The Gut Microbiota Dysbiosis in Preeclampsia Contributed to Trophoblast Cell Proliferation, Invasion, and Migration via lncRNA BC030099/NF-κB Pathway

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Background. Preeclampsia (PE) is the main reason of maternal and perinatal morbidity and mortality. Gut microbiota imbalance in PE patients is accompanied by elevated serum lipopolysaccharide (LPS) levels, but whether it affects the occurrence and development of PE, the underlying mechanism is not clear. This paper intends to investigate the relationship between lncRNA BC030099, inflammation, and gut microbiota in PE.

Methods. The feces of the patients were collected, and gut microbiota changes were assessed by 16S rRNA sequencing and pathway analysis by PICRUSt. Next, we examined LPS and lncRNA BC030099 levels in feces or placenta of PE patients. Then, we knocked down lncRNA BC030099 in HTR-8/SVneo cells and added the NF-κB pathway inhibitor JSH-23. CCK-8 and Transwell assays were performed to determine cell proliferation, migration, and invasion. Western blot was utilized to evaluate MMP2, MMP9, snail, and E-cadherin, p-IκBα, IκBα, and nuclear NF-κB p65 levels. IL-6, IL-1β, and TNF-α levels were examined by ELISA.

Results. Gut microbiota was altered in PE patients, and microbial genes associated with LPS biosynthesis were significantly elevated in gut microbiota in the PE group. LPS level in feces and placenta of PE group was significantly elevated. lncRNA BC030099 level in placenta of PE group was also notably promoted. Knockdown of lncRNA BC030099 promoted HTR-8/SVneo cell proliferation, migration, and invasion. Knockdown of lncRNA BC030099 also elevated MMP2, MMP9, and snail levels and repressed E-cadherin level. In addition, lncRNA BC030099 affected NF-κB pathway. Furthermore, NF-κB inhibitor reversed HTR-8/SVneo cell proliferation, invasion, and migration induced by LPS.

Conclusions. The gut microbiota dysbiosis in PE contributed to HTR-8/SVneo cell proliferation, invasion, and migration via lncRNA BC030099/NF-κB pathway.

1. Introduction

Preeclampsia (PE) is a hypertensive disorder of pregnancy involved in 2% to 8% of pregnancy-related complications worldwide [1]. It is the main reason of maternal and perinatal morbidity and mortality [2]. The etiology and pathogenesis of PE have not been fully clarified. Currently, it is mainly believed to be related to insufficient remodeling of uterine spiral arterioles, excessive activation of inflammatory immunity, and damage to vascular endothelial cells [3]. The clinical treatment measures for PE are limited, focusing on the control of acute hypertension, prevention of PE, and timely delivery, and the only effective treatment is delivery of the fetus and placenta [4]. Therefore, an in-depth understanding of the pathogenesis of PE can provide guidance for clinical prediction and prevention of PE, avoid serious complications in pregnant women, and improve pregnancy outcomes.

Long noncoding RNAs (lncRNAs) are a class of RNA molecules with transcripts > 200 nt, which can regulate target gene expression via epigenetic, transcriptional, and post-transcriptional regulation. It is involved in different biological processes including cell differentiation and proliferation, apoptosis, and necrosis [5, 6]. In addition, abnormal lncRNA expression is associated with disease progress [7]. Therefore, lncRNAs’ role in PE pathogenesis has also...
2. Materials and Methods

2.1. Collection of Clinical Samples. We collected feces and placental tissue from PE patients in Xiangya Hospital central South University as the PE group (n = 8). The healthy subjects during the same period were selected as the control group (C group, n = 6). The inclusion criteria of this study were as follows: aged at 25-40 years; gestational weeks from 32 to 39; singleton; no fetal abortion and stillbirth; no history of adverse pregnancy and childbirth; and no history of smoking, drinking, and other drugs. The exclusion criteria were multiple pregnancy; diabetes, chronic hypertension, and kidney disease or other pregnancy complications before pregnancy; antibiotics, glucocorticoids, and immunosuppressants within 1 month before stool collection and other drugs [12]. Fecal samples were collected in stool collection tubes and stored at -80°C until further processing.

2.2. 16S rRNA Sequencing and PICRUSt Pathway Analysis. Fecal samples from healthy subjects (n = 8) and PE patients (n = 6) were collected to detect changes in microbial diversity. Illumina NovaSeq PE250 was applied for 16S amplicon sequencing to obtain raw data. Sequence data analysis mainly used Qiime 2 (Qiime2-2020.2) and R software (4.0.2). In addition, KEGG pathway analysis was performed by PICRUSt.

2.3. Enzyme Linked Immunosorbent Assay (ELISA). According to LPS (CSB-E09945h, CUSABIO, China), IL-1β (CSB-E08053h, CUSABIO, China), TNF-α (CSB-E04740h, CSA BIO, China), and IL-6 (CSB-E04638h, CUSABIO, China) ELISA instructions, we detected LPS level in feces and placental tissues and IL-6, IL-1β, and TNF-α levels in cells. 20 μg of feces and placental tissue was taken, respectively, and the blood stains were washed with 1 × PBS. Feces and placental tissue were cut into small pieces and put into tissue grinder (homogenate tube), 200 μL 1 × PBS was added to make homogenate, then put in -20°C overnight. After repeated freeze-thaw treatment two times to destroy the cell membrane, the tissue homogenate was centrifuged at 5000 g (2-8°C) for 5 min to take the supernatant, which should be detected. For cells, centrifugation was performed at 1000 g (2-8°C) for 15 min, and the supernatant should be immediately detected.

2.4. Cell Treatment. Human trophoblast cell HTR-8/SVneo was purchased from Tongpai (Shanghai, China) Biotechnology Co., Ltd., and culture conditions were as follows: RPMI-1640 medium and 10% fetal bovine serum (FBS). LPS was applied to induce cellular inflammation [15], and in addition, IncRNA BC030099 was knocked down, which are grouped as the control, si-control, si-con+LPS, si-BC030099, and si-BC030099+LPS groups. In addition, we added 10 μM NF-κB-specific inhibitor JSH-23 (Merck, Germany) [16], grouped as the control, LPS, JSH-23, and LPS+JSH-23 groups. IncRNA BC030099 expression was knocked down. si-BC030099 and negative control si-control were synthesized by Sangon Biotechnology Co., Ltd. (Shanghai, China).

2.5. Quantitative Real-Time PCR (qRT-PCR). Total RNA was extracted using Trizol method, RNA was reverse transcribed into cDNAs using a cDNA reverse transcription kit (#CW2569, Beijing ComWin Biotech, China), and fluorescence quantification was applied by Ultra SYBR Mixture (#CW2601, Beijing ComWin Biotech, China). The relative expression of genes was examined on an PCR instrument (QuantStudio1, Thermo, USA). Using GAPDH as an internal reference, the 2ΔΔCt method was performed to calculate IncRNA BC030099 level. The primers were as follows: IncRNA BC030099-F: GCCTCCATCCTTTCCAGACC and
lnRNA BC030099-R: GCCCTTGGAAAGTGTCAAG; GAPDH-F: ACACGCTCAAGATCATCAGC and GAPDH-R: GGTCATGAGTCTCCTCCAGAT.

2.6. Western Blot. Total protein was extracted from cells by RIPA lysis buffer (P0013B, Beyotime, China), followed by protein quantification, mixed with SDS-PAGE loading buffer, and protein was adsorbed on PVDF membrane by gel electrophoresis. MMP2 (10373-2-AP, 1:500, Proteintech, USA), MMP9 (ab76003, 1:5000, Abcam, UK), snail (13099-1-AP, 1:500, Proteintech, USA), and E-cadherin (20874-1-AP, 1:1000, Proteintech, USA) primary antibodies were incubated overnight at 4°C. HRP secondary antibodies were then incubated. Visualization was performed using ECL luminescent fluid (advanta, K-12045-D50, USA). β-actin was used as an internal reference to examine protein levels.

2.7. Cell Counting Kit-8 (CCK-8) Assay. Cells grouped above were digested and counted and seeded in a 96-well plate (5 x 10^3 cells/well), with 100 μL per well. After cultivating the adherent cells, we followed the above method for the corresponding time and then added 10 μL/well of CCK-8 to each well. After incubation at 37°C and 5% CO_2 for 4 h, the absorbance (450 nm) was analyzed on the Bio-Tek microplate reader (MB-530, HAEAS, China).

2.8. Transwell Assays. 1 x 10^5/mL cells were resuspended in serum-free medium, 100 μL cell suspension was added into Transwell chamber (#33318035, Corning, USA) upper, and 10% FBS was added into lower chamber and cultured for 48 h. Chamber culture medium was discarded. Cells on upper ventricle were wiped with the wet cotton swab. 4% paraformaldehyde was fixed. Cells were stained with 0.5% crystal violet and eluted with water. Cells on the outer surface of upper chamber were observed under a microscope (Olympus, Japan) and photographed. Detection of cell invasion was performed using Transwell chamber (3428, Corning, USA) with Matrigel Basement Membrane Matrix (354262, BD Biocoat, USA). The other steps were as described above.

2.9. Statistical Analysis. Statistical analysis was performed using Graphpad Prism8.0 statistical software. Measurement data are expressed as the mean ± standard deviation. An unpaired t-test was used between groups, and one-way analysis of variance was used for comparison among multiple groups. The correlation of LPS and lnRNA BC030099 in PE placental tissues was calculated by Spearman’s correlation analysis. P < 0.05 indicated a statistically significant difference.

3. Results

3.1. Gut Microbiota in PE Patients Was Changed, and Key Pathway Targets (LPS) Were Screened. First, we analyzed changes in gut microbiota by 16S rRNA sequencing and pathway analysis by PICRUSt. Figure 1(a) shows the rank abundance curve. Venn diagram showed shared and unique ASVs between groups (Figure 1(b)). Among them, there were 314 ASVs unique to the control group, 184 ASVs unique to PE patients, and 213 ASVs shared by two groups. Relative abundance histogram of the top 20 ASVs at genus level further suggested the abundance and composition of microbial communities in each group were different (Figure 1(c)). As shown in Figure 1(d), the relative abundance of the top 20 dominant gut microbiota at phylum level was shown. Furthermore, KEGG showed that microbial genes related to LPS biosynthesis were significantly elevated in gut microbiota in PE group (Figure 1(e)).

3.2. LPS Was Positively Correlated with lnRNA BC030099. Next, we examined LPS level in feces and placenta of PE patients. LPS level in the feces and placenta of PE group were significantly elevated than control group (Figure 2(a)). Furthermore, we used qRT-PCR to evaluate lnRNA BC030099 level in PE patients’ placenta. lnRNA BC030099 level in placenta of the PE group was also strikingly elevated than that of the control group (Figure 2(b)). Spearman’s correlation analysis showed there was a significant positive correlation between LPS and lnRNA BC030099 expression in placental tissue (Figure 2(c)).

3.3. Knockdown of lnRNA BC030099 Promoted Trophoblast Cell Proliferation, Invasion, and Migration. Next, we knocked down lnRNA BC030099. We found lnRNA BC030099 expression was repressed in the si-BC030099 group compared with the si-control group. This indicated that we successfully knocked down lnRNA BC030099. Compared with the si-BC030099 group, lnRNA BC030099 expression in the si-BC030099+LPS group was facilitated (Figure 3(a)). Cell function experiments revealed compared with the si-control group; the si-BC030099 group had enhanced cell proliferation, migration, and invasion abilities. However, the si-BC030099+LPS group showed reduced cell proliferation, migration, and invasion than the si-BC030099 group (Figures 3(b)–3(d)). The above results indicated knockdown of lnRNA BC030099 promoted trophoblast cell proliferation, invasion, and migration.

3.4. Knockdown of lnRNA BC030099 Elevated MMP2, MMP9, and Snail Levels and Repressed E-Cadherin Level. In addition, Western blot analysis showed that MMP2, MMP9, and snail expressions were increased in the si-BC030099 group compared with the si-control group. But the si-BC030099+LPS group showed decreased expression of MMP2, MMP9, and snail than the si-BC030099 group. The trend of E-cadherin was opposite to that of snail (Figure 4). Collectively, knockdown of lnRNA BC030099 elevated MMP2, MMP9, and snail levels and repressed ECadherin level.

3.5. lnRNA BC030099 Affected NF-κB Pathway. Next, we examined NF-κB pathway-related protein expression. Compared with the si-control group, the si-con+LPS group showed a decrease in IκBα expression and an increase in p-IκBα and p65 expressions; IκBα expression in the si-
Figure 1: Continued.
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BC030099 group was promoted, and p-IκBα and p65 expressions were inhibited. However, the si-BC030099+LPS group had repressed IκBα expression and facilitated p-IκBα and p65 expressions than the si-BC030099 group (Figure 5(a)). In addition, we examined inflammation-related marker expression. Compared with those in the si-control group,
IL-6, IL-1β, and TNF-α levels in the si-con+LPS group were facilitated; IL-6, IL-1β, and TNF-α levels in the si-BC030099 group were repressed. The si-BC030099+LPS group had elevated IL-6, IL-1β, and TNF-α levels than the si-BC030099 group (Figure 5(b)). All in all, lncRNA BC030099 affected NF-κB pathway.
3.6. NF-κB Inhibitor Reversed Trophoblast Cell Proliferation, Invasion, and Migration Induced by LPS. Finally, we used NF-κB pathway inhibitor JSH-23. IkBα expression was suppressed, and p-IκBα and p65 expressions were facilitated in LPS group than in the control group. However, compared with the LPS group, the LPS+JSH-23 group showed an increase in IkBα expression and a decrease in p-IκBα and p65 expressions (Figure 6(a)). Cell function experiments showed compared with the control group; the LPS group had reduced cell proliferation, migration, and invasion abilities. After adding JSH-23, the LPS+JSH-23 group had enhanced cell proliferation, migration, and invasion abilities.
Figure 6: Continued.
Then, we used Western blot to detect MMP2, MMP9, snail, and E-cadherin levels. MMP2, MMP9, and snail expressions were suppressed in the LPS group than the control group. After adding JSH-23, MMP2, MMP9, and snail expressions were also repressed in the LPS+JSH-23 group. The trend of E-cadherin was opposite to that of snail (Figure 6(e)). The above results indicated that NF-κB inhibitor reversed trophoblast cell proliferation, invasion, and migration induced by LPS.

4. Discussion

PE is a devastating medical complication of pregnancy that causes severe maternal and fetal morbidity and mortality [17]. PE complicates maternal and child health management and contributes to the majority of adverse pregnancy outcomes, but the mechanisms underlying PE development remain unclear. Studies have shown gut microbiota structure of PE patients has changed significantly, which may be associated with the occurrence and development of disease [18]. However, further studies are needed to understand underlying mechanisms. In this research, we sought to explore IncRNA BC030099, inflammation, and gut microbiota relationship in PE. Our research may contribute to the early diagnosis and targeted monitoring of PE.

Colonization of neonatal gut with beneficial bacteria is essential for mucosal barrier’s establishment and maintenance, thereby protecting neonate from intestinal pathogens and local and systemic inflammation [19]. Recent advances show abnormalities in microbiome composition may play a role in various disease pathogenesis including PE [20]. Alterations in gut microbiota composition could alter short-chain fatty acid profile released by bacteria and contribute to hypertension and metabolic syndrome [21]. Chen et al. found that PE patients had repressed bacterial diversity and marked dysbiosis [22]. Our study is consistent with previous studies showing changes in gut microbiota in PE patients. It was reported maternal blood IL-6 levels in PE were positively correlated with Bilophila and Oribacterium, while LPS levels were negatively correlated with Akkermansia [23]. Chang et al. found the abundance of LPS synthesis pathway was significantly elevated in predicted PE patients, while the abundance of the G protein-coupled receptor pathway was observably repressed [11]. We also found that gut microbiota related to LPS biosynthesis were significantly elevated in gut microbiota in the PE group.

IncRNAs are noncoding transcripts, typically over 200 nt in length, that have recently emerged as one of the largest and significantly diverse RNA families [24]. IncRNAs have now become important players in almost all gene functions and regulatory levels [25]. The study showed circulating IncRNA BC030099 level in plasma of PE patients was significantly higher than non-preeclamptic healthy subjects. Elevated plasma IncRNA BC030099 level was related to an

(Figures 6(b)–6(d)). Then, we used Western blot to detect MMP2, MMP9, snail, and E-cadherin levels. MMP2, MMP9, and snail expressions were suppressed in the LPS group than the control group. After adding JSH-23, MMP2, MMP9, and snail expressions were also repressed in the LPS+JSH-23 group. The trend of E-cadherin was opposite to that of snail (Figure 6(e)). The above results indicated that NF-κB inhibitor reversed trophoblast cell proliferation, invasion, and migration induced by LPS.

**Figure 6: NF-κB inhibitor reversed trophoblast cell proliferation, invasion, and migration induced by LPS.** (a) IκBα, p-IκBα, and nucleus NF-κB p65 expressions in trophoblast cells. (b) Cell activity was determined by CCK-8 assay. (c, d) Cell migration and invasion were examined by Transwell assays. (e) Western blot was performed to determine MMP2, MMP9, snail, and E-cadherin levels in trophoblast cells. Scale bar = 100 μm. *P < 0.05 vs. control; †P < 0.05 vs. LPS.
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LncRNAs have been shown to participate in different biological processes including cell growth, antiapoptosis, migration, and invasion [27]. Chen et al. reported IncRNA KCNQ1OT1 can regulate miR-146a-3p role in HTR8/SVneo cell proliferation, invasion, and migration through the CXCL12/CXCR4 pathway, which may be involved in PE pathogenesis [14]. Our study also showed knockdown of IncRNA BC030099 promoted HTR-8/SVneo cell proliferation, migration, and invasion. Studies have shown that MMPs are vital mediators of vascular and uterine remodeling and reduced MMP2 and MMP9 expressions is thought to be involved in hypertensive pregnancy and PE [28, 29]. Placental extravillous cell invasion involves cell EMT, and several EMT regulators (including snail and E-cadherin) have been found to play vital roles in PE development [30]. Knockdown of IncRNA CRNDE has been reported to inhibit the proliferation, migration, and invasion of HTR-8/SVneo cells, inhibit the formation of EMT, and reduce the protein expression of MMP2 and MMP9 [31]. Zhou et al. reported that IncRNA SNHG12 promoted trophoblast cell migration and invasion by inducing the progression of EMT [32]. Consistent with their study, we found knockdown of IncRNA BC030099 elevated MMP2, MMP9, and snail levels and repressed E-cadherin level. NF-κB is a vital transcription factor for inflammation-related protein expression [33]. The NF-κB p65 pathway is associated with PE, and inhibition of NF-κB p65 pathway can ameliorate PE [34]. NF-κB and proinflammatory cytokines have been reported to be associated with trophoblast dysfunction [35], but the underlying mechanisms remain unclear. Vaughan and Walsh reported that placental NF-κB was activated nearly 10-fold in PE. Oxidative stress leads to NF-κB activation in trophoblast-like cells, which is enhanced by TNF-α [36]. We found through cellular experiments that LPS induced the nucleus transition of NF-κB through IncRNA BC030099. Furthermore, NF-κB inhibitor reversed trophoblast cell proliferation, invasion, and migration induced by LPS. This is also the first time that we report the mechanism of action of LPS, IncRNA BC030099 and NF-κB p65 in PE. In conclusion, we explored the evolution of LPS, IncRNA BC030099, and NF-κB p65 involved in PE progression and its mechanism. We report for the first time that gut microbiota dysbiosis in PE contributed to trophoblast cell proliferation, invasion, and migration through the IncRNA BC030099/NF-κB pathway. This study provides a reference for exploring IncRNA and gut microbiota mechanism in PE.

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 Committee of Xiangya Hospital central South University. The research was conducted according to the World Medical Association Declaration of Helsinki. All the information about the study will be fully explained to the subjects by the researchers.

Consent

All the participants provided informed consent before sampling.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this paper.

Authors’ Contributions

Rong Tang, Gong Xiao, Qiongjing Yuan, and Wei Wang contributed to the conception and design of the study. Rong Tang and Chun Jiang collected the information. Gong Xiao, Qiongjing Yuan, Chun Jiang, and Wei Wang wrote sections of the manuscript. All authors have revised, read, and approved the final version of the manuscript.

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