ORIGINAL RESEARCH

Gut dysbiosis induces the development of pre-eclampsia through bacterial translocation

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ABSTRACT

Objective  Pre-eclampsia (PE) is one of the malignant metabolic diseases that complicate pregnancy. Gut dysbiosis has been identified for causing metabolic diseases, but the role of gut microbiome in the pathogenesis of PE remains unknown.

Design  We performed a case–control study to compare the faecal microbiome of PE and normotensive pregnant women by 16S ribosomal RNA (rRNA) sequencing. To address the causative relationship between gut dysbiosis and PE, we used faecal microbiota transplantation (FMT) in an antibiotic-treated mouse model. Finally, we determined the microbiome translocation and immune responses in human and mouse placental samples by 16S rRNA sequencing, quantitative PCR and in situ hybridisation.

Results  Patients with PE showed reduced bacterial diversity with obvious dysbiosis. Opportunistic pathogens, particularly Fusobacterium and Veillonella, were enriched, whereas beneficial bacteria, including Faecalibacterium and Akkermansia, were markedly depleted in the PE group. The abundances of these discriminative bacteria were correlated with blood pressure (BP), proteinuria, aminoacids transferase and creatinine levels. On successful colonisation, the gut microbiome from patients with PE triggered a dramatic, increased pregestational BP of recipient mice, which further increased after gestation. In addition, the PE-transplanted group showed increased proteinuria, embryonic resorption and lower fetal and placental weights. Their T regulatory/helper-17 balance in the small intestine and spleen was disturbed with more severe intestinal leakage. In the placenta of both patients with PE and PE-FMT mice, the total bacteria, Fusobacterium, and inflammatory cytokine levels were significantly increased.

Conclusions  This study suggests that the gut microbiome of patients with PE is dysbiotic and contributes to disease pathogenesis.

INTRODUCTION

Pre-eclampsia (PE) is a disorder of pregnancy associated with new-onset hypertension, which occurs most often after 20 weeks of gestation combined with proteinuria. PE affects 2%–8% of pregnancies and is associated with an increased risk of adverse pregnancy outcomes, including placental abruption, prematurity and intrauterine growth restriction. Pre-eclampsia (PE) is one of the malignant metabolic diseases that complicate pregnancy. Gut dysbiosis has been identified for causing metabolic diseases, but the role of gut microbiome in the pathogenesis of PE remains unknown.

Significance of this study

What is already known on this subject?

Pre-eclampsia (PE) is one of the leading causes of maternal and perinatal morbidity and mortality and lacks reliable prediction methods and detailed pathogenesis knowledge.

What are the new findings?

Inoculation of the gut microbiome from patients with PE triggered a significantly higher blood pressure before pregnancy and PE-like phenotypes during pregnancy, including worsened hypertension, exacerbation of proteinuria and intrauterine growth restriction. Higher levels of total bacteria and Fusobacterium and an altered microbiota profile were identified in PE placenta samples compared with normotensive samples.

How might it impact on clinical practice in the foreseeable future?

Gut microbiota screening shows potential for predicting PE onset and provides a new approach for the prevention and treatment of this disease.
The presence of glucose intolerance and dyslipidaemia, along with hypertension, can contribute to endothelial dysfunction, poor placentation and abnormal placental development, which are common pathological abnormalities in PE.7

Numerous studies have suggested that gut dysbiosis plays a causative role in the development of metabolic syndrome, chronic inflammation diseases and complicated pregnancies.8 9 Gut microbiota can facilitate insulin resistance by inducing the chronic inflammation in the host and causing fat accumulation by regulating genes related to energy metabolism, which plays an important role in the development of core symptoms in metabolic disorders.10 Gut microbiome dysbiosis is a key factor in blood pressure (BP) regulation11 12 and can contribute to the development of proteinuria and subsequently lead to kidney disease.13

The composition of the gut microbiota in patients with PE and the relationship between the gut microbiota and PE have not been clarified so far. Recent studies suggested a disrupted gut microbiota composition in patients with PE in late pregnancy,14 15 but there is lack of causative analysis. In this study, we performed 16S ribosomal RNA (rRNA) gene sequencing on faecal samples to investigate the detailed profile of the gut microbiome and to determine the crucial role of the disordered gut microbiome in triggering PE-like phenotypes by faecal microbiota transplantation (FMT).

MATERIALS AND METHODS

Study subjects

Women with PE and normotensive pregnant (NP) women without previous treatments were recruited in the third trimester from March 2017 to March 2018 in the Department of Obstetrics of the Nanfang Hospital, Southern Medical University, China. According to the current American College of Obstetricians and Gynaecologists criteria,2 PE was defined as hypertension (systolic blood pressure (SBP) or diastolic blood pressure (DBP) of ≥140 or ≥90 mm Hg on two occasions at least 4 hours apart) and significant proteinuria (≥300 mg protein per 24 hours’ urine collection or protein/creatinine ratio of ≥0.3 mg/dL or dipstick reading of 2+ in a random urine sample) after 20 weeks of gestation in a woman with a previously normal BP. In the absence of proteinuria, PE was defined when hypertension was associated with any features of end organ damage. PE with severe features was considered when either severe hypertension (SBP or DBP ≥160 or ≥110 mm Hg), thrombocytopenia, impaired liver function, renal insufficiency pulmonary oedema, headache or visual disturbance was present. The clinical characteristics of the faecal and placenta donors are summarised in online supplementary tables S1 and S2, respectively.

Animals and experimental protocol

C57BL/6 mice were housed with a 12 hours’ light–dark cycle in specific pathogen-free facilities. FMT was performed according to a modified method described previously.16 Briefly, female mice aged 6–8 weeks were administered antibiotics (vancomycin, 100 mg/kg; neomycin sulfate, metronidazole and ampicillin, 200 mg/kg) intragastrically once daily for five consecutive days. We randomly selected three patients with PE and three NP controls as faecal donors for FMT (online supplementary table S3). The faecal supernatants of three PE and three NP donors were mixed and used as a single source for the PE-FMT and NP-FMT mice, respectively. Following antibiotic treatment, the recipient mice were orally inoculated daily for three consecutive days and two times per week thereafter for 59 days with prepared faecal contents. Another group of female mice was gavaged as described previously, but the donor supernatant was replaced with phosphate-buffered saline (PBS). Six weeks after the first microbial administration, all female mice were housed with male mice at a 2–3:1 ratio overnight, and pregnancy was confirmed by the presence of vaginal spermatozoa. Mouse harbouring microbiota from patients with PE and NP, as well as PBS, were referred to as PE-FMT, NP-FMT and control, respectively. The group and cage number of the mice involved in the study are shown in online supplementary table S4.

Protein expression and biochemical analysis

Protein was extracted from tissues with commercial lysis buffer. Western blotting was performed as described previously.17 Primary antibodies against zonula occluden (ZO)-1 (Invitrogen, Carlsbad, California, USA), ZO-2 (Abcam, Cambridge, UK), occludin (Invitrogen), claudin-4 (Abcam) and glyceraldehyde 3-phosphate dehydrogenase (GAPDH, Invitrogen) were used. Faecal albumin and urinary protein concentrations were determined with a mouse albumin ELISA kit (Bethyl Labs, Montgomery, Alabama, USA) according to the manufacturer’s instructions. The endotoxin level was measured with a commercial kit (Bio-Swamp, Beijing, China).

Gene expression analysis

Total RNA was extracted with Trizol reagent (Invitrogen) according to the manufacturer’s instructions. The reverse transcription reaction was carried out with reverse transcription enzyme (Toyobo, Shanghai, China). Quantitative real-time PCR was carried out on an ABI Q5 real-time PCR system and the specific primers for quantitative PCR are shown in online supplementary table S5.

Flow cytometry

Isolated immune cells were resuspended and stained with anti-CD4-FITC and/or anti-CD25-APC monoclonal antibodies (mAbs) for surface antigens and anti-Foxp3-PE or anti-IL-17A-PE mAbs for intracellular antigens in accordance with the manufacturer’s instructions (eBioscience, San Diego, California, USA). Fluorescence minus one controls were stained in parallel. Flow cytometry data acquisition was conducted using a BD LSRFortessa flow cytometer (BD Biosciences, San Jose, California, USA), and the analysis was performed by BD FACSDiva Software V8.0.1 (BD Biosciences). The gating strategies are shown in online supplementary figure S1.

In situ hybridisation

In situ hybridisation (ISH) was performed following a modified protocol described by Choi et al.18 19 Briefly, paraffin-embedded sections were subjected to deparaffinisation, rehydration and permeabilisation. The probes used in this study were as follows: Eubacterial probes (EUB338 1-II) mixture complementary to a 16S rRNA region specific for most bacteria; NON338, an oligonucleotide complementary to the probe EUB338, serving as a scrambled control for nonspecific binding; Fusobacteria-specific probe, a genus-specific probe complementary to a 16S rRNA region of Fusobacterium. All probes were 5’-labelled with digoxigenin, and antidigoxigenin/horseradish peroxidase antibodies were used as secondary antibodies. Tyramide signal amplification was performed to increase the detection sensitivity of bacterial signals. Six areas of each section were randomly photographed under a fluorescence microscope. The specific probes for ISH are shown in online supplementary table S6.
Bioinformatics processing
Using QIIME V.1.9.1, we merged, applied quality control and clustered the 16S rRNA gene reads into operational taxonomic units (OTUs). Taxonomic groups were based on the Greengenes Database V.13.8 using closed reference to perform reference-based OTU clustering. \cite{26, 27} Values for alpha diversity (Chao1 Index, Shannon’s Index, Phylogenetic Diversity (PD) Whole Tree Index and observed OTUs), beta diversity (unweighted UniFrac distance metrics) and principal coordinate analysis (PCoA) employed based on the UniFrac metrics were generated by QIIME V.1.9.1. Permutational multivariate analysis of variance was performed to determine if the compositions of microbiota differed between groups. Linear discriminant analysis effect size (LEfSe) was performed to determine the features most likely to explain the differences between groups.\cite{28} To identify and filter the potential contami- nant OTUs, decontam V.1.4.0\cite{29} and SourceTracker V.1.0.1\cite{30} were applied to the placenta samples and negative controls.

Statistical analysis
According to the different data, statistical analysis between groups were analysed using the Wilcoxon’s rank-sum test, Student’s \( t \)-test or one-way analysis of variance. Correlation analyses were performed based on the Spearman’s rho statistic. Rate com- parisons were performed with Pearson’s \( r^2 \) test or Fisher’s exact test. Multiple hypotheses were adjusted using the Benjamini and Hoch- berg method.\cite{31} In the figures, \( p<0.05 \) indicates statistical signifi- cance (\( \star p<0.05, \star\star p<0.01, \star\star\star p<0.001 \)). Statistical analyses and data visualisation were performed using R V.3.5.0 (under RStudio V.1.1.453), using the ggplot2 and corrplot packages.

Other materials and methods are described in the online supplementary materials and methods.

RESULTS
Dysbiosis of gut microbiota in patients with PE
A total of 85 normotensive women and 67 patients with PE were included in the present study. In the PE group, there were 40 patients with severe features (PESF) and 27 patients without severe features (PETSF). The PE group was further divided into an early onset of PE group (21 women) and a late onset of PE group (46 women) according to whether the disease was detected before 33 gestational weeks.

According to PCoA, the gut microbiome of the PE group differed significantly from that of the NP group using the unweighted UniFrac distance (Figure 1A). Alpha-diversity indices, including Chao1 and observed OTU, were markedly decreased in the PE group (Figure 1B,C and online supplementary figure S2A,B). We did not observe significant differences between the PE with and without severe features subgroups (online supplementary figure S3A–E) or between the early and late onset of PE subgroups (online supplementary figure S4A–E) for both alpha diversity and beta diversity.

The intestinal bacterial community consisted of 379 genera belonging to 18 phyla in patients with PE and in normotensive individuals. At both the phylum and genus levels, the micro- bial composition of patients with PE differed from that of the normotensive group (online supplementary figure S2C,D). To further identify which bacterial taxa were distinct between the PE and NP groups, we performed LEfSe analysis and identified seven genera showing significant differences (Figure 1D). Moreover, the abundance comparisons of predominant genera showed that Clostridium, Dialister, Veillonella and Fusobacte- rium were significantly enriched, whereas Lachnospira, Akkermansia and Faecalibacterium were depleted in patients with PE (Figure 1E–K). However, no significant differences of the above-mentioned genera were observed among the PESF and PETSF subgroups (online supplementary figure S3F).

Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt) analysis was performed to infer microbial metabolites and metabolic pathways. Pathways correlated with inflammation, including lipopolysaccharide biosynthesis and bacterial toxins, were enriched in patients with PE (online supplementary figure S2E,F). These key genera showed significant correlations with BP, proteinuria, impaired hepatic func- tions or other clinical manifestations based on Spearman correla- tion calculated for all participants (Figure 1L). Notably, Veillonella and Fusobacterium were significantly correlated with most clinical parameters, including SBP, DBP, proteinuria, oedema levels, alanine aminotransferase, aspartate aminotransferase, serum creati- nine and albumin. The correlation coefficients for Lachnospira, Akkermansia and Faecalibacterium were just opposite. However, no bacteria were significantly correlated with clinical parameters when testing only on the PE group, which could be due to reduced sample size. These results reveal profound changes in the intestinal microbiome structure of the PE group, indicating the importance of gut microbes in the development of PE.

FMT induces the pre-eclamptic phenotype
To further test whether alterations in the gut microbiota can cause the progression of PE in vivo, faecal microbiome from patients with PE were transplanted into mice. Following antibiotics treatment, the recipient mice were randomly divided into groups and orally inoculated with prepared faecal contents from donors with PE or NP (Figure 2A). The gut microbial profiles of recipient mice were analysed by 16S RNA sequencing at 6 weeks post-transplantation. As expected, PCoA based on unweighted UniFrac distance matrices showed a significant effect in the FMT group on the mouse faecal microbiome (online supplementary figure S5A,B). From the donors, 71 of 128 genera successfully colonised the intestine of recipient mice (online supplementary figure S5C). Additionally, 55% of the shared genera were found to vary with the same trend across the microbiota of donors and recipient mice (online supplementary figure S5D,E). Moreover, at the genus level, the abundance variations in Coprococcus, Fusobacterium, Megamonas, Clostridium, Phascolarctobacterium and Akkermansia showed the same trend between mice and donors (online supplementary figure S5F). These results suggest that the faecal microbiota composition of human donors was successful- ly reproduced in recipient mice.

At 6 weeks post-transplantation, PE-FMT mice exhibited significantly higher pregestational SBP compared with NP-FMT and control mice. After pregnancy, the SBP of the PE-FMT group was further elevated from the fifth gestational day to the end of gestation (Figure 2B and online supplementary figure S6A). In addition to worsening hypertension, the PE-FMT group showed a significantly higher urine protein concentration at 17 days of gestation, whereas urinary protein excretion was unchanged at 6 weeks post-FMT prior to mating (Figure 2C). The number of live pups, fetal weight and placental weight were also decreased in the PE-FMT group, whereas the fetal resorption rate was increased compared with that in the NP-FMT and control groups (Figure 2D,E). To determine how faecal bacteria from patients with PE affect the placenta, we examined the structure of the implantation sites. Histomorphological analysis revealed a significant structural change in the PE-FMT placenta, as the ratio between the two functional placental zones (labyrinth and junctional zones) was increased in placentas from the PE-FMT.
Figure 1  Dysbiosis of gut microbiota patterns in patients with PE and NP women. (A) Principal coordinate analysis based on unweighted UniFrac distances revealed that the NP bacterial communities clustered separately from PE bacterial communities, which were more similar to each other. Each circle represents a single sample, colour by group. The eigenvalues of axe Principal coordinate (PC)1 and PC2 were 1.27 (6.56%) and 0.73 (3.53%), respectively. Difference in beta diversity between NP and PE were tested by permutational multivariate analysis of variance (Adonis). (B,C) Comparison of alpha-diversity indices (Chao1 Index and observed operational taxonomic units) between PE and NP groups. ** indicates p<0.01 by Wilcoxon rank-sum test. (D) linear discriminant analysis effect size identified the most differentially abundant taxa between the two groups. PE-enriched taxa are indicated with a positive LD score, and taxa enriched in NP controls have a negative score. Only taxa meeting an LDA significant threshold of >3 are shown. (E–K) Relative abundances of Clostridium, Dialister, Veillonella, Fusobacterium, Lactobacillus, Akkermansia and Faecalibacterium between PE and NP groups. * indicates p<0.05; ** indicates p<0.01; and *** indicates p<0.001 by Wilcoxon rank-sum test. (L) Correlations between the relative abundance of selected bacterial genera and clinical parameters. Correlations with p<0.05 are shown. Spearman’s rho statistic adjusted using the Benjamini and Hochberg method. AT-III, antithrombin III; ALB, albumin; ALT, alanine aminotransferase; AST, aspartate aminotransferase; Cr, creatinine; DBP, diastolic blood pressure; LDA, linear discriminant analysis; NP, normotensive pregnant; PE, pre-eclampsia; SBP, systolic blood pressure; TP, total protein; TT, thrombin time.

FMT induces immunological imbalances

Decreased regulatory T (Treg) cells and/or increased T helper-17 (Th17) cells play a central role in the pathogenesis of PE.26,27 Thus, we investigated whether the adverse effect observed with FMT was associated with changes in Treg/Th17 proportions. We monitored changes in Treg/Th17 proportions. We observed a significant reduction in Treg cells and an increase in Th17 cells in PE-FMT mice compared with those in NP-FMT or control mice (figure 3A–D). Moreover, the Treg/Th17 cell ratios were lower in PE-FMT mice than in NP-FMT and control mice (figure 3E). We further examined inflammatory gene expression in the ileum to detect alterations in immunological responses. As shown in figure 3F, PE-FMT mice exhibited significantly higher Il1β, Ccl3, Cxcl1 and Vegf levels than NP-FMT or control mice. Collectively, these data indicate that the faecal microbiota from patients with PE disrupted the mucosal and systemic immunological response.

FMT in patients with PE microbiota impairs intestinal barrier function

Impaired immunological responses in the intestine are known to be associated with gut barrier dysfunction.28 We investigated if faecal bacteria from patients with PE affected gut barrier function. Intestinal permeability was monitored by faecal albumin content, which was significantly increased in the PE-FMT group.
Figure 2  Administration of faecal bacteria from patients with PE to mice induced a series of PE-like phenotypes. (A) Schematic representation of FMT. Fresh faecal from three PE and three NP donors were mixed and used as a single source for PE-FMT and NP-FMT mice, respectively. Following antibiotic treatment, the recipient mice were randomly divided and orally inoculated daily for three consecutive days and two times per week for 59 days with prepared faecal contents. (B) SBP of the three groups was evaluated pregestation and postgestation (n=8 for control and PE-FMT, n=10 for NP-FMT). α indicates p<0.01 versus NP-FMT group on the corresponding day; β indicates p<0.01 versus control group on the corresponding day. θ indicates p<0.05 for each corresponding gestational day versus the pregestational day (42 days post-FMT) by two-tailed Student’s t-test. (C) Urine protein concentration was measured at 42 days (prior to mating) and 62 days (17 days of gestation) post-FMT (n=8 for control and PE-FMT, n=10 for NP-FMT). (D) Number of pups and resorption rate per litter (n=8 for control and PE-FMT, n=10 for NP-FMT). (E) Placental weight and fetal weight significantly decreased in PE-FMT (n=41) compared with in NP-FMT (n=76) and controls (n=59) at 17 days of gestation. (F) Representative images of H&E-stained midsagittal placental tissue sections used in histomorphological analysis. Zones are marked and indicated by labyrinth zone and junctional zone. The ratio between the labyrinth/junctional zones was enhanced in PE-FMT (n=14) compared with in NP-FMT (n=16) and controls (n=16). (G) Representative H&E staining of the labyrinth zone of recipient mouse placentas. Typical lesions are marked with dotted line (original magnification, ×400; scale bar=50 µm). Data are presented as the mean±SEM (B–F). *P<0.05, **P<0.01, ***P<0.001 by one-way analysis of variance following Duncan’s multiple range test. ABX, antibiotics; BP, blood pressure; FMT, faecal microbiota transplantation; NP, normotensive pregnant; PE, pre-eclampsia; SBP, systolic blood pressure.

(Figure 4A). FMT markedly increased plasma endotoxin levels in PE-FMT mice (Figure 4B). In addition, faecal microbiota from patients with PE decreased Claudin4 and Occludin gene expression in the colon compared with that in NP-FMT or control mice (Figure 4C). Similarly, barrier-forming tight-junction protein ZO-1 and ZO-2, claudin-4 and occludin protein levels in the colon were markedly lower according to immunohistochemistry and western blot analyses (Figure 4D–H). We next examined the effect of FMT on intestinal histopathological changes. There was no obvious difference in the morphology of the epithelium in

Figure 3  FMT induced immunological imbalance. Treg and Th17 cells were analysed for coexpression of CD4+CD25+ Foxp3+ and CD4+IL-17A+ by flow cytometry, respectively. The number of Treg and Th17 cells was expressed as a percentage of CD4+ T cells for each subject and compared among different groups. (A–D) Percentages of Th17 cells in siLPL and spleen, and Treg cells in siLPL and spleen were determined (n=8 mice in each group). (E) Treg/Th17 cell ratios in siLPL and spleen of the three groups are shown (n=8 mice in each group). (F) Messenger RNA levels of key cytokines and chemokines in the ileum (n=7–8). Data are presented as the mean±SEM (A–F); *P<0.05, **P<0.01, ***P<0.001 by one way analysis of variance following Duncan’s multiple range test. FMT, faecal microbiota transplantation; NP, normotensive pregnant; PE, pre-eclampsia; siLPL, small intestinal lamina propria lymphocyte; Th17, helper 17 T; Treg, regulatory T.

Bacterial translocation is associated with inflammation in the placenta

Increased gut permeability can lead to gut bacteria or bacterial component translocation into the extraintestinal tissues.29 30 We next detected bacteria in human and mouse placentas by quantitative PCR to evaluate the bacterial 16S rRNA gene. The levels of total bacterial DNA were obviously increased in the PE and PE-FMT placentas (figure 5A,C). To confirm these results, we performed ISH using the probe EUB338 targeting bacterial 16S rRNA. Probe specificity was also tested using sections treated with a sequence scrambled probe. More bacterial signals were visualised in the villous tissue of PE placenta and junction layer of PE-FMT placenta. No staining for scrambled probe controls was observed but was detected in the EUB338-probed sections (figure 5B,D and online supplementary figures S8 and S9). As shown in online supplementary figure S10A,B, the detection rate the ileum and colon among the three groups, except for mild infiltration of inflammatory cells in the colon of PE-FMT mice (online supplementary figure S7). Furthermore, significant correlations between the abundance of the gut microbiome, gut barrier and immune index were observed (figure 4I). These data demonstrate that the intestinal barrier was disrupted by faecal bacteria from patients with PE.
of *Fusobacterium* DNA was higher in PE placental samples than in NP samples according to quantitative PCR and ISH probing for the *Fusobacterium* 16S rRNA gene.

During experimental processing, we monitored contamination by adding DNA-free water during extraction and amplification (DNA extraction from blank controls and no-template amplification controls), as well as a sterile swab during placental sampling (sampling blank controls). As shown in figure 5E, placental metagenomes from human and mouse samples were significantly distinct in their clustering from negative controls. To exclude the possibility that the placental microbiota occurrence was due to contamination by bacteria in the environment, 16S metagenomics decontamination analysis was rigorously performed using the R package decontam and SourceTracker (online supplementary figure S8C,D). According to the no-contaminant filtered data, PCoA revealed the significant separation of two clusters of PE and normotension groups and NP-FMT and PE-FMT groups (figure 5F,G). LEfSe analysis identified different taxa between the PE and NP groups, showing that *Pseudomonas* and Enterobacteriaceae were enriched, whereas Ruminococcaceae and Flavobacteriia were depleted in patients with PE (online supplementary figure S10E,F). These data suggest that a low but consistently detectable biomass of bacteria DNA present in the placenta is taxonomically distinct from potential environmental negative controls (online supplementary figure S10G).
Proinflammatory cytokine and chemokines are thought to play key roles in the pathophysiology of PE.31,32 In this study, we examined alterations in chemokine and proinflammatory cytokine expression in the placenta. Particularly, the levels of IL6, IL1β, CCL3 or CCL4 messenger RNA were significantly increased in the PE group compared with those in the NP group for both human and mouse placentas (Figure 5H, I).

**DISCUSSION**

PE is one of the most severe diseases during pregnancy that not only leads to pregnancy abortion but also is life threatening.3 Effective methods for the prediction, prevention and treatment of this disease are lacking because its aetiology remains unclear.3 The present study aimed to seek these answers from the gut. We found that the gut microbiome was dramatically shifted in patients with PE. Thus, using FMT, we proved that the gut dysbiosis can cause pre-eclamptic systems. One possible mechanism through which this happens might be gut barrier destruction and the translocation of pathogenic bacteria from the gut to the placenta, causing abnormal immune responses, as we identified higher levels of total bacteria and *Fusobacterium* in the placenta of both human and mice with PE. Our study proposed that a ‘gut–placenta’ axis could play a crucial role in understanding the aetiology of PE.

The gut dysbiosis in PE observed in our study was obvious and echoes the findings of previous reports. Because the human gut microbiome is highly heterogeneous among individuals, microbiomes in healthy and diseased individuals are usually overlapped with partial differentiation when analysed with unsupervised learning.7 Nevertheless, in this study, the gut microbiome in PE clearly deviated from that in the controls and showed very little overlap, indicating severe dysbiosis in PE. More interesting...
were the dysbiosis patterns, such that Faecalibacterium and Akkermansia were depleted, whereas Fusobacterium and Veillonella were enriched in PE. Similar to the microbiota profile of non-pregnant hypertensive patients,11 Faecalibacterium and Akkermansia, which are involved in producing intestinal epithelial nutrition, maintaining immune homeostasis and strengthening intestinal barrier functions, were depleted in patients with PE.13 34 Fusobacterium and Veillonella are mainly opportunistic pathogens and are associated with chronic inflammatory conditions and adverse pregnancy outcomes.35–37 Moreover, Liu et al found that these bacteria were changed in the intestine of patients with PE in America and South China, indicating that the dysbiosis patterns for PE are consistent across populations and regions.14 15 38 Thus, the dysbiosis pattern in PE reported by our research is reliable.

Fusobacterium could be the key regulator of the gut–placenta axis during PE. The enrichment of Fusobacterium in PE is especially worth noting because it is a newly identified key pathogen involved in promoting colorectal cancer.39 The mechanism involved is that Fusobacterium can affect apoptosis, cellular proliferation and DNA repair through its attachment to epithelial cells via Fusobacterium adhesion A and its activation of the immune response via the nuclear factor-κB signaling pathways.40 In other words, Fusobacterium is especially good at disrupting intestinal epithelia leading to gut leakage.41 A possible consequence of a leaky gut is bacterial translocation, as we have identified more total bacteria and the host. Therefore, it is not precisely clear whether the dysbiosis occurs due to hormone changes or diet shifting needs to be investigated in the future.

Excessive maternal inflammation, commonly involved in PE pathogenesis by an imbalance of Treg/Th17, is also a putative result of gut dysbiosis.26 47 In the current study, PE-FMT mice exhibited a mucosal and systemic imbalance of Treg and Th17 cells and exaggerated inflammation in the placenta. This increased the inflammatory cytokines, leading to placental oxidative stress and vascular dysfunction, resulting in rejection of the fetus and the development of hypertension.22 We found that gut dysbiosis in PE can elevate the host’s BP, which is a major manifestation of PE. It has been reported that the gut microbiome is altered on entry into pregnancy48; therefore, we deduce that in pregnant hypertensive women,11 dysbiosis during pregnancy; for example, whether the dysbiosis occurs due to hormone changes or diet shifting needs to be investigated in the future.

The present study has several limitations. First, for the mouse FMT experiment, we used whole faecal materials that included bacteria, fungi, virus, phages, proteins and metabolites from both bacteria and the host. Therefore, it is not precisely clear which component from the faeces functions in PE. It is, thus, important to use multomics techniques, inactivated faeces and isolated bacteria/metabolites to affirm the key factors contributing to PE. Second, even though we have used three methods, including sequencing, ISH and quantitative PCR, and identified bacteria in the placenta, these methods do not indicate the liveness of the detected bacteria. Future studies that isolate live bacteria from placenta can certainly add credibility to our findings. Lastly, even though we have detected the enrichment of Fusobacterium in the gut and placenta and hypothesised its translocation from the gut to the placenta, the exact route through which Fusobacterium completes its translocation also needs further investigation.

Overall, we identified here obvious gut dysbiosis in patients with PE and proved that this dysbiosis can cause pre-eclamptic systems. The possible underlying mechanisms is that dysbiosis influences the host’s BP and that gut Fusobacterium translocates into the placenta, causing local placental inflammation. Cohort studies that follow up women from early pregnancy to observe their microbiome dynamics, BP changes and PE occurrence will offer critical evidence linking the gut microbiome to PE. In conclusion, our study indicates the role of the gut–placenta axis in PE, providing new insights into PE aetiology.

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Funding This investigation was financially supported by the National Natural Science Foundation of China (81925026, 81730708, 81771609, 81800746 and 31570497), the National Key Research and Development Program of China (2018YFA0802300), the Natural Science Foundation of Guangdong Province (2017A010105025) and China Postdoctoral Science Foundation (2018M630377).

Competing interests None declared.

Patient consent for publication Not required.

Ethics approval Nanfang Hospital.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available in public, open access repository.

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