Changes of Intestinal Microflora in Colorectal Cancer Patients after Surgical Resection and Chemotherapy

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Objective. The change of bacterial flora structure in colorectal cancer (CRC) patients after treatment is not clear. The aim of this study was to explore the change and function of intestinal microflora in CRC before and after treatment.

Method. The 16S conserved region V3+V4 of intestinal flora obtained from CRC patients was sequenced and analyzed. Alpha and beta diversity indices were used to analyze the abundance and structure of gut flora. FAPROTAX, BugBase, and Tax4Fun software were used to analyze the species phenotypes and Kyoto Encyclopedia of Genes and Genomes Ontology (KO) function pathways.

Results. Total abundance and structure of species in CRC patients were significantly increased compared with healthy people (control group) \((P < 0.05)\), but there was no significant difference between CRC patients before and after treatment \((P > 0.05)\). There was significant difference in relative abundance of bacteria at different levels (phylum, class, order, family, genus, and species) between the CRC group with after operation (CRC_O group) and chemotherapy (CRC_C group) treatment, particularly Prevotellaceae_UCG-001, Akkermansia, Fusicatenibacter, Tyzzerella_4, Megamonas, etc. in genus level. The KO function analysis showed that most of the bacteria with differences were mainly involved in the biosynthesis of lipopolysaccharide (Megamonas, Megasphaera, and Ruminococcus torques group), protein digestion and absorption, renin-angiotensin system pathway (Akkermansia, Eubacterium_ruminantium_group, and Eubacterium_moitatum_group genus), adipokine signaling pathway and pexiosome pathway (Tyzzerella_4, Phascolarctobacterium, Ruminococcus_gnavus_group), and so on. Conclusion. The abundance of intestinal microflora in CRC patients was increased significantly contrasted to healthy people, and surgery and chemotherapy were hard to reduce this phenomenon. Megamonas was involved in lipopolysaccharide biosynthesis and carcinogenesis in colorectal cancer. Surgery and drug treatment did not reduced lipopolysaccharide biosynthesis but increased the number of probiotic Akkermansia population and reduced the pathogenic bacteria Tyzzerella_4, participate in adipokine signaling pathway, and affect metabolism.

1. Introduction

The incidence of colorectal cancer (CRC) ranks third in the total number of cancer patients in the world, and the mortality rate ranks fourth in the global cancer-related deaths [1]. In China, 376000 new cases and 191000 deaths of CRC occur in 2015 [2], which is the fourth largest tumor in China. With the continuous improvement of living standards, and the intake of high-fat and high sugar foods, the incidence rate of CRC is increasing year by year. The onset age of CRC is gradually younger, and there are no obvious clinical symptoms in the early stage. Advanced CRC metastasizes to distal organs, leading to important organ failure and death, which seriously threatens human health [3]. Currently, the treatment strategies of CRC include surgical resection, radiotherapy, chemotherapy, targeted therapy, and emerging immunotherapy. The effective rate of surgery combined with radiotherapy and chemotherapy is as high as 70% to 80%, and the 3-year overall survival is up to 53%. With the development of tumor, the survival rate of CRC patients is also declining [4]. In addition, 20%-30% of CRC patients receiving treatment are at risk of recurrence [5]. To evaluate the risk of formation and recurrence of colorectal cancer and to seek effective treatment strategies for intestinal cancer are the focus of clinical research.
In recent years, many studies have shown that intestinal flora disorder is an important factor leading to CRC in addition to genetics, environment, and diet. In fact, a large number of studies have confirmed the significant changes of intestinal flora composition between CRC patients with normal one. Several bacterial species have proinflammatory and carcinogenic characteristics and are involved in the occurrence of colorectal cancer [6]. Normally, the intestinal tract of human body is rich in four bacterial groups including Hirsutella, Bacteroides, Proteus, and Actinomycetes, which regulate the body’s normal nutrient absorption, metabolism, and immunity. In CRC patients, Gram-negative bacteria are enriched, while Gram-positive bacteria and Actinomycetes are decreased. For example, a large number of pathogenic bacteria such as Porphyromonas, Clostridium, Staphylococcus, Ackermann, and Methanogens genus are detected in the intestinal tract of CRC patients, while Bifidobacterium, Lactobacillus, Ruminococcus, Bacillus, Roseburia, and Treponema genus are poor. Some scholars found that the species abundance of Clostridium and Fusobacterium is negatively correlated with tumor count. Animal model experiments show that beneficial bacteria such as high level of Clostridium implanted into the intestinal tract effectively reduce the burden of CRC [1, 7, 8].

With the development of molecular technology and the improvement of microbial genomics, the causal relationship between cancer and intestinal flora has been continuously recognized. Intestinal flora has become a key component not only in tumorigenesis but also in disease-free survival after surgery. Some scholars believe that it has a potential role in reducing postoperative inflammation and recurrence rate of CRC [9, 10]. However, there are few studies on the effect of intestinal flora on recurrence and prognosis of CRC. This study focuses on the combination of microbial sequencing and bioinformatics analysis technology to explore the changes of intestinal flora in patients with CRC before and after treatment and to analyze the impact and significance of this difference on recovery or recurrence after treatment.

2. Methods

2.1. Diagnosis and Treatment of Colorectal Cancer Patients. Patients diagnosed with CRC by digital anal examination assisted by rectal endoscopy [11] were included in the study. Some intestinal hyperplasia tissues were taken with biopsy forceps for pathological examination (including hematoxylin and eosin staining) to determine the deterioration of intestinal hyperplasia. CRC patients underwent surgical resection of tumor tissue and received chemical therapy (treated with tegafur, calcium folinate, and oxaliplatin for 5 days) one month later. Patients had no obvious discomfort during chemotherapy.

2.2. Collection and Treatment of Samples. From May 2019 to October 2019, a total of 17 fecal microbiological samples were collected from healthy people and patients with colon and rectal cancer, including 5 normal samples (control group). Among the intestinal microbiological samples of colorectal cancer (CRC group), there were 3 colon cancer (CC) samples and 2 rectal cancer (RC) samples. There were 3 cases of CRC samples after operation (CRC_O) and 4 cases of CRC samples after chemotherapy (CRC_C). The samples were collected with sterile fecal collection kit and sent to the sterile operation platform. Microbial DNA was extracted immediately with fecal DNA extraction kit and then stored at -20/-80°C. The extracted total DNA samples were transported to the laboratory with dry ice for subsequent amplification, library construction, and sequencing (Figure 1).

2.3. Sequencing and Data Processing of 16S rDNA of Intestinal Flora. The V3+V4 region of 16S rDNA was amplified by specific primers with barcode. The upstream primer was 5'-CCTACGGGNGGCWGGCAG-3' (341), and the downstream primer was 5'-GGACTACHVGGGTATCTAAT-3' (806). The size of the amplified product was about 466 bp. Quantifluor™ fluorometer (Promega, Madison, WI, USA) was used to quantify the DNA of samples. The purified amplified products were mixed in the same amount, connected to the sequencing connector, and the sequencing library was constructed. Illumina PE250 (San Diego, CA, USA) was used for sequencing. The raw reads obtained by sequencing were first removed by FASTP software [12], and then tags were spliced into raw reads with Fast Length Adjustment of Short Reads (FLASH) software [13]. UCHIME algorithm [14] of USEARCH software (http://www.drive5.com/) was used to filter chimera and other non-biological data, so as to enhance the statistical reliability and biological validity of the data.

2.4. OTU Clustering and Bacterial Species Annotation. The effective reads were clustered according to sequence similarity, and the tag sequences with 100% similarity were classified as an operational tax unit (OTU). The absolute and relative abundances of each OTU in each group were calculated. The common and unique OTUs of each group were analyzed and compared. The OTU sequences were compared by SILVA software (https://www.arb-silva.de/) [15], and the bacterial population was annotated according to 97% sequence similarity.

2.5. Alpha Diversity Analysis. The Observed_OTUs (Sobs, species richness), Chao1 (bacterial diversity), and Abundance-based Coverage Estimator (ACE, bacterial abundance) indexes were calculated. Simpson and Shannon comprehensively reflected the species richness and evenness.

2.6. Beta Diversity Analysis. The Jaccard index was mainly based on the presence or absence of species to evaluate the difference of flora structure. Bray_curtis was based on the presence or absence of species and species abundance. Unweighted_unifrac was based on the evolutionary distance of species. Weighted_unifrac was based on evolutionary distance and species abundance. Unweighted pair group method with arithmetic mean (UPGMA) cluster tree and principal coordinate analysis (PCoA) analysis were conducted based on the above four distance indexes to observe the difference degree of microbial community structure among samples.
The corresponding relationship of fid databases, the phenotypic characteristics of species were classified using QIIME (http://qiime.org/) [16], KEGG (https://www.kegg.jp/), the GreenGenes database (IMG, https://img.jgi.doe.gov/cgi-bin/m/main.cgi) [17], and PATRIC (https://www.patricbrc.org/) [18, 19] databases, the phenotypic characteristics of species were classified. According to the species annotation results of GreenGenes database (http://greengenes.secondgenome.com/) [20], the corresponding relationship of “species-gene-phenotype” was constructed, and the phenotype classification and abundance of samples were predicted by BugBase software (https://bugbase.cs.umn.edu/).

2.9. Functional Enrichment Analysis of Fecal Bacteria. IMG, PICRUSt2, SILVA, and other databases were used to construct the evolutionary tree. The reference species of OTU were selected according to the tree structure, and the “species-gene” relationship network was built by Functional Annotation of Prokaryotic Taxa (FAPROTAX, http://www.loucalab.com/archive/FAPROTAX) [21] and Tax4Fun (http://tax4fun.gobics.de/) [22] software. The count values of KO (Kyoto Encyclopedia of Genes and Genomes ontology) pathway in each sample were analyzed for direct comparison between groups.

2.10. Statistical Analysis. GraphPad Prism 7.0 software (San Diego, CA, USA) was used for data analysis. The Kruskal-Wallis (KW) rank sum test was used to analyze the difference of species richness among groups. Then, the Wilcoxon rank sum test and Welch’s t test were used to test the difference between the two groups. P < 0.05 showed that there was significant difference between the groups.

3. Results

3.1. Clinical Situation of Colorectal Cancer Patients. The gastrointestinal tumors were identified by endoscopy and pathological staining (Figure 2). CRC patients included in the study collected intestinal flora samples before and after surgical eradication and after chemotherapy. The disease and treatment groups (12 samples) of 17 intestinal microbiological samples were collected from 5 cases of different CRC patients, including 3 men, 2 women, 3 colon cancer, and 2 rectal cancer. The clinical information of CRC patients is shown in Table 1.

3.2. Intestinal Bacterial Diversity in Patients with Colorectal Cancer before and after Treatment. A total of 1475040 original reads (an average of 86767.06 reads/sample) were obtained following quality control. The average effective data ratio was 80.93% (Table S1 and Figure S1). According to the statistics of total OTUs, the total number of intestinal bacteria in CRC patients increased significantly compared with the control group. There was no significant difference between CRC with CRC_O and CRC_C groups (Figure S2A). 402 of OTUs were codistributed among the four groups (control, CRC, CRC_O, and CRC_C) were used for subsequent diversity studies (Figure S2B).

The results of alpha diversity analysis showed that the Chao1 index of CRC (P = 0.0079), CRC_O (P = 0.0159), CRC_C (P = 0.0357) had a higher level than that of the control group (Figure 3(a)), and the difference was statistically significant. There was no statistically significant change in Chao1 index among the CRC, CRC_O, and CRC_C groups (P > 0.05). The change trend of Observed OTUs (Figure S3A) and ACE (Figure S3B) indexes was similar to that of Chao1 index. There was no significant difference in the Simpson (Figure 3(b)) and Shannon (Figure S3C) indexes between all study groups (P > 0.05) (Figure 3(c)).
Figure 2: Endoscopic examination of colorectal cancer tissue. The lesion site of the colon (a) and rectum (c) under endoscope. Hematoxylin and eosin (H&E) staining results of colon (b) and rectal (d) cancer tissues under light microscope (100x).

Table 1: Statistics of clinical data of colorectal cancer patients.

| Patients                          | Mean ± SD       |
|----------------------------------|-----------------|
| Age (year)                       | 58.20 ± 12.09   |
| Gender (n = 5)                   |                 |
| Female                           | 2               |
| Male                             | 3               |
| Colorectal cancer                |                 |
| Colon cancer                     | 3               |
| Rectal cancer                    | 2               |
| Tumor size (cm$^3$)              | 28.23 ± 20.91   |
| Clinical stage                   |                 |
| I                                | 1               |
| II                               | 1               |
| III                              | 2               |
| IV                               | 1               |
| MMR (n = 5)                      | 1               |
| Ki-67 (+)                         | 62.00 ± 16.00%  |
| Chemotherapy methods             |                 |
| Tegafur+irinotecan               |                 |
| Tegafur+calcium folinate+oxaliplatin |         |
| Tegafur+irinotecan+calcium folinate+oxaliplatin | |
| Times of chemotherapy            | 4.80 ± 1.64     |
| Lymphatic metastasis             | 3               |

MMR indicates CRC patients with Lynch syndrome who carry a mismatch repair (MMR) gene mutation.
Based on the distance of species evolution (Unweighted_unifrac index), UPGMA cluster tree and PCoA results showed that the sample distance was consistent with the grouping (Figures 4(a) and 4(b)). The ANOSIM test showed that there were significant differences among all groups ($P = 0.001$) (Figure S4A). Compared with the control group, the OTU composition based on Unweighted_unifrac distance of CRC ($P = 0.008$), CRC_O ($P = 0.009$), and CRC_C ($P = 0.022$) was significantly higher (Figure 4(c) and Figure S4A). Based on the Jaccard distance, beta diversity of gut flora in CRC_O and CRC_C was significantly lower than that of the control group (Figures 4(c)–4(e)). There was no significant difference in the CRC, CRC_O, and CRC_C groups (Figure 4(d)). The change of beta diversity based on Weighted_unifrac distance in each group was not statistically significant, and the change trend of beta diversity based on the Bray distance was similar to that of the Jaccard distance (Figure 4(d)). Then, the beta diversity of flora at different levels was analyzed. It was found that only the beta diversity based on the Jaccard distance at genus level was significantly reduced between the control and CRC_O groups (Figure S4C). These results indicated that the diversity of intestinal microflora in CRC patients increased significantly compared with the normal population, and surgery and chemical therapy failed to alleviate this trend. Surgical treatment had the potential to change the diversity of intestinal bacterial composition.

Notes are appended as follows:
- Raw PE: Original pair-end reads logarithm,
- Clean PE: High-quality pair-end reads logarithm after quality control filtration,
- Raw Tags: Raw tags after overlap assembling,
- Clean Tags: High-quality tags after tags filter,
- Chimera: The number of chimeric tags detected in OTU clustering,
- Effective Tags: The number of high-quality tags after removing the chimera, which is the effective tags for subsequent analysis.

Effective Ratio (%): Percentage of high-quality tags in original PE reads.

CON_1-CON_5 belongs to the control group; CC_1-CC_3 are samples from colon cancer patients, belonging to the CRC group; CCT0_1 and CCT0_2 are samples from colon cancer patient after operation treatment, belonging to the CRC_O group; CCT1_1 and CCT1_2 are samples...
Figure 4: Beta diversity analysis of intestinal bacteria before and after treatment of colorectal cancer. Unweighted pair group method with arithmetic mean (UPGMA) cluster tree (a) and principle coordinate analysis (PCoA) plot based on Unweighted_unifrac distances show distinct bacterial community clusters between the control and CRC patients before and after treatments (b), PCoA plot shows distinct bacterial community clusters based on Jaccard (c), the heat map shows the ANOSIM test results of beta diversity indexes (Unweighted_unifrac, Weighted_unifrac, Bray, and Jaccard) among each group (d), and box diagram shows the OTU diversity based on the Jaccard distance of each group (e).
from colon cancer patient after chemotherapy, belonging to the CRC_C group; RC_1 and RC_2 are samples from rectal cancer patients, belonging to the CRC group; RCT0_1 is sample from rectal cancer patient after operation treatment, belonging to the CRC_O group; RCT1_1 and RCT1_2 are samples from rectal cancer patient after chemotherapy, belonging to the CRC group.

3.3. Differences of Bacterial Phylogenetic Spectrum and Taxa in Colorectal Cancer Patients before and after Treatments.

99%, 99.98%, 99.52%, 99.32%, and 92.47% of phylum, class, order, family, and genus were successfully annotated, but only 14.60% of species were annotated. Firmicutes, Proteobacteria, Actinobacteria, and Bacteroides constitute >90% of bacteria phyla in each group, showing basically consistent phylum relative abundance between groups (Figure 5(a)). The first three most abundant families in normal people were Lachnospiraceae, Ruminococcaceae, and Enterobacteriaceae. The most abundant family of CRC and CRC_O patients was Enterobacteriaceae and the second was Veillonellaceae (CRC) and Bifidobacteriaceae (CRC_O), followed by Lachnospiraceae (CRC) and Ruminococcaceae (CRC_O), respectively. Lachnospiraceae family is the most abundant in the CRC group, followed by Enterobacteriaceae family (Figure 5(b)). In genus level, the relative abundance of Bifidobacterium (13.87%) and Escherichia_Shigella (10.44%) in the control group was higher than other genera; the relative abundance of Escherichia_Shigella (31.53%) and Megamonas (14.15%) was higher in the CRC groups. In the CRC_C group, Escherichia_Shigella (15.86%) and Ruminococcus_gnavus (14.72%) were higher. The relative abundance of Escherichia_Shigella (39.49%) and Bifidobacterium (7.81%) was higher in the CRC_O group (Figure 5(c)).

LEFSE and IndVal were used to evaluate the differences of intestinal bacterial populations in CRC patients to determine significantly enriched bacterial taxa before and after treatment. Compared with the control group, the intestinal tract of CRC patients was enriched with Negativicutes class, Clostridia class, Selenomonadales order, Clostridiales order, Veillonellaceae family, and Megamonas genus (IndVal = 14.15) (P < 0.05, abundance > 10). And normal people was enriched with Clostridia class, Negativicutes class, Clostridiales order, Selenomonadales order, and Blautia genus (IndVal = 3.91) (P < 0.05, abundance > 2.5). Compared with the CRC group, the flora of CRC_C was enriched with Verrucomicrobia phylum, Verrucomicrobiae class, Verrucomicrobiales order, Akkermansia family, and Akkermansia genus (P < 0.05, abundance > 9). The bacterial classification of the CRC_C group was similar to CRC. Compared with normal people, the microbiota of colorectal cancer patients showed different bacterial populations at the taxonomic level of phylum, class, order, family, genus, and species. Chemical therapy led to significant changes in intestinal bacterial populations of CRC patients at the levels of phylum, class, order, and family, but there was no significant difference in bacterial populations after radical surgery (Table 2 and Figure S5).
3.4. Effect of Surgery and Chemotherapy Treatment on Intestinal Microflora Phenotype in Patients with Colorectal Cancer. BugBase analyzed the bacterial phenotype of each group, and the results showed that the control group was mainly enriched in Anaerobic and Gram_positive phenotypes, while the CRC group was mainly enriched in Anaerobic, Gram_positive, and Gram_negative phenotypes. The CRC_C group was similar to the control group, and species mainly enriched in Anaerobic and Gram_positive phenotypes. The CRC_O group was mainly enriched in the Gram_negative and contains mobile element phenotypes (Figure 6(a)). There was significant difference in Aerobic phenotype between groups (P = 0.031), but the difference only existed between the CRC and CRC_O groups. There was no significant difference in species phenotype between the CRC and control groups (Figure 6(b)). This indicated that the bacteria change of CRC_O group was related to Aerobic phenotype. The Aerobic germs in the CRC_O group were mainly distributed in the Proteobacteria phylum (Figure 6(c)). For the analysis of phenotypic function among groups, the increase of Aerobic phenotype led to the enhancement of respiration of sulfur compounds (Figure 6(d)).

3.5. Effect of Chemical Drugs and Surgical Treatment on Intestinal Flora Function in Patients with Colorectal Cancer. The change of bacterial composition affects its function in the intestine. Compared with normal people, intestinal bacteria in CRC patients significantly increased the biological pathway of lipopolysaccharide biosynthesis (P = 0.02915), phosphotransfer system (PTS) (P = 0.02222), D-arginine and D-ornithine metabolism (P = 0.03061), biotin metabolism (P = 0.01746), and so on (Figure 7(a)). After chemotherapy, the biological pathways of streptomycin biosynthesis (P = 0.02499) and polyketide sugar unit biosynthesis (P = 0.04511) of intestinal bacteria in CRC patients were significantly enhanced, and the ability of vitamin B6 metabolism (P = 0.02151) was weakened (Figure 7(b)). Compared with the intestinal bacterial function of CRC patients before treatment, the metabolism of xenobiology by cytochrome P450 (P = 0.00913) biological pathway was significantly enhanced after surgical eradication (Figure 7(c)).

| Table 2: The results of bacterial classification differences in each group based on LEFSE method (P < 0.05). |
|-------------------------------|-----------------|-----------------|-----------------|
| Levels                        | Control         | CRC             | CRC_C           | CRC_O           |
| Phylum                        | —               | Verrucomicrobia, Synergistetes | —               | —               |
|                              | Class           | Acidimicrobiota, Deltaproteobacteria, Synergistetes, and Verrucomicrobiota | —               | —               |
|                              | Order           | Propionibacteriales, Desulfovibrionales, Corynebacteriales, Synergistales, IMCC26256, and Verrucomicrobiota | —               | —               |
|                              | Family          | Chloroplast, Verrucomicrobiales | —               | —               |
|                              | Genus           | Akkermansiaceae, Aerococcaceae, Synergistaceae, Cyanobacteria, and Desulfovibrionaceae | —               | —               |
|                              | Species         | Eubacterium, ruminantium group | —               | —               |
|                              |                 | Akkermansia, Clostridia, 1 | —               | —               |
|                              |                 | Clostridium, perfringens, CPE_str, F4969, Bifidobacterium, animalis, Veillonella, magna | —               | —               |
|                              |                 | GCA_900066225, Pyramidobacter, Hungatella, Eubacterium, sp, Marseille, P3202, Bacteroides, vulgatus, Parabacteroides, distasonis, Hungatella, hathewayi, Clostridium, symbiosum, ATCC_14940, Erysipelatochezridium, ramosum, Akkermansia, muciniphila, Emergencia, timonensis, and Corynebacterium, afermentans, subsb_afermentans | —               | —               |
|                              |                 | Ruminococcus, bicirulans, Alistipes, inops, Coprococcus, sp, DJF_8005, and Lachnospiraceae, bacterium, TF01_11 | —               | —               |
3.6. Changes and Functions of Key Intestinal Bacteria in Colorectal Cancer Patients before and after Treatments. The marker intestinal bacteria of each group were identified based on the genus level. *Megamonas* (IndVal = 14.15) was the indicator and dominant genus of CRC patients, and its abundance was 103.49 times of that of the control group, while *Ruminocaceae_UCG-014* (IndVal = 3.85) was the dominant germ in the control group (Figure 8(a)). Both belong to *Firmicutes* phylum. The Pearson correlation between *Megamonas* and *Ruminocaceae_UCG-014* was weak ($P > 0.05$). *Prevotellaceae_UCG-001* genus in the CRC_O group was significantly higher ($P = 0.045$, fold = 97.58) than the CRC group (Figure 8(b)). Compared with CRC group, the most significant genera ($P < 0.05$) in the CRC_C group were *Prevotellaceae_UCG-001* (IndVal = 0.062) and *Tyzzerella_4* (IndVal = 0.027) of Pasteurellaceae family (Figure 8(c)).
There was positive correlation between *Megamonas* with *Megasphaera* \((r = 0.625711, P = 0.007218)\) and *Ruminococcus_torques_group* \((r = 0.533661, P = 0.027364)\) (Figure 9(a)). *Ruminocaceae_UCG-014* was mainly associated with *Lachnospiraceae nk4a136 group*, *Lachnospiraceae nd3007 group*, and *Ruminococcus_1* \((r > 0.9, P < 0.001)\) (Figure 9(b)). They were mainly involved in metabolism, cellular processes, cancer, and other pathways. The KO pathway analysis showed that the intestinal microbial function of CRC patients was mainly involved in the biosynthesis and metabolism of glycans and the metabolism and degradation of terpenoids, polyketides, and heterobiotics. The biosynthesis of lipopolysaccharide, the biosynthesis of type II polyketide skeleton, and the degradation of nitrotoluene in CRC patients were significantly higher than those of normal people \((\text{fold} > 2, P < 0.05)\) (Figure 9(c)). *Tyzzerella_4* was positively correlated with *Phascolarctobacterium* \((r = 0.76, P = 0.00041)\), *Luminococcus gnarus group* \((r = 0.74, P = 0.00065)\), and *Bacteroides* \((r = 0.52, P = 0.031)\) in postoperative CRC patients (Figure 10(a)). These microfloras were mainly involved in adipocytokine signaling pathway, peroxisome, bile secretion, and fatty acid biosynthesis \((\text{fold} > 1, P < 0.05)\) (Figure 10(b)). *Akkermansia* was positively correlated with six bacterial genus \((P < 0.0001)\), including *Eubacterium_ruminantium group* \((r = 0.99)\), *Eubacterium_nodatum group* \((r = 0.97)\), and *Uba1819* \((r = 0.85)\) in CRC patients after chemotherapy (Figure 10(c)). Analysis of these microbiota-related KO pathways in the CRC_C group showed that the rich genera mainly enhanced protein digestion and absorption in immune, digestive, and endocrine systems, renin-angiotensin system pathway, secondary bile acid biosynthesis, and adipocytokine signaling pathway \((\text{fold} > 2, P < 0.05)\)
After chemotherapy and surgical treatment, the adipocytokine signaling pathway, peroxisome, and bill secret KO pathway were significantly enhanced in CRC patients. Chemotherapy also significantly weakened the renin-angiotensin system, primary bile acid biosynthesis, nitrotoluene degradation, flagella assembly, etc. in the CRC group.

4. Discussion

There are more than 1000 kinds of bacteria in human intestinal tract, and the number is as high as 10 trillion. These bacteria form a barrier to protect the intestine and participate in the absorption and metabolism of nutrients. Bad dietary behavior affects the composition of intestinal flora, will disturb the normal metabolism of the body, and then affects the susceptibility of intestinal diseases, thus inducing inflammation, cancer, and other diseases. It is generally believed that the mechanism of action of intestinal flora on colorectal cancer is related to the increase of toxins produced by bacteria, the decrease of beneficial bacterial-derived metabolites, the destruction of epithelial barrier, the production of cancer-promoting compounds, and the change of intestinal flora or intestinal flora. Some studies have detected that species abundance is significantly downregulated in the intestine of CRC patients. However, in this study, the intestinal flora abundance and diversity of CRC patients are increased. In addition, there is no change in the composition of species structure. There seems to be a contradiction with the current research. This is proposed that the loss of specific
beneficial bacteria or the serious loss of diversity among beneficial bacteria is the key to the occurrence of colorectal cancer.

In this study, the relative abundance differences of species composition in the colonies is further analyzed, and the significantly decreased abundance of Ruminococcaceae_UCG-014, Ruminococcus_1, Eubacterium_eligens, and Coprococcus_2 in colorectal cancer patients are found. Especially, Rucocamace_UCG-014, which belongs to the Ruminocaceae family, is related to fat metabolism. Rucocamace_UCG-014 stimulates lactic acid metabolism to produce short-chain fatty acids and plays anti-inflammatory and anticancer roles. It is a kind of beneficial anaerobic bacteria [23–26]. The lack of Ruminocaceae leads to high intestinal inflammation and the high risk of cancer. The pathogenic bacteria Allisonella and Megamonas are significantly increased in CRC, which has the potential to trigger intestinal inflammation [27]. Megamonas belongs to Clostridium family in Firmicutes phylum [28], which is a kind of anaerobic bacteria. Especially Megamonas is positively correlated with lactic acid content and negatively correlated with acetate [29]. Some studies believe that Megamonas is the characteristic flora of lymph node metastasis of colorectal cancer and one of the main floras leading to abnormal metabolism of CRC. A medical research led by Han et al. [30] shows that Megamonas genus is a characteristic microorganism of the intestinal tract in colorectal cancer patients with lymph node metastasis, which is significantly associated with poor prognosis caused by lymph node metastasis. Through the analysis of KO functional pathway, it is found that the intestinal microflora of colorectal cancer is mainly related to lipopolysaccharide biosynthesis, biosynthesis of type II polysaccharide backbone, nitrotoluene degradation, etc. In addition, the function of renin-angiotensin system and primary bile acid biosynthesis is significantly reduced. Megamonas mediates the biosynthesis of lipopolysaccharide (LPS). High levels of lipopolysaccharide are detected in patients with irritable bowel syndrome and inflammatory bowel disease. The higher level of LPS is also increased of colorectal cancer incidence rate and the proliferation of tumor cells [31]. Therefore, the increased abundance of Megamonas leads to high level of LPS, which is a risk factor for cancer.

In addition, this study focused on the diversity and structural composition of intestinal flora in CRC patients before treatment, after surgical eradication, and after chemotherapy. This observation of intestinal colonies is useful for the identification of therapeutic effectiveness and recurrence. For example, the study of Jin et al. [9] detected that the intestinal microflora of 116 CRC patients before and after operation shows that there are significant differences between Gemella, Tyzzerella 3, Unclassified Oxalobacteraceae, Howard Ella,
Figure 10: The predominant intestinal flora interactions (a, c) and their functional differences (b, d) in colorectal cancer patients after surgery and chemical drug therapy.
Lawsonella, and Parascardovia. In addition, high levels of Fusobacterium nucleatum (F. nucleatum) and Bacteroides fragilis (B. fragilis) significantly shortened the relapse-free survival. This is because these bacteria cause the decrease of butyrate metabolism, promote the proliferation of tumor cells, and ultimately increase the overall incidence rate and tumor survival rate [32]. F. nucleatum also activates autophagy to chemoresistance of colorectal cancer [33]. Therefore, intestinal flora of CRC patients after surgery and drug therapy improves lipid metabolism to a certain extent and thus participates in the prognosis of cancer patients.

The results in this study showed that there was no significant change in species diversity in the three stages (CRC, CRC_O, and CRC_C groups). There was no significant change in intestinal colony structure before and after surgical eradication, while chemotherapy affected the relative abundance of Verrucomicrobia phylum and subordinate microbiota. In fact, surgery and chemical drug treatment do not regulate the abundance of intestinal flora in CRC patients including Meganonas and Ruminocaceae_UCG-0014. Relative abundance of Prevotellaceae_UCG-001 in CRC patients is significantly increased after surgery, which is related to nutritional metabolism. Prevotellaceae_UCG-001 promotes the metabolism of cholesterol and bile acids [34] and significantly reduces [35] in colitis animal models. Upregulation of Prevotellaceae_UCG-001 promotes the production of short-chain fatty acids, improves intestinal microbial composition, regulates the immune regulatory activity related to intestinal flora, and maintains intestinal homeostasis [36]. Akkermansia genus is a kind of Gram-negative bacteria and a promising probiotic candidate [37, 38]. The results of a study in mice with colitis showed that the probiotics Akkermansia inhibits the occurrence of cancer [39]. Akkermansia is an effective colony marker for colon cancer treatment.

There are 5 CRC patients in this study, and the biological sample size is small, which needs to be further improved. In addition, the correlation between human genome and intestinal flora is ignored and does not fully explain the molecular mechanism of intestinal flora regulating colorectal cancer. This also needs to be improved in the future.

5. Conclusion

In this study, the abundance and diversity of intestinal flora in colorectal cancer patients are significantly higher than that of normal man. The abundance of Prevotellaceae_UCG-001 is significantly decreased and Meganonas is significantly increased in CRC patients, which are involved in the biosynthesis of lipopolysaccharide and have a potential role in promoting colorectal cancer. However, chemical drug and surgical treatment do not alleviate the rise trend of intestinal flora abundance but reduce Tyzzerella_4 and increase the proportion of Akkermansia to regulate intestinal immunity and metabolism.

Abbreviations

CRC: Colorectal cancer
KEGG: Kyoto Encyclopedia of Genes and Genomes
KO: Kyoto Encyclopedia of Genes and Genomes Ontology
OTU: Operational tax unit
UPGMA: Unweighted pair group method with arithmetic mean
PCoA: Principal coordinate analysis
NMDS: Nonmetric multidimensional scaling
IndVal: Indicator values
IMG: Integrated Microbial Genomes
FAPROTAX: Functional Annotation of Prokaryotic Taxa
KW: Kruskal-Wallis rank sum test
CON: Control
CC: Colon cancer
CCT: Colon cancer patients after treatment
RC: Rectal cancer
RCT: Rectal cancer patients after treatment.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Ethical Approval

The study was approved by the Ethics Association of the Guangdong Second Provincial General Hospital.

Consent

The patients were informed and approved the publication of the manuscript.

Conflicts of Interest

There is no conflict of interest for all authors.

Authors’ Contributions

All authors approved the publication of the manuscript.

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Supplementary Materials

Supplementary 1. Table S1: pretreatment and quality control of sequencing data.

Supplementary 2. Figure S1: effective reading percentage (A) and number (B) of 16S sequencing of microorganisms in each sample. Reads QC filter, low quality reads; nonoverlap, reads that cannot be spliced into tags; tag QC filter, tags that do not pass “tag filtering”; and effective tags, effective tags for subsequent analysis.

Supplementary 3. Figure S2: histogram of changes in total OTUs (A) and Venn diagram of total OTU distribution (B) of intestinal flora in patients with colorectal cancer before and after treatments. ** indicates $P < 0.01$ via comparing with the control group.
**Supplementary 4.** Figure S3: alpha diversity analysis dilution curve. (A–C) Observed OTUs, ACE, and Shannon indexes. OTUs: operational tax units. ACE, Abundance-based Coverage Estimator, indicates bacterial abundance.

**Supplementary 5.** Figure S4: box diagram of the ANOSIM test results of flora beta diversity based on the unweighted distance (A) and Jaccard distance (B) in CRC patients before and after treatments. And the heat map (C) shows the ANOSIM test results of flora beta diversity based on the Jaccard distance at different levels (phylum, class, order, family, genus, and species).

**Supplementary 6.** Figure S5: the distribution of bacterial clusters in CRC patients before and after treatment. The bar chart shows the bacterial structure composition at the (A) phylum, (B) order, (C) class, (D) family, and (D) genus levels with significant differences among the groups tested by Kruskal-Wallis (KW) method ($P < 0.05$).

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