Abstract. In the current era of precision medicine, there is a general consensus that the anatomical site is an important factor in the management of colorectal cancer (CRC). To investigate the underlying molecular mechanisms between proximal and distal CRC and to identify the responsible genes, we analyzed the gene expression patterns of colorectal tumors from two microarray datasets, GSE39582 and GSE14333, on the NCBI Gene Expression Omnibus and the RNA-seq data from TCGA. Weighted coexpression network analysis (WGCNA) was applied to construct a gene coexpression network. The red module in GSE39582 and the dark-gray module from the TCGA dataset were found to be highly correlated with the anatomical site of CRC. A total of 12 hub genes were found in two datasets, 2 of which PLAG1 like zinc finger 2 (PLAGL2) and protein O-fucosyltransferase 1 (POFUT1) were common and upregulated in tumor samples in CRC. The module with the highest correlation provided references that will help to characterize the difference between left-sided and right-sided CRC. The survival analysis of PLAGL2 and POFUT1 expression revealed differences between proximal and distal CRC. Gene set enrichment analysis based on those two genes provided similar results: GPI anchor biosynthesis and peroxisome and selenoamino acid metabolism. PLAGL2 and POFUT1, which have the highest correlation with tumor location, may serve as biomarkers and therapeutic targets for the precise diagnosis and treatment of CRC in the future.

Introduction

Colorectal cancer (CRC), which accounted for approximately 1.8 million new cases and more than 860,000 deaths in 2018, ranks as the fourth most commonly diagnosed malignancy and the second leading cause of cancer-related deaths worldwide (1). The incidence and mortality rates of CRC are still increasing rapidly in many developing countries around the world, causing a considerable public health issue (2).

Nearly three decades ago, J.A. Butill proposed sub-classifying CRC depending on the anatomical site, either proximal (right) or distal (left) to the splenic flexure (3). Subsequent research has observed distinct differences in epidemiology and pathological features according to primary tumor location in CRC. In 2000, H. Elsaleh found that the tumor site is associated with survival benefit from adjuvant chemotherapy in CRC (4). This researcher discovered that patients with right-sided tumors have better survival benefits from adjuvant chemotherapy than patients with left-sided tumors. In addition, the frequency of MSI was much higher in right-sided tumors than in left-sided tumors (5,6). It is now well established by a variety of studies that primary tumor location affects the outcome of the chemotherapy and immunotherapy of CRC patients in a large-scale population, and tumor location is a high-risk parameter for prognosis in specific stages. There is a general consensus that primary tumor location plays an important role in CRC development. We could even define right-sided and left-sided tumors as two different diseases that need different treatments (7). This influence of tumor location may be due to differences in embryological development. Specifically, the right side of the colon has historically been understood to be derived from the embryological midgut, and the left colon arises from the embryological hindgut. The transverse colon is composed of parts of both structures. These different origins could result in various clinical traits.

However, the underlying molecular mechanism governing those different behaviors and outcomes has not been fully...
research describing the use of the Chip-seq or RNA-seq sequencing technology, we currently have abundant published data as a robust measure of variability. We will be highly correlated. We chose the 75% most varying genes without notable variance in expression between patients to detect the same mRNA, the average value of the probes using Bioconductor in R. If multiple probes were applied to the same location of the genome, the average value was used. The probe information was then transformed into the official gene symbols using Bioconductor in R. If multiple probes were applied to detect the same mRNA, the average value of the probes was used. The genes that were not differentially expressed between samples have to be excluded from WGCNA, as two genes without notable variance in expression between patients will be highly correlated. We chose the 75% most varying genes to construct the weighted gene coexpression networks. Specifically, the median absolute deviation (MAD) was used as a robust measure of variability.

In addition, the level three RNA-sequencing data of both colon adenocarcinoma (COAD) and rectum adenocarcinoma (READ) patients were downloaded from The Cancer Genome Atlas data portal (TCGA; http://cancergenome.nih.gov/). In contrast to ChIP-sequencing data, we used the voom function in package limma to normalize the TCGA data and create an expression matrix for samples for which the detailed clinical information was available. Further preprocessing steps included the removal of control samples and the genes with zero counts in more than 80% of samples. As mentioned before, genes that are not differentially expressed between samples must be excluded; thus, we chose the top 12,000 genes with the highest MAD for the network building. Fig. 1 depicts a flow chart for the bioinformatic analysis.

**Materials and methods**

**Data collection.** The raw expression data of GSE39582 (11) and GSE14333 (12) were retrieved from the Gene Expression Omnibus database (http://www.ncbi.nlm.nih.gov/geo/), both based on the platform of GPL570 Affymetrix Human Genome U133 Plus 2.0 Array. We used the Affy package in R to transform the CEL files of the tumor samples into an expression matrix (13). To improve the data quality, we used the k-nearest neighbors algorithm (k-NN) from the impute package in R to impute the missing expression data (14). Meanwhile, the robust multiarray average algorithm (RMA) was utilized to adjust the data for potential batch effects and for background correcting (15). Prior to WGCNA analysis, we filtered out the probes that were absent in all samples. The probe information was then transformed into the official gene symbols using Bioconductor in R. If multiple probes were applied to detect the same mRNA, the average value of the probes was used. The genes that were not differentially expressed between samples had to be excluded from WGCNA, as two genes without notable variance in expression between patients will be highly correlated. We chose the 75% most varying genes to construct the weighted gene coexpression networks. Specifically, the median absolute deviation (MAD) was used as a robust measure of variability.

In addition, the level three RNA-sequencing data of both colon adenocarcinoma (COAD) and rectum adenocarcinoma (READ) patients were downloaded from The Cancer Genome Atlas data portal (TCGA; http://cancergenome.nih.gov/). In contrast to ChIP-sequencing data, we used the voom function in package limma to normalize the TCGA data and create an expression matrix for samples for which the detailed clinical data are available (16,17). The voom method estimates the mean variance of the log counts and generates a precision weight for each observation. Thus, the WGCNA workflows originally developed for microarray analysis can be used on the RNA-seq data. Further preprocessing steps included the removal of control samples and the genes with zero counts in more than 80% of samples. As mentioned before, genes that are not differentially expressed between samples must be excluded; thus, we chose the top 12,000 genes with the highest MAD for the network building. Fig. 1 depicts a flow chart for the bioinformatic analysis.

**Construction of weighted gene coexpression networks.** The R package ‘WGCNA’ was used in our study to construct a gene coexpression network (10). After data collection and normalization, it is crucial that outliers be excluded. However, it was difficult to distinguish outlying samples in a dendrogram when the number of samples was large. To solve this problem, we used the standardized connectivity ($Z. K$) method recommended by WGCNA authors with the default threshold, $Z. K$ score £2. After filtering out the outlying samples, expression data were tested to determine whether the samples and genes were good using the integrated function in the WGCNA package.

After filtering out the outliers and bad samples in the dataset, the next step of WGCNA is to build a scale-free network. In a scale-free network, several nodes, which are called hub nodes, are highly connected to other nodes in the network (18). In our study, we use the unsigned coexpression measure, which means that the positive correlation and negative correlation are equal. We constructed the gene coexpression network using the following steps.

First, we need a soft thresholding power $\beta$ to which coexpression similarity is raised to calculate adjacency. By raising the absolute value of the correlation to a power $\beta \geq 1$ (soft thresholding), the weighted gene coexpression network construction emphasizes high correlations at the expense of low correlations. To determine the best soft threshold power, scale independence and average connectivity degree of modules with different power values were calculated by the gradient method. We selected the power $\beta$ to ensure that the coexpression network was a ‘scale-free’ network, which was biologically close to reality. Moreover, to minimize the effects of noise and spurious associations, we subsequently constructed the Topology Overlap Matrix (TOM) from the adjacency matrix and calculated the corresponding dissimilarity (1-TOM), as well (19).

In the same way, the second coexpression network was built from TCGA data.

**Identification of coexpression modules.** The traditional static tree cut method exhibits suboptimal performance on complicated dendrograms. In WGCNA, we tend to use the dynamic tree cut method by hierarchically clustering genes using the dissimilarity matrix (1-TOM) (20). The minimal size of a module was set as 30, and modules with high similarity were identified by clustering and then merged together with a height cut-off of 0.25. To determine whether the modules are reproducible, we tested the preservation of all modules with an independent gene expression dataset, GSE14333. We used the module preservation function (number of permutations set to 100) integrated in the WGCNA package to calculate the
Enrichment Analysis.

### Validation of the hub genes
We applied Gene Expression Profiling Interactive Analysis (GEPIA) (http://gepia.cancer-pku.cn/) to detect the difference in expression levels of each hub gene between tumor and normal tissues in both the COAD and READ datasets from TCGA (22). To further validate our method, correlation plots between hub genes were generated by GEPIA, as well.

### Coexpression validation with qPCR
Twenty non-selected CRC samples were applied to perform qPCR to validate coexpression of PLAG1 like zinc finger 2 (PLAGL2) and protein O-fucosyltransferase 1 (POFUT1). These experimental samples were collected at the Sir Run Run Shaw Hospital of Zhejiang University between January 2004 and December 2006. After total RNA was isolated from tumor specimens using Trizol reagent (Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA), RNA was quantified by NanoDrop 2000c spectrophotometer (Thermo Fisher Scientific, Inc.) and reverse transcribed using RNeasy Mini Kit (Takara, Kyoto, Japan) according to the manufacturer's protocols. Quantitative real-time PCR was executed with SYBR Green Master Mix (Takara). Relative expression levels were calculated with $2^{-\Delta\Delta Cq}$ formula (23). Expression of mRNA was standardized according to β-actin. The primers were used as follows: β-actin_fwd, ACTTCTCCAGGCTTCTCTTC and β-actin_rev, CGTCTACTCTGCTGCTG; PLAGL2_fwd, GAGTCGAAGTAGGGACAAATGT and PLAGL2_rev, TGA GGGGAGCTATATGTGCTCT; POFUT1_fwd, AACCAG GCCGATCACCCTTGG and POFUT1_rev; GTTGGTGAA AGGAGGCTGTTGTG. The primers were designed on online tools (https://www.genscript.com/tools/real-time-pcr-tagman-primer-design-tool) and these were synthesized by Shanghai Generay Biotech Co. Ltd. (Shanghai, China).

### Survival analysis
We performed survival analysis for hub genes using the GSE39582 dataset because of its complete overall survival information. Kaplan-Meier analysis and log-rank test were performed to evaluate the association between hub gene expression and patient survival in left- and right-sided CRC, respectively. This procedure utilized the survival package in R (24), and the Kaplan-Meier survival curves with the at-risk table were drawn using the survminer package (25).

### Gene set enrichment analysis
To identify the possible pathway through which hub genes may play a part in the development of CRC, the expression data from GSE14333 was also used to perform Gene Set Enrichment Analysis (GSEA). The expression data of 290 cases were uniformly divided into two groups according to each hub gene's expression value.

We used the GSEA-p 2.0 software to conduct the enrichment analysis (26). For configuration, ‘c2.cp.kegg.v6.2.symbols.gmt’ from the Molecular signatures database (MSigDB) 3.0 (27) was used as the gene set, and the permutation number was set to 1,000 as the default. Finally, P-values <0.05 and FDR <25% were considered to be statistically significant (28).

### Statistical analysis
In this study, we used Pearson correlation coefficient to measure the strength of the relationship between the variables. The coexpression of mRNA expression level of PLAGL2 and POFUT1 was presented by linear
regression model. Coefficient of determination was calculated and presented. The independent samples t-test was performed for data comparison in GEPIA validation part. All statistical analyses were performed using R program. P-values <0.05 was considered to indicate a statistically significant result.

Results

Data preprocessing. A workflow of the study is shown in Fig. 1. The dataset GSE39582 contained 585 samples from CRC patients, including 19 normal tissue samples and 566 tumor samples, while GSE14333 had 290 primary CRC tissues. We used the GSE39582 data to build our network and GSE14333 for validation purposes. After data collection, a total of 436 tumor samples with complete clinical information from GSE38582 were obtained. The clinical information of GSE39582 is shown in the clustering dendrogram with the trait heatmap (Fig. 2).

For genes, we transformed the 50,362 probe ids into 22,880 official gene symbols and calculated the median absolute deviation (MAD) of each gene in all samples mentioned above. The three-quarters genes, which equals 17,160, that have the highest MAD were used to construct the final expression network. This step also ensured that the median absolute deviation was not 0, thereby avoiding further errors when constructing the gene coexpression network.

In the meantime, the preprocess of TCGA RNA-seq data was different. We combined the COAD and READ data into one matrix, which has a total of 19,754 genes and 644 samples. Then, we deleted 22 repeat samples and filtered out the genes with zero expression in more than 80% of samples. After voom normalization, we chose the top 12,000 genes with the highest MAD for further analysis.

Network construction and module identification. In choosing the best threshold, we calculated the network topology for soft-thresholding powers from 1 to 20. As shown in Fig. 3A, power value 5, which was the lowest power for the scale-free topology fit index on 0.9, was selected. Afterward, we checked the mean connectivity (Fig. 3B) and double-checked the scale-free topology $R^2$ with a linear regression plot (Fig. 3C). Fig. 3D contains a histogram of the frequency of connections. A highly skewed histogram is said to approximate a scale-free network.

The coexpression similarity matrix was then transformed into the adjacency matrix by choosing 5 as a soft threshold, and a topological overlap matrix (TOM) was subsequently computed. Using the dynamic tree cut method, a total of 38 modules were identified. The modules with higher correlation than 0.75 were subsequently merged, resulting in 31 modules at last (Fig. 4). The gray module includes genes that were not assigned to any gene modules.

In the network built by the TCGA dataset, the soft threshold was 7 by the calculation (Fig. 5A). Ultimately, 26 gene modules were recognized (Fig. 5C).

Identification of key modules. To analyze the relationship between gene modules and sample clinical information, we employed module eigengene (ME) as the average gene expression level of the corresponding modules. It can be considered a representative of the gene expression profiles in a module. The correlations between module eigengene and clinical phenotypes in GSE39582 were calculated and plotted as a labeled heatmap (Fig. 6). The red module and orange module were significantly associated with tumor location.

We calculated gene significance based on the correlation of a gene expression profile with the samples’ location traits. Then, the module significance was defined as the average absolute value of the gene significance of all genes in one module. As shown in Fig. 7A, the red and orange modules had
Figure 3. Analysis of the network topology for adjacency matrix weighting parameters (power). (A and B) The x-axis represents weighting parameters (power). The y-axis represents the scale-free fitting index and connectivity for each power. (C) The regression line with an index of $R^2=0.92$ when choosing the power of 5. The CRC network exhibits a scale-free topology. (D) The histogram of $k$ when choosing the power of 5. CRC, colorectal cancer.

Figure 4. Cluster dendrogram produced by average linkage hierarchical clustering of genes based on topological overlap matrix (TOM). Each branch in the dendrogram is a line that represents a single gene. Each color indicates a single module that contained closely conserved genes.
Figure 5. TCGA data using the same method locating a dark-gray module that is highly correlated with tumor location. (A) Analysis of scale-free topology model fit vs. the candidate soft threshold powers. (B) Gene significance (y-axis) vs. module membership (x-axis) plotted for dark-gray module in the TCGA dataset. (C) Cluster dendrogram based on topological overlap matrix (TOM) in the TCGA dataset. (D) Module-trait relationships heatmap in the TCGA dataset indicates the dark-gray module is highly related to the tumor location. TCGA, The Cancer Genome Atlas.
Figure 6. Module-trait relationships were evaluated by WGCNA using GSE39582 microarray analysis comprising 431 human colorectal cancer samples. Gene modules are denoted by an arbitrary color name. Bins show the Pearson correlation value between gene expression levels of each module within the noted clinical traits and P-values. A value of 1 (red) and -1 (blue) both quantify the strongest correlation, and 0 (white) quantifies no correlation. WGCNA, weighted gene coexpression analysis; CIMP, CpG island methylator phenotype; MMR, mismatch repair.

Figure 7. (A) Bar plot of mean gene significance across genes associated with tumor location in the module. (B) Calculations of module preservation statistics between GSE39582 and the independent dataset GSE14333. The dashed lines mark thresholds at Z=2 and Z=10, according to which >10 suggests strong evidence for preservation and >2 moderate evidence for preservation. A Z summary value <2 indicates no preservation. (C) Gene significance (y-axis) vs. module membership (x-axis) plotted for red module in the GSE39582 dataset. In this module, genes with high module membership tended to have high gene significance. The genes with the highest gene significance are labeled blue.
The genes in common are indicated in bold print. PLAGL2, PLAG1-like zinc finger 2; POFUT1, protein O-fucosyltransferase 1.

Table I. Twelve hub genes are found in the GSE39582 and TCGA dataset.

| Hub gene      | Ensemble ID | Name                                           | Cytogenetic location |
|---------------|-------------|------------------------------------------------|----------------------|
| PLAGL2        | 5326        | PLAG1-like zinc finger 2                       | 20q11                |
| POFUT1        | 23509       | Protein O-fucosyltransferase 1                 | 20q11                |
| TIT1          | 9675        | TELO2 interacting protein 1                    | 20q11                |
| ASXL1         | 171023      | Additional sex combs-like 1                    | 20q11                |
| AAR2          | 25980       | AAR2 splicing factor homolog                   | 20q11                |
| PIGU          | 128869      | Phosphatidylinositol glycan anchor biosynthesis class U | 20q11               |
| STAU1         | 6780        | Staufen double-stranded RNA binding protein 1  | 20q11                |
| DYNLRB1       | 83658       | Dynein light chain roadblock-type 1            | 20q11                |
| NELFCD        | 51497       | Negative elongation factor complex member C/D  | 20q11                |
| ZSWIM3        | 140831      | Zinc finger SWIM-type containing 3             | 20q11                |
| MOCS3         | 27304       | Molybdenum cofactor synthesis 3                | 20q11                |
| TM9SF4        | 9777        | Transmembrane 9 superfamily member 4           | 20q11                |

The genes in common are indicated in bold print. PLAGL2, PLAG1 like zinc finger 2; POFUT1, protein O-fucosyltransferase 1.

To determine the module's reproducibility, module preservation analysis was performed using an independent dataset GSE14333. As we can see in Fig. 7B, modules below the green dashed line (Z summary < 10) are poorly preserved, while the modules above the line are well-preserved in the CRC tissues. The red module, according to the preservation test, is highly preserved in CRC; however, the orange module showed moderate preservation. Thus, we chose the red module for further analysis.

Again, the same method was applied to TCGA data, locating a similar dark-gray module (Fig. 5D).

Identification of hub genes in the key module. There were 865 genes in the GSE39582 red module. After plotting the gene significance against module membership, we observed that genes with higher module memberships tended to have higher gene significance in this module (Fig. 7C). We used a relatively high criterion to select hub genes: The absolute value of gene significance > 0.4 and module membership > 0.8. Six hub genes were successfully identified. The genes with the highest gene significance were found to be POFUT1 and PLAGL2, which are labeled in blue print in Fig. 7C.

Meanwhile, in the TCGA dark-gray module, we used the absolute value of gene significance > 0.3 to filter out 8 hub genes (Fig. 5B). After combining two datasets, we determined that there were 12 possible hub genes, 2 of which are in common (Table I).

Validation of the hub genes. We concentrated on PLAGL2 and POFUT1 because of their high gene significance and their presence in both datasets. We then evaluated their expression with the online TCGA-based tool GEPIA. PLAGL2 and POFUT1 were found to be significantly differentially expressed between tumor and normal tissue in both the COAD and READ datasets (Fig. 8A and B). We also performed a correlation analysis between PLAGL2 and POFUT1. The plot shows that the Pearson correlation coefficient is tightly correlated to 0.9 in CRC (Fig. 8C).

We utilized quantitative polymerase chain reaction (qPCR) to measure the RNA expression of PLAGL2 and POFUT1 in CRC samples. PLAGL2 had a high positive correlation with POFUT1 according to the qPCR results (Fig. 8D).

Survival analysis and gene set enrichment analysis. For survival analysis, Kaplan-Meier curves were drawn for PLAGL2 and POFUT1 in both proximal and distal CRC (Fig. 9). Although the log-rank P-value of all the analyses was > 0.05 (not statistically significant), we still compared the results from different parts of the colon. In proximal CRC samples, there was a clear trend that high PLAGL2 and POFUT1 expression is related to adverse prognosis in CRC patients. However, in distal CRC samples, the expression of POFUT1 was not related to survival, and the high expression of PLAGL2 was even associated with poor survival.

We also performed a Gene Set Enrichment Analysis based on the expression level of PLAGL2 and POFUT1. As shown in Fig. 10, these two genes share a similar enriched KEGG pathway: Glycosylphosphatidylinositol GPI anchor biosynthesis and peroxisome and selenoaminoacid metabolism.

Discussion

We have only recently (over the past 5 to 10 years) determined that the parts of the colon derived from the midgut and the hindgut are different. Numerous studies have investigated this subject. In 2015, Guinney and colleagues published a leading article in Nature Medicine. These researchers divided CRC into 4 well-defined subtypes by their gene expression patterns and discovered that certain types are mainly located on one side of the colon rather than being randomly distributed (29). Moreover, behind this phenomenon, there must be gene expression patterns that we can investigate.

The information captured by microarray or RNA-seq experiments is notably richer than a list of differentially
expressed genes. Microarray and RNA-seq data are more completely represented by considering the relationships between measured transcripts, which can be assessed by pair-wise correlations between gene expression profiles. Prior bioinformatics studies have noted the importance of gene coexpression networks in various types of cancers. However, many studies used differentially expressed genes to build the coexpression network. It is not recommended by the author of WGCNA, because filtering genes by differential expression will lead to a set of correlated genes that will essentially form a single (or a few highly correlated) module. Since nonvarying genes usually represent noise, we used genes with the top 75% MAD to improve the robustness and confidence of the present analysis.

In this study, we used three different datasets to analyze the gene expression patterns of CRC. These datasets have different patient information which leads to the different clinical features. However, when we clustered every gene into the modules by WGCNA, we did not use the clinical features of any kind. Considering the number of samples in these datasets are large, together with the results from the module preservation test, we could assume the key module we identified is universal. An interesting part in the module-to-trait relationship heatmap is that the modules with high correlation with tumor location also highly correlate with mismatch repair (MMR) (30) and the CpG island methylator phenotype (CIMP) (31) status. In the last decade, extensive studies have studied this problem and found that tumors with deficient mismatch repair (microsatellite instability-high, MSI-H) and the CpG island methylator phenotype are mostly located on the right side of the colon, which matches our sample traits from GSE39582. Although dMMR or CIMP+ samples are not the majority in the dataset, this tendency may cause a bias that the correlation between tumor sites and the key module is mainly from MMR and CIMP status or other clinical features. To diminish the bias, we also used module significance to define the correlation between modules and tumor site.

Figure 8. (A and B) Expression of PLAGL2 and POFUT1 in CRC and normal tissues from GEPIA. (C) The gene expression correlation between PLAGL2 and POFUT1 from GEPIA. (D) qPCR results indicate a strong relationship between PLAGL2 and POFUT1 at the RNA level. CRC, colorectal cancer; GEPIA, Gene Expression Profiling Interactive Analysis; PLAGL2, PLAG1 like zinc finger 2; POFUT1, protein O-fucosyltransferase 1.
phonotype, both in GSE39582 and TCGA. Although other clinical information was slightly different, the key modules we found in both datasets were similar, which had several common hub genes.

The fundamental theory of WGCNA is that we assume genes interact with each other in a scale-free network. In this way, the hub genes play more important roles in the whole module than other genes. Among the cluster of genes that have a strong relationship with the tumor location of CRC, 12 hub genes with high significance were identified in the GSE39582 and TCGA datasets, which may have contributed most to the distinct behaviors. Some of the genes have been found to be critical in CRC development and prognostic biomarkers in specific stages from other publications (32,33).

As we examined these hub genes, we found they are all located on the long arm of chromosome 20 (20q11). Previous studies have confirmed that the copy number gain in 20q (mostly in 20q11 and 20q13) occurs in more than 65% of CRC patients (34). As a consequence of copy number gain of 20q, multiple genes mapping at the chromosome 20q amplicon contribute to colorectal adenoma to carcinoma progression (35). In our study here, we identified several coexpressed hub genes in 20q11 that may be attributed to the differential features of proximal and distal CRC. However, in the 12 hub genes displayed in Table I, PLAGL2 and POFUT1 were not only presented in the two datasets, but also showed the highest gene significance. We believe that they are more representative than other genes, thus we focused on them for further exploration.
PLAGL2 encodes a zinc finger transcription factor that contains seven C2H2 zinc finger motifs that exhibit DNA binding and transcriptional activation activity. Recently, Li et al found that overexpression of PLAGL2 transcriptionally activates Wnt6 and promotes cancer development in CRC (36). PLAGL2 activates the Wnt/β-catenin pathway as a transcription factor by binding to the promoter region of Wnt6.

POFUT1, on the other hand, is essential for Notch signal transduction in mammals. In 2018, Du et al discovered that POFUT1 promotes CRC development through the activation of Notch1 signaling (37). Another study by Chabanais et al also confirmed that POFUT1 is overexpressed in CRC from stage I, and its high expression is associated with the metastatic process (38). In addition, these researchers found that POFUT1 overexpression is markedly associated with rectal location, which corroborates our finding.

In all the studies reviewed in this article, PLAGL2 and POFUT1 are recognized as oncogenes that promote or at least are associated with CRC development. Furthermore, these genes are highly correlated based on our qPCR result and correlation analysis from the TCGA dataset. As we found in the GEPIA (Fig. 7), these genes were both significantly differentially expressed between tumor and normal tissue in both the COAD and READ datasets.

Moreover, our survival analysis, despite not being statistically significant, found that there were different results between left- and right-sided CRC for PLAGL2 and POFUT1 (Fig. 9). In proximal CRC patients, the red curves, which represent the low expression of PLAGL2 and POFUT1, were beneath the blue ones, and the log-rank P-value was at the verge of significance. However, in distal CRC samples, the relationship of PLAGL2 and POFUT1 expression and survival were vague and even reversed. This research showed a considerable difference between left- and right-sided survival with regard to PLAGL2 and POFUT1, which indirectly indicates that the expression of the genes is related to the tumor location in CRC patients.

According to our GSEA results, these two genes may also take effect through glycosylphosphatidylinositol (GPI) anchor biosynthesis and peroxisome and selenoamino acid metabolism pathways. When we examined the hub genes in Table I, we found that one of the hub genes from GSE39582 is associated with one of the pathways mentioned above. PIGU is a component of the GPI transamidase complex that may be involved in the recognition of either the GPI attachment signal or the lipid portion of GPI. This finding confirms that the hub genes’ functions are as tightly connected as their expression levels, which is the foundation of the WGCNA theory. However, there are few articles discussing the association of this gene with the development of CRC. This subject warrants further investigation in the future.

Another thorough study of gene expression in colon cancer from Slattery et al used Ingenuity Pathway Analysis (IPA) to determine networks associated with deregulated genes (39). In his study, PLAGL2 and POFUT1 were found to be differentially expressed genes in both MSI and CIMP status comparisons. In other words, we could assume that these genes may be related to the anatomical site of CRC through MSI and CIMP status.
The findings of these studies indicate that the hub genes that we found are oncogenes that may relate to the sidedness of CRC. Notably, PLAGL2 and POFTU1 are the centers of the module and are differentially expressed between normal and tumor tissues, which makes them promising biomarkers.

As Dr Alan P. Venook noted in Clinical Advances in Hematology & Oncology (40), what matters is not the sidedness of the tumor because sidedness is simply a surrogate for the types of tumors that tend to occur on that side. Our work, while preliminary, suggests that a weak link may exist between the oncogenesis triggered by these genes and the primary site of CRC. However, the underlying mechanism requires further investigation.

Acknowledgements

This research represents partial fulfillment of the requirements for a Master degree for YL and WZ.

Funding

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Availability of data and materials

The datasets analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions

YL, BX, XH and HZ conceived and designed the study. BB collected the data. YL and BX performed the bioinformatics analysis. LS and WZ performed the experiments. YL and BX wrote the paper. BB, LS, WZ, XH and HZ reviewed and edited the manuscript. All authors read and approved the manuscript. All authors declare that they have no competing interests.

Competing interests

The authors declare that they have no competing interests.

References

1. Bray F, Ferlay J, Soerjomataram I, Siegel RL, Torre LA and Jemal A: Global cancer statistics 2018: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin 68: 394-424, 2018.
2. Arnold M, Sierra MS, Laversanne M, Soerjomataram I, Jemal A and Bray F: Global patterns and trends in colorectal cancer incidence and mortality, Gut 66: 683-691, 2017.
3. Buttil J: Colorectal cancer: Evidence for distinct genetic categories based on proximal or distal tumor location. Ann Intern Med 113: 779-788, 1990.
4. Elsahel H, Joseph D, Griev F, Zeps N, Spry N and Lacopetta B: Association of tumour site and sex with survival benefit from adjuvant chemotherapy in colorectal cancer. Lancet 355: 1745-1750, 2000.
5. Deng G, Kakar S, Tanaka H, Matsuoka K, Miura S, Slesiemger MH and Kim YS: Proximal and distal colorectal cancers show distinct gene-specific methylation profiles and clinical and molecular characteristics. Eur J Cancer 44: 1290-1301, 2008.
6. Minyo P, Zlobe I, Peterson M, Terracciano L and Lugli A: Characterization of rectal, proximal and distal colon cancers based on clinicopathological, molecular and protein profiles. Int J Oncol 37: 707-718, 2010.
7. Lee GH, Malletzis G, Askari A, Bernardo D, Al-Hassi HO and Clark SK: Is right-sided colon cancer different to left-sided colorectal cancer? - a systematic review. Eur J Surg Oncol 41: 300-308, 2015.
8. Glebov OK, Rodriguez LM, Nakahara K, Jenkins J, Clait J, Humbyrd CJ, DeNobile J, Soballe P, Simon R, Wright G, et al: Distinguishing right from left colon by the pattern of gene expression. Cancer Epidemiol Biomarkers Prev 12: 755-762, 2003.
9. Birkenkamp-Demtroder K, Olesen SH, Sørensen FB, Laurberg S, Sorensen RA, Larsen J, Olesen BM, Kehlet H, et al: Metastasis-related gene expression in colorectal cancer of the caecum versus the sigmoid and rectosigmoid. Gut 54: 374-384, 2005.
10. Langfelder P and Horvath S: WGCNA: An R package for weighted correlation network analysis. BMC Bioinformatics 9: 559, 2008.
11. Marisa L, de Reynies A, Duval A, Seljes J, Gaub MP, Vescovlo E, Etienne-Grimaldi MC, Chiappa R, Guenot D, Ayadi M, et al: Gene expression classification of colon cancer into molecular subtypes: Characterization, validation, and prognostic value. PLoS Med 10: e1001453, 2013.
12. Jorissen RN, Gibbs P, Christie M, Prakash S, Lippton L, Desai J, Kerr D, Aaltonen LA, Arango D, et al: Metastasis-associated gene expression changes predict poor outcomes in patients with dukes Stage B and C colorectal cancer. Clin Cancer Res 15: 7642-7651, 2009.
13. Gautier L, Cope L, Bolstad BM and Irizarry RA: Affy-analysis of Affymetrix GeneChip data at the probe level. Bioinformatics 20: 307-315, 2004.
14. Hastie T, Tibshirani R, Narasimhan B and Chu G: Impute: Imputation for microarray data. Bioinformatics 17: 520-525, 2001.
15. Irizarry RA, Hobbs B, Collin F, Beazer-Barclay YD, Antonellis KJ, Scherf U and Speed TP: Exploration, normalization, and summaries of high density oligonucleotide array probe level data. Biostatistics 4: 249-264, 2003.
16. Ritchie ME, Phipson B, Wu D, Hu Y, Law CW, Shi W and Smyth GK: Limma powers differential expression analyses for RNA-sequencing and microarray studies. Nucleic Acids Res 43: e47, 2015.
17. Law CW, Chen Y, Shi W and Smyth GK: Voom: Precision weights unlock linear model analysis tools for RNA-seq read counts. Genome Biol 15: R29, 2014.
18. Zhang B and Horvath S: A general framework for weighted gene co-expression network analysis. Stat Appl Genet Mol Biol 4: 17, 2005.
19. Yip AM and Horvath S: Gene network interconnectionness and the generalized topological overlap measure. BMC Bioinformatics 8: 22, 2007.
20. Langfelder P, Zhang B and Horvath S: Defining clusters from a hierarchical cluster tree: The dynamic tree cut package for R. Bioinformatics 24: 719-720, 2008.
21. Langfelder P, Luo R, Oldham MC and Horvath S: Is my network module preserved and reproducible? PLoS Comput Biol 7: e1001057, 2011.
22. Tang Z, Li C, Kang B, Gao G, Li C and Zhang Z: GEPIA: A web server for cancer and normal gene expression profiling and interactive analyses. Nucleic Acids Res 45: W98-W102, 2017.
23. B老化 KJ and Schmittgen TD: Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) method. Methods 25: 402-408, 2001.
24. Therneau TM and Lumley T: Package ‘survival’. Survival Published on CRAN, 2014.
25. Kassambara A, Kosinski M and Biecek P: Survminer: Drawing survival curves usingggplot2. R package version 0.3.1 2017.
26. Subramanian A, Kuehn H, Gould J, Tamayo P and Mesirov JP: GSEA-P: A desktop application for gene set enrichment analysis. Bioinformatics 23: 3251-3253, 2007.
27. Liberzon A, Subramanian A, Pinchback R, Thorvaldsdóttir H, Tamayo P and Mesirov JP: Molecular signatures database (MSigDB) 3.0. Bioinformatics 27: 1739-1740, 2011.
28. Benjamini Y and Hochberg Y: Controlling the false discovery rate: A practical and powerful approach to multiple testing. J R Stat Soc: Series B (Methodological) 57: 289-300, 1995.
29. Guinney J, Dienstmann R, Wang X, de Reynies A, Schlicker A, Soneson C, Marisa L, Roeppman P, Nyamundanda G, Angelino P, et al: The consensus molecular subtypes of colorectal cancer. Nat Med 21: 1350-1356, 2015.
30. Boland CR and Goel A: Microsatellite instability in colorectal cancer. Gastroenterology 138: 2073-2087.e3, 2010.
31. Toyota M, Ahuja N, Ohe-Toyota M, Herman JG, Baylin SB and Issa JP: CpG island methylator phenotype in colorectal cancer. Proc Natl Acad Sci USA 96: 8681-8686, 1999.
32. Damas ND, Marcattia M, Côme C, Christensen LL, Nielsen MM, Baumgartner R, Gylling HM, Maglieri G, Bundsten CF, Seemann SE, et al: SNHG5 promotes colorectal cancer cell survival by counteracting STAU1-mediated mRNA destabilization. Nature Commun 7: 13875, 2016.
33. Song S, Li D, Yang C, Yan P, Bai Y, Zhang Y, Hu G, Lin C and Li X: Overexpression of NELFCD promotes colorectal cancer cells proliferation, migration, and invasion. Onco Targets Ther 11: 8741-8750, 2018.
34. Sillars-Hardebol AH, Carvalho B, Tijsen M, Bliën JA, de Wit M, Delis-van Diemen PM, Pontén F, van de Wiel MA, Fijneman RJ and Meijer GA: TPX2 and AURKA promote 20q amplicon-driven colorectal adenoma to carcinoma progression. Gut 61: 1568-1575, 2012.
35. Carvalho B, Postma C, Mongera S, Hopmans E, Diskin S, van de Wiel MA, van Criekinge W, Thas O, Matthai A, Cuesta MA, et al: Multiple putative oncogenes at the chromosome 20q amplicon contribute to colorectal adenoma to carcinoma progression. Gut 58: 79-89, 2009.
36. Li N, Li D, Du Y, Su C, Yang C, Lin C, Li X and Hu G: Overexpressed PLAGL2 transcriptionally activates Wnt6 and promotes cancer development in colorectal cancer. Onco Rep 41: 875-884, 2019.
37. Du Y, Li D, Li N, Su C, Yang C, Lin C, Chen M, Wu R, Li X and Hu G: POFUT1 promotes colorectal cancer development through the activation of Notch1 signaling. Cell Death Dis 9: 995, 2018.
38. Chabanais J, Labrousse F, Chaunavel A, Germot A and Maftah A: POFUT1 as a promising novel biomarker of colorectal cancer. Cancers 10: pii: E411, 2018.
39. Slattery ML, Pellatt DF, Mullany LE, Wolff RK and Herrick JS: Gene expression in colon cancer: A focus on tumor site and molecular phenotype. Genes Chromosomes Cancer 54: 527-541, 2015.
40. Venook AP: Right-sided vs left-sided colorectal cancer. Clin Adv Hematol Oncol 15: 22-24, 2017.

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International (CC BY-NC-ND 4.0) License.