Whole-transcriptome sequencing uncovers core regulatory modules and gene signatures of human fetal growth restriction

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Abstract

Background: Fetal growth restriction (FGR) contributes the primary cause of perinatal mortality and morbidity with impacts on the long-term health. To determine the core gene expression network and gene signatures, which in combination with ultrasound confirmation will more effectively differentiate constitutionally normal small for gestational age and pathological FGR groups, we performed RNA sequencing for protein-coding genes, IncRNAs, and small RNAs in a case–control study of umbilical cord blood.

Results: Five pairs of FGR case and control umbilical cord blood samples were used for RNA sequencing and weighted gene co-expression network analysis (WGCNA). Results showed that 339 mRNAs, 295 lncRNAs, and 13 miRNAs were significantly differentially expressed between FGR cases and controls. Bioinformatics analysis indicated that these differentially expressed molecules were mainly involved in metabolism, neural, cardiac, and immune systems, and identified 18 WGCNA modules for FGR. Further quantitative verification was performed using umbilical cord blood and maternal peripheral blood from 12 pairs of FGR cases and controls. The logistic regression and receiver operating characteristic curve indicated that RP11_552M6.1, LINC01291, and Asgr1 in umbilical cord blood, while Sfrp2, miR-432-5p, and miR-1306-3p in maternal peripheral blood had potential significance for FGR.

Conclusions: We comprehensively profiled the whole-transcriptome landscape of human umbilical cord blood with FGR, constructed the core WGCNA modules, and delineated the critical gene signatures of FGR. These findings provide key insight into intrauterine perturbations and candidate signatures for FGR.

Keywords: Fetal growth restriction, Case–control study, Whole-transcriptome, IncRNAs, miRNAs, Gene co-expression network
we comprehensively profiled the transcriptome-wide and prevention at an early time-point. In this study, and provide candidate approaches for disease interven-
small for gestational age and pathological FGR groups, more effectively differentiate constitutionally normal
umbilical cord blood, the ultrasound confirmation could
dation network obtained from a case–control study of
dating fetal growth-related processes.

Recent developments in high-throughput sequenc-
ing enable the assessment of the entire transcriptome of
mRNAs, long noncoding RNAs (lncRNAs), and micro-
RNAs (miRNAs) with the potential to uncover the bi-
ological processes driving complex phenotypes. Systems
biology methods can better capture the complexity of
inter-gene relationships and the signaling pathways asso-
ciated with diseases, and offer the opportunity to better
define the co-regulatory patterns that underlie complex
phenotypes. Weighted gene co-expression network anal-
ysis (WGCNA) has been successfully applied in several
studies to facilitate the systems-level characterization of
expression pattern by clustering highly-correlated genes
into co-expression modules with conserved biological
functions [10, 11].

Studies assessing the transcriptome-wide profile of
human placentas with FGR are beginning to emerge, and
a few protein-coding genes and noncoding RNAs have
been assessed [12, 13]. However, the inconsistent find-
ings suggest that placental biomarkers have a low reli-
ability, limiting their clinical ability [14]. FGR, especially
abnormal fetal growth confirmed by repeated ultrasound,
has a multifactorial nature with many causes, including
maternal, fetal, and placental factors [15]. Birth weight
is correlated with the maternal body mass index (BMI)
and delivery gestational age [16]. Moreover, information
from previous studies is limited by sample heterogeneity,
such as placental differences, and a focus on univariate
gene expression analyses contrasting normal and adverse
phenotypic outcomes. Thus, a case–control study with a
matching control to each FGR infant, could eliminate the
confounding factors and provide more persuasive evi-
dences. Furthermore, high-throughput sequencing for
the entire transcriptome of umbilical cord blood as the
origin of FGR infant could provide an exemplary oppor-
tunity to demonstrate the core gene networks by eluci-
dating fetal growth-related processes.

Therefore, in combination with the core gene expres-
sion network obtained from a case–control study of
umbilical cord blood, the ultrasound confirmation could
more effectively differentiate constitutionally normal
small for gestational age and pathological FGR groups,
and provide candidate approaches for disease interven-
tion and prevention at an early time-point. In this study,
we comprehensively profiled the transcriptome-wide
landscape of human umbilical cord blood in a case–con-
control study by implementing a network-based approach to
construct the core gene co-expression network and deline-
ate the critical gene signature of FGR.

Methods
Study participants
Using a case–control study, FGR cases and correspond-
ning controls were matched according to gestational age,
maternal BMI and age (Additional file 1: Table S1) to
exclude the maternal factors. The inclusion criteria of
FGR were based on the birth weight reference percentiles
as an estimated weight below the 10th percentile for gestational age [17]. These FGR and control infants
had a birth score not less than 9 and no birth defects. The
women had no smoking history and no other preg-
nancy complication, and the women with preeclampsia
and other complications of pregnancy were excluded. Five pairs of FGR cases and controls obtained at Shang-
hai First Maternity and Infant Hospital (Tongji Uni-
versity, Shanghai, China) between 2017 and 2018 were
used for RNA sequencing. Further quantitative RT-PCR
verification was performed in the umbilical cord blood
and maternal peripheral blood obtained from the 12
FGR cases and 12 controls (Additional file 2: Table S2).
All women provided written informed consent, and the
study protocol was approved by the Ethics Committee
of Shanghai First Maternity and Infant Hospital (No.
KS17115).

Sample collection
In total, 2.5 mL of umbilical cord blood and 2.5 mL of
maternal peripheral blood were collected at the time
of delivery into PAXgene whole blood RNA tubes (Pre-
Analytix) and stored at 25 °C for at least 2 h, at −20 °C
for 24 h, and at −80 °C until processing. The total RNA
was extracted using a RNeasy Protect Animal Blood Kit
(Qiagen) according to the manufacturer’s instructions.
The RNA concentration and purity were measured using
a NanoDrop ND100 spectrophotometer (Thermo Scien-
tific) and BioAnalyzer 2100 system (Agilent).

RNA-sequencing workflow
For lncRNAs and mRNA, the RNA-sequencing library
generation, workflow, and data analysis were performed
as previously described [18]. The small RNA sequencing
including miRNAs was also performed. After the auto-
matic quality control and adapter trimming by Trim_-
Galore (http://www.bioinformatics.babraham.ac.uk/
projects/trim_galore/), the RNA paired-end reads were
mapped to the human genome hg38 by HISAT2 [19],
and quality controlled using RSeQC [20]. Based on the
annotation file (Homo_sapiens.GRCh38.83.gtf) in the
Ensembl database, the read counts were calculated by featureCounts [21], and normalized by rlogTransformation. The differentially expressed genes (DEGs) were analyzed by DESeq 2 package in R [22].

Bioinformatics and stability analyses
The lncRNA targets were predicted with LncTar, RNAplex, and Rsearch, while the miRNA targets were predicted with miRanda, PITA, and RNAhybrid. The gene ontology (GO) and signaling pathway enrichment analyses were completed using DAVID web servers [23]. Based on the log2(Fold Change), we performed the gene set enrichment analysis (GSEA) for the significant signaling pathways and imprinted genes [24–26]. The gene co-expression network was generated using the WGCNA package in R [11]. The parameters were as follows: networkType = unsigned, corType = Pearson, Power = 9, minModuleSize = 50, mergeCutHeight = 0.35, reassignThreshold = 0.99 and the remaining default parameters. The regulatory network was illustrated by the igraph package in R, and a circos graph was obtained by the RCircos package [27]. The logistic regression analysis and receiver operating characteristic (ROC) curve were performed by glm and pROC packages, and 95% CI of area under the curve (AUC) were calculated by boot package in R. The stability of gene modules was also assessed [28].

Quantitative RT-PCR analyses
The first-strand cDNA was synthesized by M-MLV Reverse Transcriptase (TaKaRa) with 500 ng total RNA. The qRT-PCR assays were performed with SYBR® Premix Ex Taq™ II (TaKaRa) on an Mx3000P QPCR System (Agilent). The qRT-PCR primers for the candidate protein-coding genes and lncRNAs were shown in Additional file 3: Table S3. The Bulge-LoopTM miRNA qPCR Primer Sets (RiboBio) were used for the expression detection of miRNAs. The expression levels of protein-coding genes and lncRNAs were normalized to GAPDH, while the miRNAs were normalized to U6. The statistical significance was analyzed by Student’s t-test.

Results
Significantly different transcriptome of FGR and control umbilical cord blood
To examine differences in the entire transcriptome between FGR cases and normal controls, we performed RNA sequencing for protein-coding genes, lncRNAs, and miRNAs in a case–control study with five pairs of umbilical cord blood samples (Additional file 1: Table S1). The differential expression analysis and hierarchical clustering showed that the protein-coding genes mostly tended to be down-regulated (Fig. 1a), while the IncRNAs (Fig. 1b) and miRNAs (Fig. 1c) tended to be up-regulated in the FGR cases as compared to the corresponding controls. All of the differentially expressed genes including protein coding genes, lncRNAs, and miRNAs were shown in Additional file 4: Table S4. The representative differentially expressed protein coding genes and lncRNAs were shown in Fig. 1d, e, respectively. These findings indicate that the entire transcriptome of umbilical cord blood from FGR case is significantly different from the corresponding control.

Differentially expressed protein-coding genes and physiological functional signaling pathways
Among the 339 differentially expressed protein-coding genes, 224 genes were down-regulated, and 115 genes were up-regulated in FGR cases (Fig. 2a, Additional file 4: Table S4). The GO enrichment analysis showed that these down-regulated genes were mainly enriched in plasma membrane, arachidonic acid binding, and neutrophil degranulation (Fig. 2b). Further signaling pathway analysis indicated that these genes were mainly involved in osteoclast differentiation, phagosome, and lysosome (Fig. 2c). Additionally, the up-regulated genes were mainly enriched in MHC class I protein complex binding, the cellular defense response (Fig. 2d), natural killer cell-mediated cytotoxicity and Graft-versus-host disease (Fig. 2e). Further GSEA results (Additional file 5: Table S5, Additional file 6: Table S6, Additional file 7: Table S7) showed that these genes were mainly enriched in glutathione metabolism (Fig. 2f), Alzheimer’s disease, Parkinson’s disease, Huntington’s disease (Fig. 2g), cardiac muscle contraction (Fig. 2h), systemic lupus erythematosus and oxidative phosphorylation (Fig. 2i). These findings indicate that the differentially expressed protein-coding genes are not only enriched in known FGR-related processes, such as metabolism and neural and cardiac systems, but also significantly associated with the immune system.

Differentially expressed miRNAs and physiological functional signaling pathways
Based on the top-30 highly varied miRNAs with larger log2(Fold Change) including the 13 significantly differentially expressed miRNAs (Fig. 3a, Additional file 4: Table S4), we predicted miRNA targets by miRanda, PITA, and RNAhybrid. The top-30 popular target genes as regulated by these miRNAs were shown in Fig. 3b, including the suppressor of cytokine signaling 1 (SOCS1), which is required for normal postnatal growth and survival [29, 30]. Further GO (Fig. 3c) and signaling pathway (Fig. 3d) analyses showed that the predicted targets of these miRNAs mainly focused on DNA binding, mitosis, integrated pancreatic cancer pathway, DNA damage response only ATM dependent, insulin...
signaling, and type II diabetes mellitus. An integrated analysis of the predicted targets and human diseases using the Human MicroRNA Disease Database (HMDD, v2.0) indicated that these miRNAs were significantly correlated with immune cells and published diseases (Fig. 3e). The clustering and family analysis showed that these miRNA precursors focused on the miR-194 family and Chr14_100911139-100911213 cluster (Fig. 3f), and neoplasm, leukemia, inflammation, and prolactinoma (Fig. 3g). For the 13 significantly differentially expressed miRNAs, we also observed the significance of the Chr14_100911139-100911213 cluster and prolactinoma.
Fig. 3 Differentially expressed miRNAs and physiological functional signaling pathways. 

a. Volcano plot of the differentially expressed miRNAs (11 up-regulated and 2 down-regulated).

b. The top-30 popular targets as predicted for the critical miRNAs (top-30).

c. GO-enrichment of the predictive targets of the critical miRNAs (top-30).

d. Signaling pathway enrichment of the predictive targets of the critical miRNAs (top-30).

e. Specific diseases in TCGA database correlated with the predictive targets of the critical miRNAs (left panel, top-30; right panel, 13 differentially expressed miRNAs).

f. Clustering and family analyses of the precursors of the critical miRNAs (top-30, red for 13 differentially expressed miRNAs).

g. Specific diseases correlated with the precursors of the critical miRNAs (top-30, red for 13 differentially expressed miRNAs).
These findings indicate that the significantly varied miRNAs are not only involved in FGR-related diseases but also focused on specific gene clusters and various diseases.

Core regulatory network and imprinted genes of FGR
To systematically investigate the core regulatory network of FGR at the entire transcriptome level, we performed RNA sequencing for the expression profile of lncRNAs, and found 79 lncRNAs significantly down-regulated and 216 lncRNAs significantly up-regulated (Fig. 4a, Additional file 4: Table S4). Further analyses of the regulatory relationship among the 339 protein-coding genes and 295 lncRNAs showed that seven lncRNAs were predicted to cis-regulate their neighboring protein-coding genes within a 2-kb region (Fig. 4b), and 7616 trans-regulatory relationships were obtained. To clearly demonstrate the trans-regulatory relationships, the top-35 significantly differentially expressed lncRNAs and protein-coding genes with larger degrees was shown (Fig. 4c). In combination of lncRNA-mRNA, miRNA-mRNA, and miRNA-lncRNA analyses, we observed 7 cis-, 59 trans-, and 2 miRNA regulatory relationships after screening with the Pearson correlation coefficient (PCC) (absolute value > 0.9) (Fig. 4d). The lncRNAs and protein-coding genes in cis-regulatory relationships had a positive PCC, while the lncRNAs and their trans-regulated protein-coding genes had a negative PCC (Fig. 4e).

As imprinted genes are critical in growth and development [31], we performed further analysis based on 240 imprinted genes obtained from two existed public databases (http://www.geneimprint.com and http://igc.otago.ac.nz/home.html) (Fig. 4g). The expression levels of these imprinted genes were significantly negatively associated with FGR (Fig. 4h), and six imprinted genes (Col9a3, Dlk1, Fucal1, Lirb4, Sfrp2, and Ventx) were significantly differentially expressed between FGR cases and controls (Fig. 4i). These findings observed a cluster of imprinted genes as correlated with FGR and might provide potential signatures for FGR.

Critical gene co-expression network modules closely correlated with FGR
To clarify the significant gene co-expression network involved in FGR, we performed WGCNA and clustered the entire transcriptome of FGR cases and controls (Fig. 5a). All genes focused on 18 modules (Additional file 8: Table S8), and most genes were clustered in the turquoise module as enriched in neutrophil degranulation (Fig. 5b). The module salmon was mainly enriched in metabolic process, while module green focused on regulation of immune response. The Pearson correlation analysis of the relationships between the network modules and sample characteristics showed that the turquoise and purple modules were significantly positively correlated with birth weight, while the turquoise and midnightblue modules were significantly different between FGR cases and controls (Fig. 5c). Similarly, we found that the cyan module was negatively correlated with maternal BMI and positively correlated with infant gender, and the tan module was negatively correlated with maternal age. Further investigation of the module stability suggested that the first four modules, including the turquoise, blue, brown, and yellow modules, exhibited much higher stability than the other modules (Fig. 5d).

Further hierarchical clustering indicated that the turquoise module was significantly correlated with birth weight and could clearly separate the FGR and control groups (Fig. 5e). The GO enrichment and signaling pathway analyses suggested that the turquoise module was mainly enriched in osteoclast differentiation and transcriptional misregulation in cancer (Fig. 5f). These findings indicate that the critical module turquoise is significantly correlated with FGR, but further confirmation of these genes in the module by a large sample size will provide more evidence for elucidating FGR.

The protein-coding genes, lncRNAs, and miRNAs as potential signatures for FGR
To detect the significance of differentially expressed transcriptome, we detected the expression level of the top-10 up-regulated and top-10 down-regulated molecules for protein coding genes and lncRNAs, six differentially expressed imprinted genes, and 13 differentially expressed miRNAs (Additional file 4: Table S4) in 12 FGR and 12 control umbilical cord blood samples (Additional file 2: Table S2) by quantitative RT-PCR analyses. The representative two imprinted genes (Sfrp2 and Dlk1), Slpi, and five lncRNAs (LINC01291, RP11_552M6.1, RP11_588G21.1, CTD_2083E4.5, and AMZ2P2) were shown in Fig. 6a. Furthermore, we performed a forward stepwise logistic regression analysis, and the ROC plot showed that RP11_552M6.1, LINC01291, and Asgr1 in final model could make the AUC value to 0.958 (Fig. 6b), indicating significantly predictive potential for FGR. To observe the application potential in clinic, we examined the expression level of the differentially expressed molecules in the 12 pairs of FGR case and control maternal peripheral blood samples. The ROC plot showed that the expression pattern of Sfrp2, miR-432-5p, and miR-1306-3p had significantly predictive power (AUC = 0.882) in maternal peripheral blood of FGR (Fig. 6c). These findings suggest that a cluster of protein-coding genes, lncRNAs, and miRNAs that is critically correlated with FGR
Fig. 4 Core regulatory network and significant imprinted genes of FGR. **a** Volcano plot of the differentially expressed IncRNAs (216 up-regulated and 79 down-regulated). **b** Predicted cis-regulatory relationships among the differentially expressed protein-coding genes and IncRNAs. **c** Representative trans-regulatory relationships among the top 35 differentially expressed protein-coding gene and IncRNAs. **d** Regulatory network among the differentially expressed protein-coding genes, IncRNAs, and miRNAs after screening with the correlation coefficient (>0.9). **e** PCC plots of all differentially expressed IncRNAs, protein-coding genes, and miRNAs. **f** Regulatory network among the differentially expressed protein-coding genes, IncRNAs, and miRNAs after screening with the correlation coefficient (>0.9). **g** Circos plot of all differentially expressed protein-coding genes, IncRNAs, and miRNAs. **h** GSEA plot of all imprinted genes in FGR and control umbilical cord blood samples. **i** Six imprinted genes (Col9a3, Dlk1, Fuca1, Lilrb4, Sfrp2, and Ventx) significantly differentially expressed in umbilical cord blood of FGR versus control.
and may provide potential signatures of FGR. Further investigation of their diagnostic potential in a large sample size at early stage of pregnancy, and the functional study on significant signatures may improve the understanding of FGR.

Discussion

Due to the lack of case–control studies investigating fetal-originated umbilical cord blood, the regulatory networks of FGR remain unclear, and the early prediction and diagnosis of FGR are challenging. In this case–control study, we performed a systematic whole-transcriptome profiling of the functional regulatory networks in human FGR case and control umbilical cord blood samples. The differentially expressed whole-transcriptome, imprinted genes, and weighted gene co-expression analyses revealed an enrichment in functional processes and critical modules related to growth, development, and the immune system.

Several studies have demonstrated that maternal factors such as age, BMI, preeclampsia, chronic hypertension, and anemia are associated with FGR [16, 32]. Fetal factors and placental factors, including malformation, infections, and abnormal placental vascular system are also associated with FGR. The measurement of placental biomarkers in maternal blood is a common method used to evaluate placental functions related to pregnancy outcomes [33]. However, these placental molecules are heterogeneous and identified in various locations, such as placental tissue, amniochorionic membranes, amniotic fluid, cord blood and maternal blood [34]. The sample heterogeneity in the placenta and multifactorial nature of FGR especially abnormal fetal growth, should be confirmed by repeated ultrasound. Thus, matching controls to each FGR infant based on these characteristics could eliminate their potential influences on fetal weight and yield more constructive evidences. Our study focused on a case–control study of the entire transcriptome of umbilical cord blood to demonstrate the most core gene networks in FGR. The combination of the core gene expression signatures and ultrasound confirmation could more effectively separate normal small infants from pathological FGR infants, and lead to an effective strategy for FGR intervention or even prevention at an early time-point.

As FGR confers a high risk of increased perinatal, childhood, and adulthood complications, effective screening and treatment procedures are critical for avoiding adverse health outcomes in neonates born with FGR [35]. Here, we used RNA sequencing to assess the entire transcriptome (mRNA, IncRNA, and miRNAs), then uncovered the significant signatures and potential signaling pathways of FGR. It has been reported that FGR may lead to a cardiovascular risk, metabolic problems, and poor neurodevelopmental outcomes in adulthood [36, 37]. Based on system biology and network-based analyses of the differentially expressed whole-transcriptome, this study not only confirmed that the differentially expressed genes of FGR at the whole-transcriptome level are significantly enriched in metabolism and neural and cardiac systems but also elucidated the critical correlation between the differentially expressed transcriptome and immune system diseases, such as Graft-versus-host disease and systemic lupus erythematosus. These findings provide more potential and effective approaches for immune treatment of FGR. Since human umbilical cord blood includes various cell types such as lymphocytes, monocytes, and mesenchymal stem cells [38–40], which may have different effects on the transcriptome profile of protein-coding genes, IncRNAs, and miRNAs. To clearly explore the exact contribution of each cell source of whole blood on RNA profile may provide more specific evidence in the future.

To better capture the inter-gene relationships and define the co-regulatory patterns involved in FGR, we performed WGCNA to explore the systems-level expression changes and clustered the highly-correlated genes into co-expression modules [10, 11]. This study identified five modules related to maternal–infant demographic variables. Interestingly, we found that the turquoise module could clearly separate the FGR group from the controls and was positively correlated with birth weight among the FGR cases. Similarly, we found that the cyan module was negatively correlated with maternal BMI and positively correlated with infant gender, while the tan module was negatively correlated with maternal age. It has been reported that infant gender may affect the placental gene expression and function [41, 42]. Whether infant gender affects the gene expression profile of umbilical cord blood needs further study with a large sample size.
And the effects of infant gender on these genes enriched in the tan module also needs further confirmation. To evaluate the utility of these modules in elucidating the molecular underpinnings of FGR, we performed GO and signaling pathway enrichment analyses of each module. These analyses indicated that the significant FGR modules focused on metabolism, immune systems and transcriptional misregulation. However, further studies are warranted to determine whether these gene signatures are relevant to postnatal health, and to provide further
mechanistic insight into the consequences of abnormal fetal growth.

Furthermore, to systematically investigate the core regulatory network of FGR at the whole-transcriptome level, we performed an integrated analysis of protein-coding genes, lncRNAs, and miRNAs. By combining the lncRNA-mRNA, miRNA-mRNA, miRNA-lncRNA, and protein–protein interaction (PPI) analyses, we observed 7 cis-, 59 trans-, 2 miRNA, and 70 PPI regulatory relationships. LncRNAs and protein-coding genes in cis-regulatory relationship had a positive PCC, while lncRNAs and trans-regulated genes had a negative PCC, which is similar to miRNAs and their targets. It has been reported that imprinted genes can regulate growth and development [31]. Here, we found that the expression levels of six imprinted genes (Col9a3, Dlk1, Fuc1, Lirlb4, Sfpr2, and Ventx) significantly differed between FGR cases and controls. Furthermore, we confirmed the expression level of the top-10 differentially expressed protein-coding genes and lncRNAs, six imprinted genes and 13 miRNAs in umbilical cord blood and maternal peripheral blood. The ROC plots showed that RP11_552M6.1, LINC01291, and Asgr1 in umbilical cord blood, while Sfpr2, miR-432-5p, and miR-1306-3p in the maternal peripheral blood had significantly predictive power for FGR. These findings suggest a cluster of molecular signatures that are potentially diagnostic and predictive markers of FGR.

Since the limited sample size used in this study, the stability of the gene modules defined by the WGCNA has been further investigated by the bootstrap method [28]. In consistent with Shannon et al., the distribution of the Jaccard similarity coefficients of each module between the real data and bootstrapped set suggested that the gene modules with large gene sizes had higher stability than the small sized modules. Furthermore, we confirmed the turquoise, blue, brown, and yellow modules exhibited much higher stability than the other modules as compared with the best-case stability of the random modules in the simulation. Furthermore, considering with the limited sample size in this study, these 95% CI AUCs of logistic regression models were also calculated by 1000 bootstraps. Then, these findings need further investigation of the diagnostic potential of these potential molecules in a large sample size, and their relevance to postnatal health in childhood and adulthood for the consequences of abnormal fetal growth.

Conclusions
This study comprehensively profiled the transcriptome-wide landscape of human umbilical cord blood, constructed the core gene co-expression network, delineated the critical gene signatures including imprinted genes correlated with FGR, and provided key insight into intrauterine perturbations and candidate signatures of FGR. However, it needs further exploration for the diagnostic significance in a large sample size during early stage of pregnancy, and functional and mechanistic study to provide more evidences for elucidating FGR.

Supplementary information
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Authors’ contributions
GYW, CQL, JY, TD, and JHK designed the study. GYW, JY, and XBZ collected samples. GYW and CQL ran the analyses. YWY and XQL performed verification experiments. GYW, CQL, XDG, TD, and JHK interpreted the results and wrote the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
The RNA sequencing data were deposited in the National Omics Data Encyclopedia (NODE) database (https://www.biosio.org/node) under project ID: OEP000732. The data that support the findings of the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
All women provided written informed consent, and the study protocol was approved by the Ethics Committee of Shanghai First Maternity and Infant Hospital (No. KS17115).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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