4,562 cases and 382,756 controls. After functional annotation analyses, we annotated 131 candidate SNPs that passed the genome-wide significance threshold ($P < 5.00 \times 10^{-8}$), and three independent lead SNPs were identified located at three genomic risk loci. These 131 independently significant SNPs were found in introns (32.3%, $P_{enrichment} = 0.36$) and noncoding RNA intronic (31.5%, $P_{enrichment} = 9.39 \times 10^{-10}$), intergenic (15.4%, $P_{enrichment} = 7.68 \times 10^{-14}$), and UTR3 (15.4%, $P_{enrichment} = 1.54 \times 10^{-18}$) regions that showed enrichment, while only 2.31% of the 131 independently significant SNPs were located in exonic regions and noncoding RNA exonic regions (Table S1, Figure S2).

One of the interesting genomic risk loci is 8q24.21, which contained 42 GWAS SNPs, representing 6 unique genes (LGOLGA8N, ARHGAP11A, SCG5, GREM1, and FMN1) and 52 GWAS SNPs (Figure S3a). Consistent with the previous study, SNPs near GREM1 and FMN1 were highly correlated with elevated CRC risk [30], while the difference is that the rs4779584 reported is strongly related to an elevated risk of CRC, but the opposite in the present study ($P = 1.73 \times 10^{-9}$, OR = 0.85) [30]. In addition to the above two risk loci, there is also another genomic risk locus at 18q21.1. CTIF, SMAD7, and DYM, 3 protein-coding genes, were included in this locus. rs6507874 is the lead SNP of the genomic risk locus and is found in the intron of the SMAD7 gene (Figure S3c), while the previous research that studied the relationship between rs6507874 and CRC risk showed that rs6507874 did not show a statistically significant connection ($P = 0.075$) with increased SMAD7 expression [31].

3.3. Gene-Based and Gene-Set Tests Implemented in MAGMA. All SNPs found inside genes were assigned to 19,252 protein-coding genes in the gene-based analysis conducted with MAGMA. After gene-based analysis, 19,252 genes had at least one SNP covering. The gene-based analysis for CRC summary statistics identified 5 genes (SMAD7, COLCA1, COLCA2, POUS5F1B, and LAMAS5) at a stringent Bonferroni correction for significance at $P < 2.60 \times 10^{-6}$ (Figure S4, Table S2). Immediately after, the results of the gene-set analysis conducted by MAGMA showed nonsignificant results after Bonferroni correction for numerous testing (Table S3). But it is worth noting that several suggested gene sets, such as neuroendocrine cell differentiation, linoleic acid (LA) metabolism, are known CRC-related pathways [32, 33].

3.4. Transcriptome-Wide Association Study. To identify potential genes associated with CRC risk, we collected a publicly available GWAS dataset from a European case-control cohort. Of the 3 SNP weight sets (colon sigmoid, colon transverse, and whole blood), we identified 6 transcriptome-wide significant features (6 unique genes), with the colon sigmoid transcript-level weights yielding the most significant relationships (Table 1). Four transcriptome-wide significant loci were detected for 6 distinct genes (Figure 2).

3.5. Chromosome 1q41. Only one relevant feature in the locus, correlating to the DUSP10 gene, was shown to be transcriptome-wide significant ($P_{TWAS} = 8.72 \times 10^{-7}$) (Table 1): rs12125368, an intergenic variant, was the SNP in the locus most highly correlated with CRC (odds ratio (OR) = 1.34, $P_{GWAS} = 2.65 \times 10^{-6})$. And the best
Figure 1: Schematic workflow of this study. We performed a TWAS for the CRC based on the publicly available GWAS datasets and the eQTL datasets. The GWAS datasets were derived from CRC GWAS summary statistics from UK Biobank analysis results; participants (n = 4,562 cases, n = 382,756 controls) were of European genetic ancestry. The eQTL dataset was from GTEx (v7). Follow-up analyses, including the SMR, colocalization analyses, conditional analysis, fine-mapping analysis, and pheWAS, were performed to extensively characterize the identified associations. CRC: colorectal cancer; eQTL: expression quantitative trait loci; GTEx: Genotype-Tissue Expression Project; GWAS: genome-wide association study; pheWAS: phenome-wide association study; SMR: summary data-based Mendelian randomization; SNP: single nucleotide polymorphism; UKBB: UK Biobank; WGS: whole-genome sequencing.
eQTL in the locus correlated with the expression level of the DUSP10 gene \( (P_{eQTL} = 3.86 \times 10^{-4}) \) was rs6695584, which was in moderate LD with rs12125368 \( (0.6 \geq r^2 \geq 0.4) \) (Figure 3(a)). Then, formal Bayesian colocalization indicated a moderate shared signal with a PP4 of 0.52 (Table 1), confirming the general likelihood that the GWAS and colon sigmoid eQTL signals may share the same variants at this locus.

To determine whether the signals of this locus were independent, we conducted conditional and joint analyses.
Figure 3: Continued.
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Figure 3: The locus-compare scatter plot for the association signals at DUSP10, POU5F1B, C11orf53, COLCA1, COLCA2, and GREM1-AS1 in the European cohort. Colocalization analyses results are shown for (a) DUSP10, (b) POU5F1B, (c) C11orf53, (d) COLCA1, (e) COLCA2, and (f) GREM1-AS1 genes. The locus-compare scatter plot compares the expression quantitative trait loci (eQTL) results and the genome-wide association study (GWAS) results, which indicates whether the GWAS top locus is also the leading SNP in the eQTL result. The eQTL results were from Genotype-Tissue Expression (GTEx) (v7). The GWAS results were from the European cohort (n = 4,562 cases, n = 382,756 controls). The gene prioritized in each locus is shown on the y-axis of the corresponding figure label. Chr.: chromosome.
The result showed that the DUSP10 gene describes all of the signals at its loci (best SNP: rs12125368, $P_{\text{GWAS}} = 2.65 \times 10^{-6}$; conditioned on DUSP10, $P_{\text{GWAS}} = 0.89$) (Figure 3(a) and Figure S5a). FOCUS was utilized to assign a PIP to genes at each transcriptome-wide significant loci with relevant tissue in order to select putatively causal genes. Unfortunately, the PIP of DUSP10 was not available in the three eQTL tissues from GTEx (v7) for this genomic locus 1q41 (Table 1).

### 3.6. Chromosome 8q24.21. POU5F1B (colon transverse) achieved transcriptome-wide significance, located at 8q24.21 locus ($P_{\text{TWAS}} = 4.32 \times 10^{-7}$). rs6983267, a noncoding transcript variant in the CCA2 gene or intron variant in the CASC8 gene, was the most significant SNP associated with CRC at this locus (OR = 0.71, $P_{\text{GWAS}} = 6.98 \times 10^{-12}$). At this locus, rs7014346 was the best eQTL correlated with the expression level of the POU5F1B gene in the colon transverse tissue from the GTEx database (v7) ($P_{\text{eQTL}} = 6.65 \times 10^{-7}$), which was in moderate LD with rs6983267 (0.6 $\geq r^2 \geq 0.4$) (Figure 3(b)). Colocalization analysis supported the fifth hypothesis with a PP4 of 0.73, offering suggestive evidence that the significant CRC GWAS signal and the expression level of the POU5F1B gene were driven by the same causal variant.

For the genomic locus 8q24.21, POU5F1B was added in the 90%-credible gene set with a PIP of 1 in the colon transverse (Table 1). At this locus, conditional and joint analyses found that POU5F1B explains 0.53 of the variance at its loci (best SNP: rs6983267, $P_{\text{GWAS}} = 6.98 \times 10^{-12}$; conditioned on POU5F1B, $P_{\text{GWAS}} = 2.30 \times 10^{-6}$) (Figure 3(b) and Figure S5b).

### 3.7. Chromosome 11q23.1. Three transcriptome-wide significant genes (C11orf53, COLCA1, and COLCA2) were observed within the q23.1 region of chromosome 11 ($P_{\text{TWAS}} = 2.80 \times 10^{-7}$; 1.08 $\times 10^{-6}$; and 1.11 $\times 10^{-6}$, respectively). rs7130173 is an intron variant, most significantly associated with CRC at its loci (OR = 0.76, $P_{\text{GWAS}} = 2.52 \times 10^{-7}$). Meanwhile, our study, respectively, identified rs6589218, rs6589220, and rs3087967 as the best eQTL in the locus correlated with the expression level of C11orf53, COLCA1, and COLCA2 genes in the colon transverse tissue ($P_{\text{eQTL}} = 3.46 \times 10^{-16}$, 2.71 $\times 10^{-20}$, and 3.98 $\times 10^{-27}$, respectively) (Figures 3(c)-(e)). In addition, colocalization analysis identified a strong LD between the best eQTL (rs7130173) and the best eQTL (rs6589218, rs6589220, and rs3087967) (1.0 $\geq r^2 \geq 0.8$) (Figures 3(c)-(e)), and with PP4s for causality (C11orf53, PP4 = 0.99; COLCA1, PP4 = 0.99; and COLCA2, PP4 = 0.99, respectively), confirming that the significant CRC GWAS signal and colon transverse eQTL signals share the same variant at its locus.

Similarly, conditional and joint analyses showed that conditioning on C11orf53 completely described the variance of the loci on chromosome 11 (best SNP: rs7130173, $P_{\text{GWAS}} = 2.52 \times 10^{-7}$; conditioned on C11orf53, $P_{\text{GWAS}} = 1$). The fine-mapping findings revealed that the PIP values for C11orf53, COLCA1, and COLCA2 were 0.36, 4.77 $\times 10^{-5}$, and 0.48, respectively. However, only C11orf53 and COLCA2 were included in the 90%-credible gene set.

### 3.8. Chromosome 15q13.3. The GREM1-AS1 gene reached transcriptome-wide significance ($P_{\text{TWAS}} = 1.54 \times 10^{-11}$) and was the only significant association signal at this locus. Among this locus, rs1919360 was the most significant SNP associated with CRC, and the closest to the GREM1 gene (OR = 1.51, $P_{\text{GWAS}} = 1.91 \times 10^{-10}$). The best eQTL in the locus associated with the expression level of GREM1-AS1 was rs2611583 ($P_{\text{eQTL}} = 4.42 \times 10^{-4}$), which is shown to be weak LD with rs1919360 (0.2 $\geq r^2 \geq 0$) (Figure 3(f)), while the fifth hypothesis was supported by formal Bayesian colocalization with a PP4 of 0.70 (Table 1), providing modest evidence that significant CRC GWAS signaling and colon sigmoid eQTL signaling share the same variants at their loci.

For the genomic locus 15q13.3, GREM1-AS1 was included in the 90%-credible gene set with a PIP of 1 in the colon sigmoid (Table 1). At this locus, conditional and joint analyses found that GREM1-AS1 explains 0.83 of the variance at its loci (best SNP: rs1919360, $P_{\text{GWAS}} = 1.90 \times 10^{-10}$; conditioned on GREM1-AS1, $P_{\text{GWAS}} = 8.10 \times 10^{-7}$) (Figure 3(f) and Figure S5d).

### 3.9. Summary Data-Based Mendelian Randomization Results. We applied SMR and HEIDI to investigate whether gene expression levels are mediated by genetic variation, testing causal associations between CRC susceptibility gene expression levels and CRC (Figure S6). We identified significantly associated gene expression levels in the colon transverse data from GTEx (v7) that passed the HEIDI test at chromosome 11q23.1 (Figure S7a) ($P_{\text{SMR}} = 3.59 \times 10^{-6}$), and there has been no substantial heterogeneity underlying the eQTL signals ($P_{\text{HEIDI}} > 0.05$). Also, this Mendelian randomization study showed a causal relationship between low COLCA2 expression and CRC risk (Figure S7b).

We continued with SMR analyses of the eQTL summary data conducted by Westra et al. and Lloyd-Jones et al., respectively [34, 35]. However, after correcting for multiple testing, we were unable to detect any significant pleiotropic connection (Table S4).

### 3.10. Comparison with Previous Literature. Our TWAS study identified 6 genes significantly associated with CRC (GREM1-AS1, C11orf53, POU5F1B, DUSP10, COLCA1, and COLCA2) with a Bonferroni-corrected threshold of $P < 1.11 \times 10^{-6}$ (0.05/44,953). Compared with the previous largest TWAS study for CRC, which detected 25 associated genes with CRC risk at a Bonferroni-corrected threshold of $P < 9.10 \times 10^{-4}$ by MetaXcan software, we found that three of the significant TWAS genes (C11orf53, COLCA1, and COLCA2) were overlapped with the TWAS result performed by Guo et al. [36]. In addition, SFMBT1 was almost close to the Bonferroni-corrected threshold in our research ($P = 5.19 \times 10^{-4}$), overlapping with the largest TWAS results for CRC. Obviously, in addition to the three significant genes mentioned above, we also identified three novel associations (GREM1-AS1, POU5F1B, and DUSP10). The above differences may be due to different GWAS summary
statistics data or the type and quantity of SNP weight sets used, algorithm of the TWAS software and statistical thresholds.

### 3.11. Phenome-Wide Association Study

A PheWAS was performed for each best eQTL in 6 transcriptome-wide important features to further identify phenotypes that may be related or comorbid with CRC. In the process, we found that most of the eQTL were significantly associated with CRC, so we excluded the associated CRC traits to be able to effectively identify other phenotypes associated with them. Several best eQTL-related phenotypes were discovered to be strongly linked or comorbid with the risk of CRC, including bowel movement, alcohol consumption, C-C motif chemokine 22, family history of primary malignant neoplasm, cholelithiasis, and helicobacter pylori infection (Table S5) [14, 37–40].

### 4. Discussion

CRC is one of the most often diagnosed cancers, and it has a significant impact on cancer morbidity and mortality globally [41]. It is generally known that genetic factors play a significant role in the etiology of both familial and sporadic CRC [8, 42–44].

In this investigation, we first performed post-GWAS analyses of CRC GWAS summary data, including SNP annotation and gene-based and gene-set tests analysis using MAGMA, which was implemented in FUMA. In the follow-up gene-based analysis of GWAS, we identified 5 genes with significant association with CRC (SMAD7, COLCA1, COLCA2, POU5F1B, and LAMA5). In addition, gene-set analyses also identified neuroendocrine cell differentiation, LA metabolism, and other pathways associated with CRC, but they were not significant after multiple corrections. Second, we conducted a new TWAS on CRC of the European populations, which combined the CRC GWAS summary statistics and SNP weight sets to map four associated genes, including COLCA1, COLCA2, POU5F1B, and LAMA5. In addition, gene-set analyses also identified neuroendocrine cell differentiation, LA metabolism, and other pathways associated with CRC, but they were not significant after multiple corrections.

Confusingly, the PIP value for the GWAS and eQTL signals were identical. Specifically, Bayesian colocalization analyses were performed by COLOC package at the transcriptome-wide significant loci identified in this TWAS, and the PP4 of each gene sharing signal was calculated, providing evidence for whether the GWAS and eQTL signals share the same associations. We observed that no transcription-CRC signal was obtained from the same causal polymorphisms associated with SNP-CRC correlations, suggesting that most of the observed genes constituted linkage effects rather than pleiotropy. While these data show that transcription mediates the link between genetic vulnerability and CRC, neither TWAS nor colocalization can determine the causal. Therefore, we used SMR software to investigate the causal relationship between gene expression and CRC and only found that the COLCA2 gene identified in colon transverse map to 11q23.1 with COLCA1 and C11orf53 closing the threshold of pSMR value, suggesting multiple causal signals at this locus.

We also utilized a TWAS fine-mapping approach called FOCUS to obtain additional insight into which genes are likely causative for CRC, fine-mapping causal genes from several TWAS correlations at a locus and highlighting a single feature as the causal relation. Fine-mapping of the corresponding genomic loci prioritized GREM1-AS1 and POU5F1B in the 90%-credible gene set with a PIP of 1 in the colon sigmoid and colon transverse. For the genomic locus 11q23.1, COLCA2 was included in the 90%-credible gene set with the highest PIP in the colon transverse tissue. Confusingly, the PIP value for the DUSP10 gene was not available. Accordingly, we speculate that it may be due to the GWAS signal around the DUSP10 gene not reaching the threshold of significance and not meeting the computational inclusion requirements of the FOCUS software [22]. The above findings largely reflected local patterns of LD and indicated the requirement for further functional identification at several of these complicated loci.

Based on the above-mentioned various analyses, we found that the COLCA2 gene has an outstanding performance. COLCA2 was recognized as a colorectal cancer-associated gene, like COLCA1, and they were coregulated genes transcribed from opposite strands of a region of chromosome 11q23 associated with colon cancer [45]. COLCA2 is predominantly expressed in cells of epithelial, mesenchymal, and hematopoietic origin and has orthologs in a variety of mammals. Since COLCA2 expression is reduced in tumor cells from subjects with higher risk alleles, COLCA2 may play an important role in suppressing tumor formation in epithelial cells [45].

In compiling the results of this study’s analysis, we also raise some limitations that were worth discussing. First, the limited sample size of the GTEx (v7) gene expression reference sample may have hampered the identification of subtle transcriptome effects on CRC heredity, emphasizing the need for bigger samples [21, 23]. Second, our TWAS methodology only examined cis-eQTL of gene expression and did not account for trans-eQTL effects [15, 34]. Future studies should devote resources to building larger gene expression reference plates to be able to investigate trans-QTL effects. Third, the samples in this study were exclusively
European; therefore, the generalization of the findings to other ethnic groups was limited because of ethnic specificity.

5. Conclusion

In conclusion, we present the evidence for broad genomic and transcriptome alterations in colorectal cancer. Our study allows for the discovery of new connections as well as the elucidation of the genomic and transcriptome alterations that previously identified risk genes go through. We highlight genes that may be important for SMAD7, LAMAS5, GREM1-AS1, C11orf53, POU5F1B, DUSP10, COLCA1, and COLCA2. These results suggest that GWAS and TWAS are effective statistical methods to observe small- and large-effect genes correlated with CRC, providing a testable target for further functional validation of CRC, assisting in the knowledge of the molecular basis of the disease.

Abbreviations

CRC: Colorectal cancer  
eQTL: Expression quantitative loci  
FOCUS: Fine-mapping Of CaUsal gene Sets  
GWAS: Genome-wide association studies  
GTex: Genotype-Tissue Expression  
HEIDI: Heterogeneity independent instrument  
LD: Linkage disequilibrium  
Mb: Megabase  
MR: Mendelian randomization  
OR: Odds ratio  
pheWAS: Phenome-wide association study  
PPIP: Posterior inclusion probability  
PP: Posterior probability  
SNPs: Single nucleotide polymorphisms  
SMR: Summary data-based Mendelian randomization  
TWAS: Transcriptome-wide association studies.

Data Availability

The original contributions presented in the study are included in the article/supplementary material; further inquiries can be directed to the corresponding author.

Conflicts of Interest

The authors declared no competing interests.

Authors’ Contributions

Ruimin Yin and Binbin Song contributed equally to this work and share first authorship.

Acknowledgments

This study is funded by (1) the Key Discipline of Jiaxing Medicine Construction Project (No. 2019-zc-11) and (2) the Jiaxing Key Laboratory of Precision Treatment for Lung Cancer. We would like to thank the researchers and study participants for their contributions.

Supplementary Materials

Table S1: functional consequences of SNPs on genes. Table S2: summary statistics for significant genes ($P < 2.60E - 6$) identified in the gene-based. Table S3: results of the gene-set analyses. Table S4: the top ten probes were identified in the SMR analysis for 5 eQTL summary data. Table S5: phenotypes associated with top eQTLs derived from TWAS. Figure S1: Manhattan plot and quantile-quantile plot of the genome-wide $P$ values in the GWAS analysis. Figure S2: functional consequences of SNPs on genes. Figure S3: regional plots for associations in the regions surrounding the rs6983267, rs58658771, and rs6507874 in the GWAS analysis. Figure S4: gene-based genome-wide analysis for CRC by MAGMA to each locus in the GWAS analysis. Figure S5: conditional transcriptome-wide association study analysis for loci showing statistically significant associations with multiple genes. Figure S6: schematic diagram of the summary data-based Mendelian randomization model. Figure S7: prioritizing genes at the COLCA2 locus for CRC. (Supplementary Materials)

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