ORIGINAL RESEARCH

Alterations of the Gut Microbiota in Response to Total Sleep Deprivation and Recovery Sleep in Rats

Zhong Wang,^{1,*} Kai Yuan,^{1,*} Yan-Bin Ji,^{2,*} Su-Xia Li, ¹⁰ Le Shi,¹ Zhe Wang,¹ Xin-Yu Zhou,⁴ Yan-Ping Bao,³ Wen Xie,⁵ Ying Han,³ Jie Shi,³ Lin Lu,^{1,3,6} Wei Yan,¹ Wen-Hao Chen ¹⁰

¹Peking University Sixth Hospital, Peking University Institute of Mental Health, NHC Key Laboratory of Mental Health (Peking University), National Clinical Research Center for Mental Disorders (Peking University Sixth Hospital), Peking University, Beijing, 100191, People's Republic of China; ²Department of Neurology, Qilu Hospital of Shandong University, Shandong University, Jinan, 250012, People's Republic of China; ³National Institute on Drug Dependence and Beijing Key Laboratory of Drug Dependence, Peking University, Beijing, 100191, People's Republic of China; ⁴Department of Psychiatry, The First Affiliated Hospital of Chongqing Medical University, Chongqing, 400016, People's Republic of China; ⁵Mental Health Center of Anhui Province, Hefei, 230032, People's Republic of China; ⁶Peking-Tsinghua Center for Life Sciences and PKU-IDG/McGovern Institute for Brain Research, Peking University, Beijing, 100871, People's Republic of China

*These authors contributed equally to this work

Correspondence: Wen-Hao Chen; Wei Yan Peking University Sixth Hospital/Peking University Institute of Mental Health, 51 Huayuanbei Road, Haidian District, Beijing, 100191, People's Republic of China Tel +86-10-82801342; Tel +86-10-62723704 Email chenwenhao@bjmu.edu.cn; weiyan@bjmu.edu.cn **Introduction:** Accumulating evidence suggests that both sleep loss and gut dysbiosis can lead to metabolic disorders. However, less is known about the impact of total sleep deprivation (SD) and sleep recovery on the composition, function, and metabolic dynamics of the gut microbiota.

Methods: Specific-pathogen free Sprague–Dawley rats were subjected to 48 h of SD with gentle handling and then allowed to recover for 1 week. Taxonomic profiles of fecal microbiota were obtained at baseline, 24 h of SD, 48 h of SD, and 1 week of recovery. We used 16S rRNA gene sequencing to analyze the gut microbial composition and function and further characterize microbiota-derived metabolites in rats.

Results: The microbiota composition analysis revealed that gut microbial composition and metabolites did not change in the rats after 24 h of SD but were significantly altered after 48 h of SD. These changes were reversible after 1 week of sleep recovery. A functional analysis was performed based on Kyoto Encyclopedia of Genes and Genomes (KEGG) annotations, indicating that 19 KEGG pathways were significantly altered in the gut microbiota in SD rats. These functional changes occurred within 24 h of SD, were more apparent after 48 h of SD, and did not fully recover after 1 week of sleep recovery.

Conclusion: These results indicate that acute total SD leads to significant compositional and functional changes in the gut microbiota, and these changes are reversible.

Keywords: gut microbiota, sleep deprivation, short-chain fatty acid, function prediction

Introduction

Sleep plays a critical role in physical and mental health maintenance and well-being throughout an individual's lifetime. Sleep loss is typically associated with impairments in performance, negative health consequences, and lower survival.^{1,2} Sleep deprivation (SD) is a widespread phenomenon that is associated with adverse metabolic consequences,^{3–5} such as obesity, insulin resistance, diabetes, and cardiovascular disease, that ultimately promote the emergence of metabolic dysfunction for reasons that are still unclear.⁶

The gut microbiome is a vital component of the intestinal environment. It has been identified as a critical environmental factor that contributes to the development of obesity,⁷ insulin resistance,⁸ and diabetes.⁹ The gut microbiota-brain axis has become an increasingly popular research focus.¹⁰ Alterations of the intestinal microbiota have been linked to the pathophysiology of many psychiatric disorders, such as autism spectrum disorder,¹¹ schizophrenia,¹² and depression,^{13–15} suggesting that the gut microbiota affects multiple aspects of brain development and neuroendocrinological function. Research also demonstrates that microbiota-derived metabolites, such as

Nature and Science of Sleep 2022:14 121-133

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short-chain fatty acids (SCFAs),¹⁶ are primarily produced in the large intestine through anaerobic bacterial fermentation and can maintain both intestinal immune function and gut barrier function.¹⁷ Accumulating evidence indicated that SCFAs played an important role in maintaining health and preventing disease.¹⁸ Recent studies showed that butyrate treatment led to rapid and robust increases in non-rapid-eyemovement (NREM) sleep in rats,¹⁹ highlighting the associations between sleep and gut microbial metabolites.

The gut microbiota fluctuates in response to circadian rhythm and routine schedules.^{20,21} A widely held presumption is that there is a relationship between sleep and the gut microbiome. Both sleep loss and dysbiosis of the gut microbiome are associated with metabolic diseases, such as obesity and diabetes.^{7,9,22} Several studies reported preliminary evidence that the gut microbiota was involved in sleep disorders in both murine models and human patients. Insomnia was shown to lead to significant structural and functional changes in the gut microbiota.^{23,24} A previous study found that the composition of the gut microbiota was subtly affected, with a higher Firmicutes/Bacteroidetes ratio in human subjects with sleep restriction.²⁵ In contrast, recent work suggested that a single brief period of SD (5 h) did not alter the overall microbial composition.²⁶ These preliminary studies highlighted the potential relationship between SD and changes in the gut microbiota. To date, no studies have comprehensively examined the impact of total SD and sleep recovery on the composition, function, and metabolic dynamics of the gut microbiota. The present study investigated the effects of acute total SD and 1 week of sleep recovery on microbiota populations. We used 16S ribosomal RNA (rRNA) gene sequencing to evaluate the compositional and functional changes in the gut microbiota in response to SD and sleep recovery.

Materials and Methods Rat Sleep Deprivation and Sample Collection

Adult male specific-pathogen free Sprague-Dawley rats (7–8 weeks old) were housed individually under a 12 h/12 h light/ dark cycle (lights on from 8:00 AM to 8:00 PM) and a controlled temperature of 21–22°C and humidity of 55% \pm 5%. Standard laboratory irradiated chow food (Cobalt 60) and sterile drinking water were provided ad libitum. The rats (n = 8) were subjected to 48 h of SD and then allowed to recover for 1 week. The rats were in their home cages during SD. To reduce the rats' stress, they were kept awake by gently

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tapping and shaking the cage by an experimenter.^{27,28} The control group (n = 8) had a regular sleep-wake cycle in the same environment (Figure 1A). The weight of the rats was recorded over the deprivation period. Each day, the same amount of food was provided to the SD and non-SD groups. Fecal pellets were freshly collected (8:00 AM of d4, d5, d6 and d13), snap-frozen, and stored at -80° C for later analysis. The animal protocols were carried out in accordance with the Administration System of Laboratory Animals in China and were approved by the Research Ethics Committee of Peking University Sixth Hospital.

16S rRNA Gene Sequencing

The procedures for preparing fecal samples were described previously.¹⁴ Briefly, DNA from fecal samples was extracted using the Qiagen QIAmp Fast DNA Stool Mini Kit (Qiagen, Valencia, CA, USA) coupled with an initial bead-beating step as previously described.²⁹ DNA was quantified with a Qubit fluorometer using the Qubit dsDNA BR Assay kit (Invitrogen, Carlsbad, CA, USA). Quality was checked by running an aliquot on 1% agarose gel. Polymerase chain reaction (PCR) enrichment was performed in a 50 µL reaction that contained 30 ng of the template, fusion PCR primer, and PCR master mix. The PCR products were purified with Ampure XP beads and eluted in elution buffer. Total RNA was quantified by spectrofluorimetry and qualified by capillary electrophoresis using Agilent Bioanalyzer 2100 (Agilent, Palo Alto, CA, USA). The validated libraries were used for 2×300 bp paired-end sequencing with the Illumina HiSeq platform (BGI, Shenzhen, China) following the standard pipelines.

Determination of Short-Chain Fatty Acid Concentrations

SCFA levels were determined by gas chromatography-mass spectrometry (GC-MS) in frozen fecal samples that were freshly collected from the rats. The method was adopted from Zhang et al.³⁰ Helium was used as the carrier gas at a constant flow rate of 1 mL/min. The initial oven temperature was held at 60°C for 5 min, increased to 250°C at 10°C/min, and finally held at 250°C for 5 min. The temperatures of the front entrance, transmission line, and electron impact (EI) ion source were set to 280°C, 250°C, and 230°C, respectively. Data processing was performed using an Agilent MSD ChemStation (Agilent). The reported values were normalized to the weight of the original samples that were used.

Statistical Analysis

The statistical analyses were performed using SPSS 20.0 software (IBM, Chicago, IL, USA). The data were assessed for a normal distribution and are expressed as the mean \pm SEM. Differences were assessed using repeated-measures analysis of variance (ANOVA) followed by the Bonferroni post hoc test. Values of p < 0.05 were considered statistically significant.

Results

Effects of Sleep Deprivation on Microbial α -Diversity

Different diversity indices (ie, Observed species index, Chao1 index, Shannon index, and Simpson index) were used to assess gut microbiota α -diversity (Figure 1B-E). During SD, the mixed-design ANOVA, with group (sleep deprivation and normal sleep control) as the between-groups factor and time of day as the within-groups factor, revealed a main effect of time of day ($F_{3.42} = 9.26, p < 0.001$) on the Shannon index of α -diversity and a significant time of day \times group interaction ($F_{3,42} = 6.54$, p < 0.001; Figure 1D). The ANOVA analysis also showed a significant main effect of time of day ($F_{3,42} = 5.56$, p = 0.003) on the Simpson index and a significant time of day \times group interaction ($F_{3,42}$ = 4.95, p = 0.005; Figure 1E). We also found a significant effect of time of day on the Observed species index ($F_{3.42} = 9.23$, p < 0.001). The post hoc tests revealed no difference in the α diversity of the gut microbiota between baseline (Shannon index: p = 0.471; Simpson index: p = 0.318) and after 24 h of SD (Shannon index: p = 0.223; Simpson index: p = 0.329) in SD rats compared with control rats but a significant reduction (Shannon index: p = 0.017; Simpson index: p = 0.021) after 48 h of SD, and this difference disappeared (Shannon index: p = 0.693; Simpson index: p = 0.529) after 1 week of sleep recovery. These results indicate that 48 h of SD reduced adiversity of the gut microbiota, which fully recovered after 1 week of sleep recovery.

Effects of Sleep Deprivation on Microbial β -Diversity

We further examined the effects of SD on microbial β diversity, which was performed based on a weighted UniFrac analysis.³¹ We applied the weighted UniFrac rank (Figure 2A), which provided a measure of the evolutionary distance between microbiotas. We found that the difference between groups was significantly greater than the difference within groups (R = 0.16, p = 0.001). Microbial communities clustered using principal coordinates analysis (PCoA) of the weighted UniFrac matrix. The percentage of variation that was explained by the principal coordinates was indicated on the axes in Figure 2B. PCoA1 represented the main percentage variation (57.82%), revealing significant main effects of time of day ($F_{3,42} = 7.43$, p < 0.001) and group ($F_{1,14} = 5.41$, p = 0.036) and a significant time of day \times group interaction $(F_{3,42} = 2.92, p = 0.045)$. The PCoA weighted Unifrac matrix analysis showed that the gut microbiome exhibited a trend towards differences (p = 0.068) after 24 h of SD, which became more significant after 48 h of SD (p =0.004), compared with the normal sleep control group. These differences disappeared (p = 0.597) after 1 week of sleep recovery. These results indicated that 48 h of SD altered β -diversity of the gut microbiota, which fully recovered after 1 week of sleep recovery.

Effects of Sleep Deprivation on Microbial Composition at the Genus Level

To more closely examine the impact of SD on the microbiome, we examined the changes in microbiota composition at the genus level in response to our intervention. The mixed-design ANOVA, with group (SD and normal sleep control) as the between-groups factor and time of day as the within-groups factor, revealed six genera that significantly changed after SD, with a significant time of day \times group interaction. There were no significant differences in these altered genera after 24 h of SD (all p > 0.05). After 48 h of SD, the relative abundance of g Butyricicoccus (p = 0.028), g Butyricimonas (p = 0.020), g Alistipes (p = 0.028)(0.006), g Intestinimonas (p = 0.001), and g Lactobacillus (p = 0.014) decreased (Figure 3A-E), whereas the relative abundance of g Streptococcus (p = 0.040) increased (Figure 3F). After 1 week of sleep recovery, most of the different microbial species recovered such that no differences from the control were found.

Changes in the Stool Content of Short-Chain Fatty Acids Following Sleep Deprivation

Our results showed that the abundance of SCFAs-producing microbes, such as $g_Butyricicoccus$,³² $g_Butyricimonas$,³³ and $g_Alistipes$,³⁴ markedly decreased after SD. We further analyzed the levels of total SCFAs and five different SCFAs in samples that were obtained before and after SD and after 1 week of sleep recovery (Figure 4). We found a significant



Figure I Effects of sleep deprivation (SD) on microbial α -diversity in rats. (A) Experimental design and protocol. (B–E) Comparison of (B) Observed species index, (C) Chaol index, (D) Shannon index, and (E) Simpson index between SD and control groups following SD and recovery. The data are presented as mean ± SEM. *p < 0.05.

effect of time of day on acetate ($F_{3,42} = 6.00$, p = 0.002), propionate ($F_{1.82,25.48} = 12.61$, p < 0.001), valerate ($F_{3,42} = 4.73$, p = 0.006), and the total amount of fecal SCFAs ($F_{3,42} = 6.27$, p = 0.001). We found a significant time of day × group interaction for butyrate ($F_{3,42} = 2.86$, p = 0.048), which was significantly reduced after 48 h of SD (p = 0.012). After 1 week of sleep recovery, butyrate concentrations returned to similar levels as controls (p = 0.227; Figure 4D). The correlation analysis showed that fecal butyrate concentrations positively correlated with the relative abundance of $g_Butyricimonas$ (r = 0.39, p = 0.001; Figure 4G). We did not find any interaction for the total amount of SCFAs and other four SCFAs.

Sleep Deprivation Led to Significant Functional Changes in the Gut Microbiota

A phylogenetic investigation of communities by the reconstruction of unobserved states (PICRUSt) algorithm was performed to assess the functional differences by plotting differential pathways against the KEGG database with significantly different abundances between SD and control rats.^{35,36} A total of 637 taxonomies and KEGG ontologies were parsed and mapped into 328 KEGG modules. Compared with the control group, a total of 17 KEGG pathways (energy metabolism, oxidative phosphorylation, carbohydrate metabolism, glycolysis/gluconeogenesis,

lipid metabolism, fatty acid metabolism, propanoate metabolism, butanoate metabolism, amino acid metabolism, and tryptophan metabolism, among others) significantly decreased, and two KEGG pathways (lipopolysaccharide biosynthesis and lipopolysaccharide biosynthesis proteins) significantly increased in SD rats (Figure 5A). These differential pathways are mainly involved in four biological processes, especially in various metabolic pathways, such as amino acid metabolism. These results suggested that amino acids, carbohydrates, and lipid metabolism were predominantly disturbed after SD. These functional changes appeared after 24 h of SD and were even more significant after 48 h of SD but with partial restoration after 1 week of sleep recovery. To explore functional correlations between gut microbiota dysbiosis and alterations of fecal metabolites, a correlation matrix was generated using Spearman correlation coefficients between microbial communities at the genus level, SCFAs, and significantly altered KEGG pathways. Overall, the cocorrelation analysis showed that KEGG metabolism pathways formed strong and broad co-correlation relationships with both bacterial species and fecal metabolites (Figure 5B).

Discussion

In the present study, we found that total SD led to significant compositional and functional changes in the gut microbiota, and these changes were partially reversible after 1 week of sleep recovery. We examined the impact of 24 h and 48 h of SD and 1 week of sleep recovery on gut microbiota composition in rats. There were no significant differences in gut microbiota composition after 24 h of SD. Significant alterations of gut microbiota composition were observed after 48 h of SD, and this change disappeared after 1 week of sleep recovery. We also found that functional changes in the gut microbiota occurred within 24 h of SD, which were more apparent after 48 h of SD and not fully recovered after 1 week of sleep recovery.

The microbial composition analysis revealed significant alterations of α - and β -diversities after 48 h of SD, indicating that SD may be linked to dynamic changes in intestinal microbiota composition. Similar to the present study, previous studies found that α - and β -diversities of the gut microbiota were significantly altered in patients with insomnia.^{23,24} A previous study that used an 1-week sleep restriction protocol in both rats and humans found no significant changes in β-diversity or changes in Operational Taxonomic Unit abundance of the gut microbiome.³⁷ A recent study examined a single short period of SD in mice and found no major shifts in gut microbiota composition.²⁶ Consistent with these results, in the present study, no significant difference in gut microbiota composition was found after 24 h of SD. Another study examined the microbiome in humans after 2 days of partial sleep restriction and found that the Firmicutes/ Bacteroidetes ratio but not β -diversity was altered by sleep restriction.²⁵ Furthermore, Poroyko et al. found a significant shift in the microbiome in mice after longterm (4 weeks) sleep fragmentation.³⁸ These findings indicated that the microbiome might be resistant to a certain degree of sleep loss, and discrepancies in changes in the gut microbiota might be attributable to the differences in the duration of sleep loss. A short period and slight degree of SD may not be sufficient to exert apparent consequences with regard to dysbiosis of the gut microbiota.

We also identified six significantly altered microbiota at the genus level after 2 days of SD, in which we compared relative abundance between SD and control rats. Sleep-deprived rats exhibited significantly lower proportions of g Butyricicoccus, g Butyricimonas, g Alistipes, g Intestinimonas, and g Lactobacillus and significantly higher proportions of g Streptococcus. Correlation analysis showed that the relative abundance of g Streptococcus was negatively correlated with the concentration of fecal butyrate and positively correlated with the KEGG pathway of lipopolysaccharide biosynthesis. g Streptococcus are gram-positive aerobic organisms that cause many disorders, including pharyngitis, pneumonia, wound and skin infections, sepsis, and endocarditis.³⁹ The sleep-wake cycle likewise regulates the normal functioning of the immune system, people have been aware of the sleep loss that accompany many diseases, including multiple microbial infections.⁴⁰ Sleep loss may enhance the translocation of viable bacteria from the intestine, which provides another means by which sleep-microbe interactions may impact health.⁴¹

Total SD is the most common laboratory-based method for studying sleep. Many people often experience SD as a consequence of shift work (eg, truck drivers, physicians, etc.).^{28,42} Compared with chronic sleep restriction and sleep fragmentation, total SD causes more acute and severe impairment.⁴³ However, the effect of total SD with more than one circadian rhythm on the gut microbiota



Figure 2 Effects of sleep deprivation on microbial β -diversity in rats. (A) Weighted Unifrac Anosim analysis of the microbiome in rats following SD and recovery. (B) Weighted Unifrac PCoA analysis of microbiome in rats following SD and recovery. Boxes represent the interquartile ranges. Lines inside the boxes denote medians. Circles are outliers. **p < 0.01.



Figure 3 Effects of SD and recovery on the relative abundance of gut microbiota at the genus level, including (A) $g_{Butyricicoccus}$, (B) $g_{Butyricimonas}$, (C) $g_{Alistipes}$, (D) $g_{Intestinimonas}$, (E) $g_{Lactobacillus}$, and (F) $g_{Streptococcus}$. The data are expressed as mean \pm SEM. *p < 0.05, **p < 0.01.

remains unclear. In previous studies, several methods were used for total SD, such as rotating drums and gentle handling. In the present study, the rats were subjected to 48 h of total SD by gentle handling, a technique of gentle stimulation with physical contact used to disrupt the circadian system and sleep. This method of SD may reduce the animals' stress.⁴⁴ Other techniques that have been used for SD in rodents included using a water platform and the presentation of unpleasant stimuli. These techniques,

however, are more stressful for the animals than gentle handling.⁴⁵ Furthermore, environmental factors can contribute to variations in the gut microbiota, leading to inconsistent results.²³ To eliminate the influence of the environmental factors and dietary habits on gut microbial communities, we maintained a strict, standard laboratory environment during the experiment.

Evidence suggested that gut bacteria were a source of sleep-inducing signals.⁴⁶ To determine whether our total



Figure 4 Effects of SD and recovery on the fecal content of SCFAs, including (A) total short-chain fatty acids, (B) acetate, (C) propionate, (D) butyrate, (E) isobutyrate, and (F) valerate. (G) Correlation between fecal butyrate content and the relative abundance of $g_Butyricimonas$. The data are expressed as mean \pm SEM. *p < 0.05.



Figure 5 Sleep deprivation and recovery lead to significant functional changes in the gut microbiota. (A) To predict the metagenome function, the PICRUSt analysis heatmap showed several significantly changed KEGG pathways between groups after SD and recovery. (B) Correlation analysis between signature genus, functional changes, and metabolites of the gut microbiota. Edges between nodes indicate Spearman's negative (light red) or positive (light blue) correlations. Edge thickness indicates a range of correlation coefficient values (all p < 0.05).

SD paradigm functionally affected the intestinal microbiome, we assessed the abundance of fecal SCFAs produced by microbes, such as $g_Butyricicoccus$,³² $g_Butyricimonas$,³³ and $g_Alistipes$,³⁴ markedly decreased after SD, which might potentially influence the production of SCFAs. We found a significant effect of time of day on acetate, propionate, valerate, and the total amount of fecal SCFAs. We also found a significant time of day \times group

interaction for butyrate, in which fecal butyrate significantly decreased after SD compared with the control group. Butyrate is a four-carbon SCFA produced from the fermentation of dietary fibers by microbiota.⁴⁷ Previous studies found that oral or intraportal butyrate administration robustly increased NREM sleep in rats and induced slow, high-amplitude electroencephalographic wave sleep in rabbits.^{19,48} Our results were different from previous studies that imposed sleep restriction. In these previous studies, SCFAs did not change after partial SD. This discrepancy in SCFA levels may be attributable to differences in the degree of sleep loss. The present study also found that fecal SCFA content did not change after 24 h of SD.

Sleep deprivation is a biological stressor that alters metabolic parameters. The gut microbiota is now widely accepted to significantly shape several metabolic pathways in the host.^{12,49} The biological and physiological functions of the gut microbiota can be defined in several ways. Taxonomic composition and diversity are poorly conserved across individuals, whereas genetic composition and functional capacity are evolutionarily conserved across individuals.⁵⁰ Thus, to decipher the metabolic switch of the gut microbiota, the PICRUSt algorithm was applied to map the bacterial genetic pathway against the KEGG database.²³ Compared with the control group, a wide range of pathways were altered in the SD group. These functional changes appeared after 24 h of SD and were even more significant at 48 h of SD. This suggested that some functional changes already appeared during SD before changes in the composition of the gut microbiota were evident. Interestingly, energy metabolism and oxidative phosphorylation pathways significantly decreased in the gut microbiota in SD rats, indicating that SD disturbed energy metabolism.⁵¹ Previous study found that SD affected the capacity of mitochondrial bioenergetics, decreasing respiration with the oxidative phosphorylation and electron transport systems.⁵² The final common metabolic pathway for carbohydrates, lipids, and amino acids is the tricarboxylic acid (TCA) cycle, which is the most effective way for the body to obtain energy through metabolism.⁵³ We found that the carbohydrate, lipid, and amino acid metabolism pathways in the gut microbiota decreased after SD. We also found that glyoxylate, dicarboxylate, and pyruvate metabolism, which belong to the energy and carbohydrate metabolism that were critical for the TCA cycle, were downregulated in the gut microbiota after SD.^{51,54} Sleep disruption was reported to decrease energy metabolism in rats, suggesting a causal effect of sleep loss on weight gain.55,56

Moreover, lipopolysaccharide biosynthesis pathways also increased in the gut microbiota in SD rats. Lipopolysaccharide causes an inflammatory reaction mainly via activation of the Toll-like receptor 4/nuclear factor kB inflammatory pathways. Human and rodent studies found that sleep loss increased peripheral markers of inflammation.^{57–59} Chronic sleep loss can alter the gut microbiota and induce systemic inflammation in mice.³⁸ Disturbances in the metabolism of amino acid neurotransmitters, such as tryptophan, glutamate, and γ -aminobutyric acid (GABA), are prominent after SD. Recent studies showed that the gut microbiome might indicate the onset of mental disorders by modulating amino acid neurotransmitters, such as GABA and serotonin. As a precursor for production of the neurotransmitter serotonin, tryptophan metabolism has also been found to be altered by SD and linked to various neurodegenerative diseases.^{60,61} In the present study, we found that the levels of fecal amino acid neurotransmitters (ie, glutamate, tryptophan, aspartate, and glycine) decreased in SD rats, suggesting a decrease in amino acid neurotransmitter content in the intestine after SD. These findings suggested that fecal amino acid metabolism might be modulated by gut microbes, which might reflect detrimental consequences of sleep loss.

There are some limitations of the current study. First, this study used 16S rRNA gene sequencing analysis without further metagenomic analysis. Although more data and information can be obtained through the shotgun metagenomics, 16S rRNA gene sequencing is currently more commonly used and popular in microbiome research.⁶² Second, unique perturbations to the ileal microbiota that register even after acute SD suggest that analysis of fecal material alone may not be appropriate when assessing dysbiosis of the gut microbiome.²⁰ Third, we have only conducted correlation analysis for gut microbiota and metabolism pathways, which could not provide evidence of causality. We also recognize that the sample size in each group was limited. We plan to explore the mechanism using large sample sizes in future studies.

Conclusion

In the present study, we examined the impact of 24 and 48 h of total SD and 1 week of sleep recovery on gut microbiota composition and function. We found that 48 h but not 24 h of total SD significantly altered composition of the gut microbiota, which were reversible after 1 week of sleep recovery. We also found that functional changes in the gut microbiota occurred within 24 h of SD, which was more apparent

after 48 h of SD and did not fully recover after 1 week of sleep recovery. These findings indicated that SD might be linked to dynamic changes in composition and function of the intestinal microbiota. Future studies should elucidate the molecular mechanisms that are responsible for these changes and how they contribute to behavioral and neurophysiological phenotypes that are associated with sleep loss.

Acknowledgments

This work was supported in part by the National Key Research and Development Program of China (no. 2020YFC2003600), National Natural Science Foundation of China (no. 81901347, 82171477, 31800897, 81821092, and 82001404), Peking University Medicine Fund for Fostering Young Scholars' Scientific and Technological Innovation, and Fundamental Research Funds for Central Universities (no. BMU2018PYB010). We thank Shi-Qiu Meng and Ya-Bin Wei for their help in revising our manuscript. We are grateful to BioNovoGene (Suzhou) for assistance with GC-MS.

Disclosure

The authors declare no conflicts of interest in this work.

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