Changes in Rats’ Gut Microbiota Composition Caused by Induced Chronic Myocardial Infarction Lead to Depression-Like Behavior

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Depression is common among patients who have chronic myocardial infarction (CMI). Despite their frequency, depression and CMI are bidirectional related conditions, each is a risk for the other, and they often co-exist, suggesting shared or interacting pathomechanisms. Accumulating data revealed the effects of gut microbiota in terms of regulating depression via the gut–brain axis. Thus, we investigated the role of gut microbial dysbiosis in CMI-induced depression-like behavior. Hierarchical cluster analysis of sucrose preference test (SPT) results was adopted to classify the CMI rats into depression-like behavior (CMI + Dep) or non-depression-like behavior (CMI + Non-Dep) phenotypes. First, 16S ribosomal RNA sequencing analysis showed both β-diversity and relative abundance of several gut bacteria significantly differed between the CMI + Dep and CMI + Non-Dep rats. Next, transplantation of fecal microbiota from CMI + Dep rats visibly altered the relative abundance of gut microbiota and also induced depression-like behavior in the antibiotics-treated pseudo-germ-free rats. In conclusion, these findings suggested that dysbiosis in gut microbial composition contributed to the onset of CMI-induced depression-like behavior and that exogenous regulation of gut microbiota composition could be a potential therapeutic strategy for CMI and related depression-like behavior.

Keywords: chronic myocardial infarction, gut microbiota, depression-like behavior, microbiome-gut-brain axis, hierarchical cluster analysis

INTRODUCTION

With the optimization of evidence-based treatments in modern society, the mortality of myocardial infarction significantly decreased (Saar et al., 2015; Gabet et al., 2016). However, a pile of literature revealed that myocardial infarction survivors were more prone to psychological disorders, such as depression (van Dijk et al., 2016). Over the past 20 years, it has been found that depressed chronic myocardial infarction (CMI) patients have a high risk of cardiac independent risk factors (Brborović et al., 2009) and poor prognosis of further adverse cardiac events
controlled relative humidity (60±5%). Experimental protocols and animal handling procedures were allowed to acclimate for 7 days before experiments. All animals were housed under controlled temperature (22±1°C), ad libitum access to food and water. Animals were weighed to ensure consistent body weight. Prior to modeling, animals were fasted for 12 h and deprived of water for 4 h in order to prepare them for the surgery.

MATERIALS AND METHODS

Animals

One hundred male SD rats (age, 6–8 weeks) were purchased from Hunan SJA Laboratory Animal Co., Ltd. (Hunan, China). Animals were housed under controlled temperature (22 ± 2°C), controlled relative humidity (60 ± 5%), and a 12-h/12-h light/dark cycle with ad libitum access to food and water. Animals were allowed to acclimate for 7 days before experiments. All experimental protocols and animal handling procedures were carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals, published by the National Institutes of Health (NIH Publications No. 80–23, revised in 1996). This study was ethically approved by the Animal Care and Use Committee of Renmin Hospital of Wuhan University (Wuhan, China).

This study was divided into two parts. The first part of the study was designed to compare the difference of the gut microbiota between the Sham group, CMI + Dep group, and CMI + Non-Dep group. Fifty male SD rats were randomly used to establish the CMI models and 10 to the sham models. In the second part, 30 male SD rats were randomly used to establish pseudo-germ-free rat models and 10 to the control group. Then fecal microbiota transplantation was performed on the 30 pseudo-germ-free rats. The fecal microbiota for transplantation was collected from the CMI + Dep group and CMI + Non-Dep group rats. Detailed experimental schedule is shown in Figure 1.

Chronic Myocardial Infarction Model

Left anterior descending coronary artery ligation was performed to produce a CMI rat model (Wang et al., 2013; Li, 2018). The rats fasted for 12 h and were deprived of water for 4 h before modeling. We used sodium pentobarbital (50–60 mg/kg intraperitoneal) as anesthetic and confirmed the adequacy of anesthesia by the absence of a withdrawal response to hindpaw nociceptive stimulation. Under sodium pentobarbital anesthesia and intubation, 2-cm incisions were made to the left of and parallel to the sterna in rats. Then, the third and fourth ribs were separated with a clamp, and the hearts were exposed. The pericardium was torn open, and the anterior descending branch of the coronary artery was ligated with 6/0 suture approximately 1–2 mm below the left auricle. Sham group rats underwent an identical surgery but did not sustain a left descending coronary artery ligation. Rats were monitored by electrocardiogram (ECG) before and after coronary artery ligation. Ischemia was confirmed by the myocardial surface becoming blanched, and ST–T segment elevation in ECG recordings was performed on the rats. After the surgery, lidocaine cream was administered to the wound locally twice daily for 3 days for incision pain. Four weeks later, we performed the following evaluations (Lin et al., 2009; Ito et al., 2013).

Cardiac Function and Cardiac Morphology

Cardiac function was evaluated by using echocardiography (Lin et al., 2009; Ito et al., 2013). Under light sodium pentobarbital anesthesia with spontaneous respiration, we performed serial M-mode echocardiography by an echocardiography system (VividTM E95, French). We measured left ventricular internal diameter at end-diastole (LVIDd), left ventricular internal diameter at end-systolic (LVIDs), and left ventricular ejection fraction (LVEF), the posterior left ventricular (LV) wall, and the percent fractional shortening (%FS). The criterion of a successful CMI model was LVEF less than 60% and the percent fractional shortening (%FS) less than 30% (Zhang et al., 2010).
After all the experiments were done, rats were sacrificed, and their hearts were repeatedly rinsed with phosphate-buffered saline (PBS) buffer solution to remove the redundant tissues such as right ventricle, right atrium, and great vessels. Ventricular muscle tissues were taken and fixed with 4% paraformaldehyde solution. Samples were alcohol-dehydrated and paraffin-embedded, sections were made, and Masson staining was performed. Masson stain blue was used for collagen fibers and red for cardiomyocytes. The experimental process is shown in Figure 1.

Behavioral Tests
Depression-like behavior was evaluated by SPT and FST. All tests were conducted individually, in the morning (between 9 and 11 a.m.). The detailed schedule is shown in Figure 1.

Sucrose Preference Test
This test consisted of a two-bottle procedure in which rats were given the choice between water and 1% sucrose solution. Rats were habituated to drink water from two bottles during days 1 and 2. At day 3, rats were deprived of water and food. At day 4, rats were exposed to two identical bottles again: one was water, and another was 1% sucrose solution. The bottles containing water and sucrose were weighed before and at the end of this period. The sucrose preference [sucrose consumption/(sucrose consumption + water consumption)] was used as a measure for depression-like behavior (Iniguez et al., 2014). The liquid used in the experiment was sterile.

Forced Swimming Test
In FST, rats were forced to swim under inescapable conditions. Initially, rats tried to escape but eventually adopted a static posture in which they make only the movements necessary to maintain their head above water. The immobility time was used as a measure for depression-like behavior. Specifically, all rats were placed individually in a clear plastic cylinder (diameter: 25 cm; height: 35 cm) containing 20 cm of water, maintained at 23 ± 1°C. Rats were forced to swim for five consecutive minutes, and immobility time was recorded (Iniguez et al., 2014; Yang et al., 2019).

Pseudo-Germ-Free Rats Modeling
Forty male SD rats were acclimated to a new environment for 7 days. Thirty rats were randomly chosen to perform the pseudo-germ-free rat models and 10 to the control group. Broad-spectrum antibiotics [ampicillin 1 g/L, neomycin sulfate 1 g/L, metronidazole 1 g/L (Amresco Co., Ltd., United States)] were dissolved in drinking water and administered *ad libitum* to the pseudo-germ-free group rats for 14 consecutive days. At the same time, the control group rats were administered to drinking water *ad libitum*. All drinking water was sterile and renewed every day.

Fecal Samples Collection
After finishing the behavioral tests, feces from each group of rats were collected individually. All rats were separately placed in a clean cage containing sterilized filter paper. Fecal samples were collected immediately after defecation in a sterilized centrifuge tube. The filter paper was replaced for each rat. Feces were
stored in a −80°C freezer until analysis or transplantation (Yang et al., 2017).

Fecal Microbiota Transplantation

Fecal microbiota was prepared by diluting 1 g of fecal sample obtained from CMI + Dep group rats in 10 ml of sterile saline. The fecal material was suspended, and 0.2 ml of the suspension was administered by gavage to each recipient pseudo-germ-free rats for 14 consecutive days (Ge et al., 2017). Fecal samples from the CMI + Dep in the first experiment were randomly mixed. Rats that received fecal microbiota transplantation of the CMI + Dep group were marked as the Dep group. Fecal microbiota transplantation of the CMI + Non-Dep groups was performed in the same way. And the rats that received fecal microbiota transplantation of the CMI + Non-Dep group were marked as the Non-Dep group. Subsequently, behavioral tests were measured, and fecal samples of the Dep and Non-Dep group rats were collected for 16S rRNA gene sequencing analysis. The experimental process is shown in Figure 1.

16S rRNA Gene Sequencing of Fecal Samples

16S rRNA gene is mainly composed of defensive zone and variable zone. The conserved region represents the affinity between the microbiome, and the variable region represents the difference. Among them, V3–V4 had good specificity and complete database information (Claridge, 2004). In view of this, we chose to use regions V3–V4 for bacterial diversity analysis. Fecal samples were collected after all behavioral tests, placed in 1.5-ml tubes, snap frozen on dry ice, and stored at −80°C prior to 16s rRNA gene sequencing at Shanghai OE Biotech. Co., Ltd (Shanghai, China). We used Vsearch (Version 2.4.2) (Rognes et al., 2016) to classify operational taxonomic units (OTUs) according to 97% similarity of valid tags obtained from quality control and selected the sequence with the maximum abundance in each OTU as the representative sequence of this OTU. The RDP Classifier Naive Bayesian classification algorithm (Wang et al., 2007) was used to compare and annotate the representative sequence with the database to obtain the annotation information of OTUs. All sequences were classified using the National Center for Biotechnology Information BLAST and SILVA databases. Distance calculation, OTU clustering, rarefaction analysis, and estimator calculation (α-diversity and β-diversity) were performed using the MOTHUR program (Sun et al., 2016).

Receiver Operating Characteristic Curve Analyses

Receiver operating characteristic (ROC) curves illustrated the diagnostic ability of a binary classifier system with the true positive rate (sensitivity) as the ordinate and the false positive rate (1-specificity) as the abscissa. The ROC curves were used to distinguish rats with CMI-induced depression-like behavior from all other rats. The value of the area under the curve (AUC) represents the accuracy of the diagnosis.

Statistical Analysis

Data was presented as the mean ± standard error of the mean (SEM). All statistical analyses were performed using GraphPad Prism 8 (GraphPad Software, San Diego, CA, United States). For the hierarchical cluster analysis of SPT results to define CMI + Dep and CMI + Non-Dep groups, the data were first standardized as Z scores. Hierarchical cluster analysis was performed using Ward’s method and applying squared Euclidean distance as the distance measure. Correlation analysis was conducted using Pearson’s product-moment coefficient. The diagnostic cutoff value, sensitivity, specificity, and accuracy of each bacterium were determined using ROC curve analysis. Group means were compared by one-way analysis of variance (ANOVA), followed by post hoc Tukey’s tests for pairwise comparisons. A p < 0.05 (two-tailed) was considered significant for all tests.

RESULTS

Differences in Depression-Like Behaviors Among Control, CMI + Dep, and CMI + Non-Dep Rats

Echocardiography revealed that LVEF and %FS were both significantly greater in Sham rats than in CMI rats (84.83 ± 3.22% vs. 52.35 ± 6.06%, 48.77 ± 4.2% vs. 22.97 ± 3.04%, p < 0.05; Figures 2A,B). In the Sham group, Masson staining results showed that left ventricular myocardial cell aligned neatly with no obvious pathological changes. But the CMI group rats’ left ventricular myocardial cells were arranged in a disorderly manner, shapes were irregular, and there were a lot of blue collagen fibers and extensive fibrosis between myocardial cells (Figures 2C,D). All of these indicated the success of CMI modeling. And then CMI rats were divided into depression susceptible (CMI + Dep) and depression unsusceptible (CMI + Non-Dep) groups according to the results of hierarchical cluster analysis of SPT (Figure 3A). Compared to the Sham and CMI + Non-Dep groups, the immobility times of FST were significantly increased in CMI + Dep group (48.17 ± 13.55 s vs. 62.29 ± 19.01 s vs. 108.67 ± 34.31 s, p < 0.05; Figure 3C). And in SPT, sucrose preference was significantly decreased in CMI + Dep group but not in the sham or CMI + Non-Dep groups (69.33 ± 8.25 % vs. 88.90 ± 4.56% vs. 85.00 ± 1.85%, p < 0.05; Figure 3B). Therefore, CMI + Dep rats showed depression-like behavior.

Differences in Gut Microbiota Profile Among Control, CMI + Dep, and CMI + Non-Dep Rats

We used 16S rRNA gene sequencing to determine alterations in gut microbiota composition among the Sham group,
the CMI + Dep group, and the CMI + Non-Dep group rats after left coronary artery ligation. After quality control, the data volume of clean tags was distributed between 38,106 and 41,514. Valid tags were obtained by removing chimerism from clean tags and were finally used for analysis. The data volume of valid tags was distributed between 34,129 and 37,599, and the average length ranged from 429.12 to 433.94 bp. In the first experiment, the number of OTU in each sample ranged from 698 to 1,201 (Figure 4E). The number of differential OTUs was 100, and the number of differential genera was 10. Differential profiles of gut microbiota existed among the three groups. Unweighted unifrac diversity distance suggested remarkable differences in gut microbiota among the three groups (Figure 4A). Although Shannon and Simpson indices failed to show such a difference (Figures 4B,C), three-dimensional data of principal component analysis (PCA) revealed separated positions among groups (Figure 4D). PCA was a visualization method to make a comparison between sample diversity. In three-dimensional data of PCA, each dot represented a sample, and similar samples would be clustered together. Most of the dots from the sham, CMI + Dep, and CMI + Non-Dep groups were aggregated separately (Figure 4D).

**Differential Levels of Gut Microbiota Among Sham, CMI + Dep, and CMI + Non-Dep Rats**

16S rRNA gene sequencing revealed that a total of eight gut bacteria at six phylogenetic levels (phylum, class, order, family, genus, and species) significantly differed among the Sham group, the CMI + Dep group, and the CMI + Non-Dep group (Figures 5A–H). At the class level, the relative abundances of *Actinobacteria* were significantly lower in the CMI + Dep group than Sham group (0.000299 ± 0.000138 vs. 0.000635 ± 0.000234, p < 0.05; Figure 5B).
Correlations Between Depression-Like Behaviors and Specific Gut Bacteria
Sucrose preference test and FST were adopted as measures to evaluate the effects of gut microbiota composition on depression-like behaviors. Correlations between the sucrose preference percentage of the CMI + Dep group and the CMI + Non-Dep group and the relative abundance of eight bacteria were analyzed. Correlations between the immobility time and the relative abundance of eight bacteria were also analyzed. Although the results were not statistically significant, they showed clear trends in the influence of gut microbiota on depression-like behaviors. The relative abundance of class Alphaproteobacteria, order Rhodospirillales, family Rs-E47_termite_group, and genus Roseburia negatively influenced depression-like behavior in rats (Figures 6A,B).

Evaluation of Gut Bacteria for the Diagnosis of Chronic Myocardial Infarction-Induced Dep Using Receiver Operating Characteristic Curve Analysis
Receiver operating characteristic curves were constructed to evaluate the diagnostic ability of the gut bacteria in depression (Figure 7). The best cutoff values, sensitivity, specificity, and accuracy as well as the positive and negative predictive values of gut bacteria for the diagnosis of CMI-induced Dep are summarized in Table 1.

Effects of CMI + Dep and CMI + Non-Dep Gut Microbiota Transplantation on Depression-Like Behaviors in Pseudo-Germ-Free Rats
Pseudo-germ-free models were established by administering antibiotics for 14 days. Gut microbiota from CMI + Dep group and CMI + Non-Dep group rats were transplanted into the gastrointestinal tract of pseudo-germ-free rats through feces for another 14 consecutive days (Figure 1D). FST and SPT were performed on days 29–33 (Figure 1C). Compared to the control, vehicle, and Non-Dep group, sucrose preference percentage obviously decreased in Dep group, and the immobility time of FST also significantly increased (77.61 ± 14.63% vs. 89.36 ± 5.95% vs. 92.34 ± 3.88% vs. 38.95 ± 17.76%, 39.00 ± 9.68 s vs. 27.83 ± 11.72 s vs. 39.83 ± 10.09 s vs. 63.50 ± 5.44 s, p < 0.05, Figures 8A,B). Therefore, gut microbiota from CMI + Dep rats appeared to induce depression-like behavior in pseudo-germ-free rats.
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FIGURE 4 | Differences in gut microbiota profiles among Sham, CMI + Dep, and CMI + Non-Dep group rats. (A) Unweighted unifrac diversity distance. (B) Shannon index (one-way ANOVA; F2, 16 = 0.3482, p > 0.05). (C) Simpson index (one-way ANOVA; F2, 16 = 0.1202, p > 0.05). (D) PCA analysis of gut microbiota. (E) Flower plot of OTUs distribution. The numbers in the core represented the OTUs shared by all samples, and the numbers on the petals represented the total OTUs of each sample minus the number of OTUs shared. The α-diversity was shown as mean ± SEM. (n = 6–7 individual fecal samples/group). N.S., not significant. CMI, chronic myocardial infarction; Dep, depression-like behavior; Non-Dep, non-depression-like behavior; PCA, principal component analysis.

FIGURE 5 | Differences in the relative abundance of various gut microbes among the Sham group, the CMI + Dep group, and the CMI + Non-Dep group. (A) Class Alphaproteobacteria (F2, 16 = 7.358, p < 0.05). (B) Class Actinobacteria (F2, 16 = 4.512, p < 0.05). (C) Order Rhodospirillales (F2, 16 = 7.270, p < 0.05). (D) Family Streptococcaceae (F2, 16 = 4.685, p < 0.05). (E) Family Rs-E47_termite_group (F2, 16 = 3.540, p < 0.05). (F) Genus Roseburia (F2, 16 = 5.120, p < 0.05). (G) Genus Streptococcus (F2, 16 = 4.685, p < 0.05). (H) Genus Prevotellaceae_UCG-001 (F2, 16 = 3.831, p < 0.05). Data are shown as mean ± SEM. (n = 6–7) by one-way ANOVA. CMI, chronic myocardial infarction; Dep, depression-like behavior; Non-Dep, non-depression-like behavior. *p < 0.05, **p < 0.01, N.S., not significant.
Differences in Gut Microbiota Composition Between Pseudo-Germ-Free Rats Receiving CMI + Dep or CMI + Non-Dep Rats Fecal Bacteria

16S rRNA gene sequencing result showed that the number of OTUs ranged from 688 to 1,135. A flower plot showed that gut microbiota in the four groups were distinct (Figure 9E). There were 625 different OTUs, 57 different genera, and 6 different phyla. A plot of unweighted unifrac diversity distance demonstrated notable differences in gut microbiota composition among control pseudo-germ-free rats and those receiving vehicle, CMI + Dep rats fecal bacteria, or CMI + Non-Dep rats fecal bacteria (Figure 9A). α-Diversity refers to the diversity of species and bacteria within a community or habitat. Shannon as well as Simpson indices are commonly used to evaluate the α-diversity of gut microbiota (Zhan et al., 2018). Dep group remarkably restored the decrease of Shannon index compared to vehicle groups (6.56 ± 0.17 vs. 5.77 ± 0.20, p < 0.05, Figure 9B), whereas there was no significant change in Simpson index between Dep groups and vehicle groups (Figure 9C). β-Diversity using principal coordinates analysis (PCoA) demonstrated the differentiation among habitats (Guida et al., 2018). Clearly, the dots of the control group were individually separated from the others. And the dots of Dep and Non-Dep groups were also far away from the vehicle group. Compared with Non-Dep group, the composition of gut microbiota in Dep group showed a different profile (Figure 9D). Therefore, it was extremely possible that the composition of the gut microbiota was significantly different between control, vehicle, Dep, and Non-Dep groups.

Differential Levels of Gut Microbiota in Pseudo-Germ-Free Rats

Fecal transplantation significantly altered the relative abundances of gut microbiota in Dep and Non-Dep rats. At six levels, a total of 25 gut bacteria in the fecal samples had substantial differences
among control rats and rats receiving vehicle, Dep rats fecal bacteria, or Non-Dep rats fecal bacteria (Figure 10). Compared to the Non-Dep group, the relative abundance of the phylum Gemmatimonadetes, the other orders, the family Rikenellaceae, the family Bacteroidales RF16 group, the other families, the genus Rikenellaceae_RC9_gut_group, and the genus Ruminococcus_1 significantly increased in Dep group (Figures 10B,J,L,M,O,S,T), whereas the relative abundance of the family Streptococcaceae and the species Lactococcus_garvieae_subsp._garvieae decreased in Dep group (Figures 10N,X).

**DISCUSSION**

Left anterior descending coronary artery ligation has been commonly used to create myocardial infarction models (Wang et al., 2013; Li, 2018). In the present study, results of cardiac function and cardiac morphology suggested successful creation of CMI models. In humans, depression is a common psychiatric symptom of CMI and is an independent risk factor for disease prognosis. A higher risk of developing depression in patients with CMI suggests a shared pathogenesis, although the mechanisms remain unknown (Penninx et al., 2001). The SPT and FST could well assess the symptoms of depression-like behavior in rodents. And several lines of evidence observed depression symptoms in rodents approximately 4 weeks after left anterior descending coronary artery ligation (Lin et al., 2009; Ito et al., 2013). In this study, CMI rats could be stratified into CMI + Dep and CMI + Non-Dep groups by using the hierarchical cluster analysis of SPT results. The CMI + Dep rats also demonstrated longer immobility time in FST compared with both sham-operated rats and CMI + Non-Dep rats.
Gut microbiota diversity has been strongly associated with mood-relating behaviors, such as depression (Lin et al., 2009; Winter et al., 2018). Accumulating evidences indicated that gut microbiota influenced central neurochemistry and behavior, which is called the microbiota–gut–brain axis (Cryan and O’Mahony, 2011; Dinan and Cryan, 2017). In the present study, we observed no significant difference in α-diversity (consisting of Shannon and Simpson indices) among the Sham, CMI + Dep, and CMI + Non-Dep groups. However, the separation of groups according to β-diversity (PCA) suggested that the microbiota composition was significantly altered by the comorbidity of CMI and depression-like behavior. To the best of our knowledge, this was the first study to demonstrate the role of gut microbiota in individual differences of depression-like phenotype in rats with CMI. Additionally, this was also the first study adopting hierarchical cluster analysis to study the comorbidity of CMI and depression-like phenotypes in rodents.
In the present results, gut microbiota was significantly altered in the CMI + Dep or CMI + Non-Dep rats compared to sham-operated rats, suggesting a role of gut microbiota in the individual differences of depression-like phenotype. Furthermore, transplantation of fecal microbiota from CMI + Dep rats induced depression-like behavior in the antibiotics-treated pseudo-germ-free rats. However, transplantation of fecal microbiota from Non-Dep rats showed no tendency to depression-like behavior in the antibiotic-treated pseudo-germ-free rats, compared to the control group. Distinct gut microbial compositions were identified following fecal microbiota transplant. Taken all together, these results suggest that change in gut microbiota played an important role in the depression-like phenotypes in rats with CMI.

*Streptococcus* accounted for a large amount of deaths worldwide through diverse disease manifestations. Cardiac injury and life-threatening acute cardiac complications are more common in *Streptococcus* infection, compared to other bacterial infections (Alhamdi et al., 2015). Intriguingly, *Streptococcus* was recently reported to be higher in depression patients compared to healthy people (Barandouzi et al., 2020). This result was consistent with our findings that the phylum *Streptococcus* and the genus *Streptococcus* were enriched in the fecal samples of Dep rats. Therefore, *Streptococcus* infection was possibly associated with the onset and symptom of depression-like behavior in CMI rats.

Our study revealed significant inconsistencies in the abundance of *Rikenellaceae*, *Bacteroidales_RF16_group*, *Ruminococcaceae_UCG-014*, and *Lactobacillus_gasseri*. High abundance in *Rikenellaceae*, *Bacteroidales_RF16_group*, and *Ruminococcaceae_UCG-014* was observed in antibiotics-treated pseudo-germ-free rats treated with fecal microbiota from CMI + Dep rats. In contrast, *Lactobacillus_gasseri* observably decreased in antibiotics-treated pseudo-germ-free rats treated with CMI + Dep rats' fecal microbiota,
In conclusion, the CMI-induced depression-like behavior was notably associated with dysbiosis in gut microbiota, and change in specific gut bacteria may be involved in the pathogenesis of this comorbidity. In light of the links between depression and CMI, the regulation of gut microbiota may be a potential therapeutic target for depression. However, further detailed studies are clearly needed to validate the role of gut microbiota in the pathogenesis and therapeutic mechanisms of the depression-like behavior in CMI patients.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The animal study was reviewed and approved by Renmin Hospital of Wuhan University.

AUTHOR CONTRIBUTIONS

All authors critically reviewed and approved the final version for submission.

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