Study on the Effect of Huanglian Jiedu Decoction on the Composition of Gut Microflora in SD Rats Based on 16S rRNA Sequencing

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Objective. To study the effects of Huanglian Jiedu Decoction (HJD) on gut microflora of SD rats. Method. After 3 days of adaptive feeding, 36 rats were randomly divided into 4 groups, namely, the normal control group (NC, 10 mL/kg, distilled water), high_HJD dose group (H_HJD, 6.25 g/kg, weight ratio between crude drug and rat), medium_HJD dose group (M_HJD, 3.125 g/kg), and low_HJD dose group (L_HJD, 1.56 g/kg), and each group consisted of 9 mice. The HJD groups were then treated with orally administered HJD for 21 days, while the NC group was treated with distilled water. After 7 days, 14 days, and 21 days of the experiment and after 12 hours of fasting and water deprivation, 3 SD rats in each group were randomly sacrificed by cervical dislocation and sterile operation to collect stool faces. Sample DNA was extracted by Fecal total DNA extraction kit, sequenced using Illumina MiSeq platform, and analyzed. Results. The abundance of Romboutsia, Turicibacter, Lactobacillus, and Gemella decreased, while that of Escherichia_Shiella, Caprobacillus, Blautia, Akkermansia, Klebsiella, Rhodococcus, Parabacteroides, Citrobacter, Enterococcus, Bacteroides, and Erysipelotrichaceae_incertae_sedis increased after using H_HJD. The abundance of Gemella, Turicibacter, Romboutsia, and Lactobacillus decreased, while that of Blautia, Akkermansia, Escherichia_Shiella, Thioecoccus, Rothia, Enterococcus, Lactobacillus, and Erysipelotrichaceae-incertae-sedis increased after using M_HJD. The abundance of Romboutsia, Turicibacter, Lactobacillus, and Gemella decreased, while that of Blautia, Akkermansia, Escherichia_Shiella, Bacteroides, Akkermansia, Enterococcus, Rhodococcus, Parabacteroides, Desulfovibrio, Blautia, Fusobacterium, Rothia, and Streptococcus increased after using L_HJD. Conclusion. HJD can regulate gut microbiota, and its effect varies with different dosage and duration of medication.

1. Introduction

As the normal microbiota in the intestine, gut microbiota not only plays an important role in the physiological functions of the intestine but also plays an important role in the host’s physiology, metabolism, immunity, digestion, and nutrition uptake [1]. And the composition of gut microbiota is altered in many disease states, including type 2 diabetes mellitus (T2DM), high blood pressure (BP), colitis, cardiovascular disease, and obesity [2]. Dysbiosis in the gut microbiota of type 2 diabetes mellitus (T2DM) patients is characterized by a higher proportion of Bacteroides [3]. Li and his colleagues found that the gut microbiota in patients with hypertension had a higher percentage of an enterotype rich in bacteria from the genus Prevotella [4]. In the gut microbiota of colitis patients, there was imbalance in Bacteroidetes, and species diversity decreased [5]. The stable balance between host and gut microflora plays an indispensable role in maintaining human health and has become a research object of widespread concern. Huanglian Jiedu Decoction (HJD), first recorded in the book “Wai-Tai-Mi-Yao” in the Tang dynasty, is a representative agent for clearing away heat and detoxification in traditional Chinese medicine [6]. It has been clinically practiced in the treatment...
of T2DM, cerebrovascular disease, hypertension, cancer, and other diseases in China. [7–11]. However, the mechanism of HJD for the treatment of diseases is currently lacking. Exploring the role and mechanism of traditional Chinese medicine through the gut microflora has become a research hotspot in the treatment of disease [12]. Therefore, 16S rRNA sequencing technology was used to analyze the effect of HJD on the structure of gut microflora in SD rats and to clarify the effect of HJD on gut microflora and provide reference for clinical application.

2. Methods

2.1. Preparation of HJD. The traditional Chinese medicine (TCM) formula in our study was HJD, composed of four herbs, namely, *Rhizoma Coptidis*, *Radix Scutellariae*, *Phellodendron amurense* Rupr, and *Gardenia jasminoides* Ellis. Herbs were purchased from Beijing Tongrentang Guiyang Branch. We prepared HJD referred to the optimized method described by Guang Chen [13]. Details are as follows: distilled water was used for the preparation of samples. *Rhizoma Coptidis* (9 g), *Radix Scutellariae* (6 g), *Phellodendron amurense* Rupr (6 g), and *Gardenia jasminoides* Ellis (9 g) were crushed into small pieces with scissors and mixed. The mixture was soaked in 1000 mL of cold water for 40 min before being boiled for 1.5 h. Filtrates were collected, and the residues were then boiled with an addition of 300 mL water for 40 min. Finally, the two filtrates were combined and concentrated to a final volume of 48 mL (0.625 g/mL) by rotary evaporation instrument (model: RE5203, Shanghai Yarong biochemical instrument factory). The filtrate was stored in seal at 4°C.

2.2. Animal Experiments

2.2.1. Animals. Specific-pathogen-free (SPF) Sprague Dawley rats were purchased from Chongqing Tengxin Biotechnology Co., Ltd. (age, 20 ± 2 d, weight, 200 ± 20 g, 50% female; n, 36). Rats were acclimatized to the laboratory for 3 days prior to the experiments.

2.2.2. Experimental Processes. The animal experiments were conducted by referencing the methods of a similar study with minor modifications [14]. We randomly divided the 36 rats into 4 groups, namely, the normal control group (NC, 10 mL/kg, distilled water), high_HJD dose group (H_HJD, 6.25 g/kg, weight ratio between crude drug and rat), medium_HJD dose group (M_HJD, 3.125 g/kg), low_HJD dose group (L_HJD, 1.56 g/kg), and each group consisted of 9 mice. The HJD groups were then treated with orally administered HJD for 21 days, while the NC group was treated with distilled water. After 7 days, 14 days, and 21 days of the experiment, after 12 hours of fasting and water deprivation, 3 SD rats in each group were randomly sacrificed by cervical dislocation for aseptic operation to collect faeces. Faeces were frozen immediately with liquid nitrogen after sampling and stored at −80°C (model: DW-86L490J, Qingdao Haier biomedical Co., Ltd.) until the DNA extraction.

2.3. Analysis of Fecal 16S rRNA. The composition of gut microbiota was detected by 16S rRNA sequencing analysis. Microbial genomic DNA was extracted from fecal samples using a fecal total DNA extraction kit (batch number: DP328-2, Tiangen Beijing Biochemical Technology Co., Ltd.) according to the manufacturer’s protocols. The extracted DNA from each fecal sample was amplified at the V4 region of the 16S rRNA genes by using specific primers: 338F (5’-ACTCCTACGGGAGGCAGCAG-3’) and 806R (5’-GGACTACHVGGGTWTCTAAT-3’). The qualified amplicons were pooled into equimolar fractions and paired-end sequenced (PE250) on an Illumina MiSeq platform (Shanghai Biotree Biotech Co., Ltd, Shanghai, China).

2.4. Bioinformatics Analysis. Reads were assembled and analyzed using Vsearch (version 2.14.1) and USEARCH10 [15, 16]. After quality filtering and chimera removal, clean sequences and exact amplicon sequence variants (ASVs) were resolved using Usearch3. Taxonomy was assigned to ASVs using USEARCH10 -sintax with -strand_both and -sintax_cutoff 0.6. ASV abundance was normalized using a standard of sequence number corresponding to the sample with the least sequences. Subsequent analysis of alpha-diversity and beta-diversity were all performed basing on these output normalized data. Linear discriminant analysis effect size (LEfSe) was performed to find biomarker(s) differentially represented between NC and HJD groups [17]. PICRUSt2 (Phylogenetic Investigation of Communities by Reconstruction of Unobserved States 2) is used to impute MetaCyc pathway abundances from the original microbial abundance data [18]. And NC group data were used in our previous paper [19].

3. Results

3.1. Bacterial Alpha- and Beta-Diversity. Rarefaction analysis revealed that our population captured most gut microbiota members from each SD rat. Measurement of within-sample diversity (α-diversity) revealed a significant difference between HJD and NC varieties (Figures 1(b) and 1(c)). The gut microbiota of L_HJD_14d had higher abundance than NC (Figure 1(b)), indicating that taking low-dose HJD for 14d can increase species richness. The gut microbiota of L_HJD_7d, M_HJD_21d had higher diversity than NC (Figure 1(c)), indicating that taking low-dose HJD for 7d and L_HJD_7d can increase species diversity. It is interesting that the species diversity of L_HJD_14d was lower than that of NC. Unconstrained principal coordinate analysis (PCoA) of Bray–Curtis distance revealed that the gut microbiota of different dose and time of HJD and NC formed distinct clusters, in which the separation was not well along the first coordinate axis. The constrained principal coordinate analysis with Bray–Curtis distance showing that the contribution rate of current conditions to the total difference among samples was 43.7.2%, and there was significant difference among groups (P < 0.01).
3.2. Gut Microflora Structure Analysis. Next, we further analyzed the relative abundances of differential bacteria in the gut microbiota at phylum and genus levels. At the phylum level, phylum Firmicutes, Proteobacteria, and Bacteroidetes were most abundantly presented which accounted for the majority of the population in all groups (Figure 1(f)), and at the genus level, Lactobacillus and Romboutsia were most abundantly presented which accounted for the majority of the population in most groups (Figure 1(g)). The abundance of Firmicutes, Proteobacteria, and Bacteroidetes changes in different groups over time (Figure 1(h)). We can find that no matter which
bacteria, the final abundance of the M_HJD tends to be normal.

To identify specific bacterial taxa associated with HJD, we compared gut microbiota using LEfSe, and the criteria were set at a linear discriminant analysis (LDA) > 2. In the first week, at the phylum level, compared with NC, the abundance of Firmicutes was significantly reduced in H_HJD_7d and the abundance of Verrucomicrobia and Bacteroidetes was significantly increased; at the genus level, the abundance of Romboutsia, Turicibacter, and Clostridium sensu stricto was significantly reduced and the abundance of Escherichia-Shigella, Coprobacillus, Enterococcus, Blautia, Bacteroides, and Akkermansia increased significantly in H_HJD_7d (Figure S1).

Compared with NC, at the phylum level, Proteobacteria significantly decreased and Verrucomicrobia increased significantly in M_HJD_7d; at the genus level, the abundances of Gemella, Clostridium sensu stricto, Turicibacter, and Romboutsia decreased significantly and the abundances of Blautia, Akkermansia, Escherichia-Shigella, and Thio- bacillus increased significantly in M_HJD_7d (Figure S2).

Compared with NC, L_HJD_7d has no significant difference at the phylum level; at the genus level, the abundances of Romboutsia, Turicibacter, and Clostridium sensu stricto were significantly reduced and the abundances of Escherichia-Shigella, Bacteroides, Akkermansia, and Enterococcus were significantly increased in L_HJD_7d (Figure S3).

In the second week, at the phylum level, compared with NC, the abundance of Firmicutes was significantly reduced, and the abundances of Fusobacteria, Verrucomicrobia, and Bacteroidetes were significantly increased in H_HJD_14d; at the genus level, the abundances of Lactobacillus, Clostridium sensu stricto, and Turicibacter were significantly decreased and the abundances of Coprobacillus, Escherichia-Shigella, Klebsiella, Bacteroides, Fusobacterium, Enterococcus, Blautia, and Akkermansia increased significantly in H_HJD_14d (Figure S4).

Compared with NC, at the phylum level, the abundance of Firmicutes was significantly reduced; in L_HJD_14d; at the genus level, the abundances of Rothia, Gemella, and Lactobacillus were significantly reduced and Akkermansia, Bacteroides, Blautia, Escherichia-Shigella, Clostridium, XIVA, Erysipelotrichaceae incertae sedis, Enterococcus, and Clostridium XVIII increased significantly in M_HJD_14d (Figure S5).

Compared with NC, at the phylum level, the abundance of Firmicutes was significantly reduced and the abundances of Verrucomicrobia and Bacteroidetes were significantly increased in L_HJD_14d; at the genus level, the abundances of Lactobacillus, Romboutsia, Rhodococcus, and Gemella were significantly reduced and the abundances of Prevotella, Clostridium_XIVA, Akkermansia, Parabacteroides, Escherichia-Shigella, Eubacterium, Bacteroides, Desulfovibrio, and Blautia increased significantly in L_HJD_14d (Figure S6).

In the third week, at the phylum level, compared with NC, the abundance of Firmicutes was significantly reduced and the abundances of Verrucomicrobia, Proteobacteria, and Bacteroidetes were significantly increased in H_HJD_21d; at the genus level, the abundances of Romboutsia, Turicibacter, Clostridium sensu stricto, and Gemella were significantly reduced and the abundance of Clostridium_XVIII, Erysipelotrichaceae incertae sedis, Parabacteroides, Citrobacter, Enterococcus, Coprococcius, Eubacterium, Escherichia, Shigella, Akkermansia, Blautia, Klebsiella, and Bacteroides increased significantly in H_HJD_21d (Figure S7).

Compared with NC, M_HJD_21d has no significant difference at the phylum level; at the genus level, the abundance of Gemella is significantly reduced and the abundance of Enterococcus is significantly increased in M_HJD_21d (Figure S8).

At the phylum level, the abundance of Firmicutes decreased significantly and the abundances of Fusobacteria, Bacteroidetes, and Actinobacteria increased significantly in L_HJD_21d; at the genus level, the abundances of Lactobacillus, Romboutsia, and Turicibacter decreased significantly and the abundances of Blautia, Fusobacterium, Rhodococcus, Escherichia-Shigella Enterococcus, Bacteroides, Clostridium XIVA, Rothia, and Streptococcus increased significantly in L_HJD_21d (Figure S9).

3.3. Metabolic Pathway Predictions. To determine whether the observed taxonomic shifts in gut microbiota might alter their functional capabilities, we used PICRUSt2 to infer genomic content combined with STAMP to determine whether functions were differentially abundant. A total of 8 MetaCyc pathways were generated using the composition of the gut microbiota based on PICRUSt2 in M_HJD versus NC (Figure 2). PWY-6147 (6-hydroxymethyl-dihydropterin diphosphate biosynthesis I) and PWY-7539 (6-hydroxymethyl-dihydropterin diphosphate biosynthesis III) were increased in M_HJD_7d (Figure 2(a)). PENTOSE-P-PWY (pentose phosphate pathway) was increased in M_HJD_14d (Figure 2(b)). SER-GLYSYN-PWY (superpathway of L-serine and glycine biosynthesis I) decreased in M_HJD_21d (Figure 2(c)).

4. Discussion

TCM treatment of diseases emphasizes “the coordination of the five zang-viscera and the unity of man and nature,” and its views are highly similar to the system theory and balance theory of microecology [20]. Syndromes have the same treatment and different treatments are different in treatment based on syndrome differentiation. For this reason, the same prescription can treat different diseases. In recent years, the mechanism of action of TCM through the gut microbiota has been widely recognized with the continuous in-depth research on the gut microbiota.

The work explored the effect of HJD on gut microbiota from different doses, the results show that the abundance of Romboutsia, Turicibacter, Lactobacillus, and Gemella decreased, while that of Escherichia, Shigella, Coprococcius, Blautia, Akkermansia, Klebsiella, Rhodococcus,
Parabacteroides, Citrobacter, Enterococcus, Bacteroides, and Erysipelotrichaceae_incertae_sedis increased after using H_HJD. The abundance of Gemella, Turicibacter, Romboutsia, and Lactobacillus decreased, while that of Eshibacter_Shigella Blautia, Akkermansia, Thiobacillus, Rothia, Enterococcus, Lactobacillus, and Erysipelotrichaceae_incertae_sedis increased after using M_HJD. The abundance of Romboutsia, Turicibacter, Lactobacillus, and Gemella decreased, while that of Eshibacter_Shigella Blautia, Bacteroides, Akkermansia, Enterococcus, Rhodococcus, Parabacteroides, Desulfovibrio, Blautia, Fusobacterium, Rothia, and Streptococcus increased after using L_HJD.

Although different doses of HJD have different effects on gut microbiota of SD rats, there are still some of the same effects on Romboutsia, Turicibacter, Escherichia coli-Shigella, Braute, Akkermansia, Lactobacillus, Gemella, and Enterococcus. Studies have confirmed that the abundance of Akkermansia and Braute in diabetic patients has significantly decreased, while the abundance of Lactobacillus has increased [21]. This may be one of the mechanisms of HJD in treating diabetes.

Academician Wang Yongyan proved through experiments that HJD has a good preventive effect on Alzheimer’s disease [22]. Ketogenic diet can effectively reduce the risk of Alzheimer’s disease by increasing the abundance of Akkermansia and Lactobacillus and reducing the abundance of Turicibacter [23]. In this study, different doses can positively regulate Akkermansia, Braute, and Lactobacillus, suggesting that HJD may be effective in preventing diabetes and Alzheimer’s disease by acting on this type of bacteria.

HJD also has a certain preventive effect on colorectal cancer [24–26]. Metagenomic analysis shows that Gemella is significantly increased in the gut microbiota of patients with colorectal cancer [27]. Qin Huanlong’s research team showed that the abundance of Escherichia-Shigella in the intestine of sporadic colorectal cancer was significantly decreased [28]. The results of this experiment show that after the intervention of HJD, Gemella was significantly decreased, and Escherichia-Shigella was significantly increased, which further suggested the application value of HJD in the prevention and treatment of colorectal cancer.
4.1. Analysis from the Specificity of Each Dose Group. *Coprobacillus*, *Klebsiella*, and *Citrobacter* were only significantly affected in the H_HJD. *Thiobacillus* was only significantly affected in the M_HJD. *Streptococcus*, *Fusobacterium*, and *Desulfovibrio* were only significantly affected in the L_HJD.

Studies have shown that the abundance of *Coprobacillus*, in patients with acne, is significantly reduced, and it has been reported that HJD can treat acne [29, 30]. The abundance of *Coprobacillus* increased in H-HJD that explained part of the mechanism of HJD in the treatment of acne from the gut microbiota.

Bad breath is generally produced by heat evil in the stomach and intestines, and its main odor comes from sulfide [31]. *Thiobacillus* can oxidize sulfide (H$_2$S) to sulfate [32]. The increase of *Thiobacillus* in the middle-dose group suggests that *Thiobacillus* may be one of the targets of HJD in the treatment of bad breath.

4.2. Analysis from the Duration of Medication. For *Akkermansia*, the significant differences in the M_HJD_21d and L_HJD_21d groups disappeared, suggesting that H_HJD is needed to improve the abundance of *Akkermansia* for a long time.

For *Blautia*, it only began to increase significantly at 14 days in the L_HJD group, suggesting that *Blautia* needs a cumulative effect.

However, in the 7-day period of *Lactobacillus*, there is no significant difference in each dose group, suggesting that HJD has little correlation with its effect and dose but is related to the duration of medication.

Studies have shown that *Erysipelotrichaceae_iae* is a beneficial bacteria, and its abundance is negatively correlated with the blood sugar level, insulin level, and fat mass of female offspring [33]. The increase in the abundance of the bacteria in the H_HJD_14d and H_HJD_21d groups in this experiment further shows that HJD has a therapeutic effect on diabetes, but it requires 14 days of medication.

5. Conclusions

To sum up, this study comprehensively studied the effect of HJD on gut microbiota from different doses and different administration time. At the same time, it expands some potential target bacteria of HJD in the prevention and treatment of diabetes, Alzheimer’s disease, acne, bad breath, and colorectal cancer, but its causality needs to be further studied.

Data Availability

The sequence data used to support the findings of this study have been deposited to the NCBI Sequence Read Archive under the BioProject accession number PRJNA664442.

Disclosure

H. Du and G. Zhang are joint first authors.

Conflicts of Interest

The authors declare that there are no conflicts of interest associated with the manuscript.

Authors’ Contributions

G. He obtained funding and supervised the work. G. He, G. Yang, and W. Tian designed the study. P. Wang, W. Wang, and G. Zhang collected the data. H. Du analyzed the data and drafted the manuscript. All authors read and approved the final version of the manuscript prior to submission. H. Du and G. Zhang contributed equally to this work.

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Supplementary Materials

Figure S1: lefse analysis result figure of HJD and NC group. (Supplementary Materials)

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