Prenatal inflammation exposure-programmed hypertension exhibits multi-generational inheritance via disrupting DNA methylome

Xiao Guan1, Guo-rong Dan1, Yao Yang1, Yan Ji1, Wen-jing Lai1, Fang-jie Wang1, Meng Meng1, Bang-hui Mo1, Pei Huang1, Ting-ting You1, Ya-fei Deng1, Liang Song1, Wei Guo1, Ping Yi2, Jian-hua Yu3,4, Yuan Gao5, Wei-nian Shou1,6, Bing-bo Chen7, You-cai Deng1 and Xiao-hui Li1

The multi-generation heredity trait of hypertension in human has been reported, but the molecular mechanisms underlying multi-generational inheritance of hypertension remain obscure. Recent evidence shows that prenatal inflammatory exposure (PIE) results in increased incidence of cardiovascular diseases, including hypertension. In this study we investigated whether and how PIE contributed to multi-generational inheritance of hypertension in rats. PIE was induced in pregnant rats by intraperitoneal injection of LPS or Poly (I:C) either once on gestational day 10.5 (transient stimulation, T) or three times on gestational day 8.5, 10.5, and 12.5 (persistent stimulation, P). Male offspring was chosen to study the paternal inheritance. We showed that PIE, irrespectively induced by LPS or Poly (I:C) stimulation during pregnancy, resulted in multi-generational inheritance of significantly increased blood pressure in rat descendants, and that prenatal LPS exposure led to vascular remodeling and vasoconstrictor dysfunction in both thoracic aorta and superior mesenteric artery of adult F2 offspring. Furthermore, we revealed that PIE resulted in global alteration of DNA methylome in thoracic aorta of F2 offspring. Specifically, PIE led to the DNA hypomethylation of G beta gamma (Gβγ) signaling genes in both the F1 sperm and the F2 thoracic aorta, and activation of PI3K/Akt signaling was implicated in the pathologic changes and dysregulated vascular tone of aortic tissue in F2 LPS-P offspring. Our data demonstrate that PIE reprogrammed DNA methylome of cells from the germline/mature gametes contributes to the development of hypertension in F2 PIE offspring. This study broadens the current knowledge regarding the multi-generation effect of the cumulative early life environmental factors on the development of hypertension.

Keywords: hypertension; multi-generational inheritance; prenatal inflammation exposure; DNA methylation; Gβγ; PI3K/Akt signaling; LPS; poly (I:C)

Acta Pharmacologica Sinica (2022) 43:1419–1429; https://doi.org/10.1038/s41401-021-00772-8

INTRODUCTION
Hypertension is the leading risk factor for cardiovascular diseases (CVDs), affecting the health of more than 1 billion people worldwide [1]. A recent multi-generation cohort study of Framingham Heart Study participants has demonstrated that early-onset hypertension not only in parents but also in grandparents could raise the risk for hypertension in their descendants, indicative of multi-generation heredity of hypertension [2]. However, the detailed mechanisms are largely unknown.

Prenatal inflammation exposure (PIE), a momentous event during pregnancy, could be caused by various pathogenesis, including “hot” inflammation after pathogens infections (virus, bacterial, and parasites infection) and also “cold” inflammation in several chronic diseases (such as obesity, maternal diabetes or hyperglycaemia, hypertension, pre-eclampsia, and smoking) [3]. Decades of research works have shown the close association between PIE and neuropsychiatric disorders, such as autism and schizophrenia [4]. Recent epidemiological and experimental studies have also revealed that prenatal influenza or bacterial exposure results in increased incidence of CVDs, including hypertension (summarized in our and others’ recent published reviews) [3, 5–7], which provide a close linkage between PIE and CVDs in adult offspring (called as PIE-programed CVDs). Due to the pandemic, over 150 million people have Coronavirus disease (COVID-19). Pregnant women with COVID-19 also showed an obvious proinflammatory status, such as increased levels of IL-6, IL-8, and TNF-α [8].
IFNγ, C-reactive protein, and D-dimer, in periphery blood serum [8]. All these evidence demonstrate a great need to elucidate the detailed mechanisms for PIE-induced CVDs, which may provide new insight into preventing the continuous increasing rate of CVDs incidence post COVID-19 pandemic [9].

Rats have been used as the primary model for studying the genetic mechanisms of hypertension [10]. A rat model of PIE-programed CVDs has been used previously by treating pregnant Sprague–Dawley (SD) rats with lipopolysaccharide (LPS, the main component of gram-negative bacteria cellular wall) and polyribinosinosic-polyniroctidylic acid (Poly (I:C)), a substitute for viral dsRNA and a ligand of toll-like receptor 3), during the secondary semester, a similar strategy that is extensively used in studying the mechanisms of PIE-induced neuropsychiatric disorders [4]. Irrespective of transient stimulus (one time on gestational day 8.5, 10.5, and 12.5) or persistent stimulus (three times on gestational day 8.5, 10.5, and 12.5) of LPS or Poly (I:C) challenge, the F1 offspring showed aortic dysfunction and eventually developed hypertension at elder age [11–13]. However, whether PIE could contribute to the multi-generation heredity of hypertension remains largely unknown.

Recent studies have demonstrated that epigenetic features, such as DNA methylation by covalent modifications, can be transmitted for several generations. DNA methylation pattern could better reflect the cumulative effects of environmental and lifestyle factors than expression profiles that are subject to transient changes [14–16]. To this end, in this current study, we investigated the phenotype of transgenerational inheritance in multiple generations relevant to PIE-programmed hypertension and explored the potential mechanisms regarding DNA methylation. We found that PIE-programed hypertension, irrespective of induced by prenatal LPS or Poly (I:C) stimulation and irrespective of stimulation duration, was inherited by several generations, accompanied by altered DNA methylome. Specifically, PIE led to the DNA hypomethylation of G beta gamma (Gβγ) signaling genes in both the sperm of F1 offspring and the thoracic aorta of the second generation (F2) offspring, accounting for the pathologic changes and dysregulated vascular tone of aortic tissue via phosphatidylinositol 3-kinase/protein kinase B (PI3K/Akt) signaling pathway. These data demonstrated that PIE reprogrammed the DNA methylome of cells from the germline/mature gametes, which could be inherited by several generations, contributing to the increased risk of hypertension.

MATERIALS AND METHODS

Experimental animals
Nonparous SD rats were purchased from the Animal Centre of Army Medical University (Chongqing, China). All rats were bred and housed in pathogen-free conditions, at 22–25 °C, periodic air changes, and free access to water and food, and had 12-h light/dark cycle, at the Experimental Animal Center of the Animal Centre of Army Medical University (Chongqing, China), according to ARRIVE guidelines and the guidelines of Institutional Animal Care and Use Committee. All experiments performed were approved by the Institutional Animal Care and Use Committee at the Animal Centre of Army Medical University (Chongqing, China), and complied with the NIH guidelines or the guidelines from Directive 2010/63/EU of the European Parliament on the protection of animals used for scientific purposes.

Experimental design
F0 maternal inflammatory stimulation: time-dated nulliparous pregnant SD rats (250–300 g) were randomly divided into five groups (n = 8 for each group), intraperitoneally (i.p.) injected with saline on embryonic day 8.5, 10.5, and 12.5 (E8.5, 10.5, and 12.5); LPS (0.79 mg/kg, Sigma-Aldrich, L2630, St. Louis, MO, USA) on E8.5, 10.5, and 12.5; LPS (0.79 mg/kg) on E10.5; Poly (I:C) (4 mg/kg, Sigma-Aldrich, P1530) (only included the weight of the Poly (I:C) itself on E8.5, 10.5, and 12.5); and Poly (I:C) (4 mg/kg) (only included the weight of the Poly (I:C) itself on E10.5, named as Control, LPS-P, LPS-T, Poly (I:C)-P, and Poly (I:C)-T group, respectively). "P" indicates persistent stimulation, and "T" indicates transient stimulation, respectively [12]. For LPS-T and Poly (I:C)-T groups, pregnant SD rats were also i.p. injected with saline on E8.5 and E12.5. No convulsions, deaths, or miscarriage occurred after LPS or Poly (I:C) injection in all the treated pregnant rats. Pups were delivered around 21 days of pregnancy and no obvious deformities were found.

After delivery, litter size was adjusted to eight pups per dam by random selection to avoid the disproportionate of nutrient availability during lactation. Offspring rats from F0 mothers were designated as F1 control or PIE offspring. Multi-generational inherited rat model of PIE referred to the previously reported mouse model of intrauterine growth restriction [17]. F1 pups were weaned freely and weaned at the age of 4 weeks and provided with standard chow ad libitum. As it is difficult to rule out the influence of intrauterine microenvironment changes, maternal metabolism, and behavior on the offspring physiology by using maternal transmission model to clarify transgenerational inheritance [18], we chose male offspring to study the paternal inheritance. External control virgin females (250–300 g) were bred with either F1 control or PIE males, randomized selected from each litter of rats, to generate F2 PIE descendants, named as F2 control, F2 LPS-P, F2 LPS-T, F2 Poly (I:C)-P, F2 Poly (I:C)-T, respectively (Fig. 1a). At birth, F2 litters were adjusted to eight pups per dam and had standard chow ad libitum at weaning at the age of 4 weeks. F2 control and F2 LPS-P males were bred with external control females to generate F3 offspring, while F3 LPS-P males were further bred with external control females to generate F4 offspring (Fig. 1a). For F1–F4 offspring, pregnant rats and pups were no longer subjected to the stimulation with LPS or Poly (I:C). Rats were anesthetized with sodium pentobarbital (240 mg/kg, i.p.) and were sacrificed by cervical dislocation if necessary.

Blood pressure measurement
Systolic blood pressure (SBP) was measured by the tail-cuff system (Softren BP-2010A, Softron Biotechnology Co., Beijing, China) every 4 weeks starting at 8 and ending at the age of 28 weeks, as described previously [12]. The investigators were blinded for measuring the blood pressure.

Histological analysis
The rats were anesthetized via intra-abdominal injection with pentobarbital (100 mg/kg, Sigma-Aldrich, P-010) and euthanized by cervical dissociation. The thoracic aortas and superior mesenteric arteries were fixed in 4% paraformaldehyde (Sigma-Aldrich, 158127) for 7 days, soaked in 30% sucrose, embedded in paraffin wax, sectioned (4 μm), and then stained with haematoxylin and eosin as described previously [19]. The sections were visualized under an Olympus BH-2 microscope. The values of vascular wall thickness, cross-sectional area, and vascular smooth muscle cells (VSMCs) width were quantified by using ZEN 2.3 (blue edition). The investigators were blinded for acquiring and quantifying the images.

Immunofluorescence staining
The protein levels of eNOS in thoracic aortas were determined by immunofluorescence staining, as described previously [19]. Goat anti-mouse IgG H&L (Alexa Fluor® 647) (ab150115) antibodies (Abcam, La Jolla, CA, USA) were used for an additional 1 h incubation after anti-eNOS (ab76198) antibodies (Abcam, La Jolla, CA, USA) incubation and washing. Sections were then incubated with 1 mg/mL DAPI for 10 min, washed three times with PBS, and mounted into Vectashield™ mounting medium (Vector Laboratories, Burlingame, CA, USA). The coverslips were visualized under a Leica confocal laser scanning microscope (Leica, Wetzlar, Germany). The investigators were blinded for acquiring the images.
Vascular reactivity assessment

The procedure of vascular reactivity assessment was performed as described previously [20]. Briefly, rats were euthanized, thoracic aortas and superior mesenteric arteries were cut into 4-mm rings, stored in an oxygenated Krebs–Henseleit (K–H) solution (mM: NaCl, 11.8; KCl, 0.47; KH2PO4, 0.13; MgSO4, 0.045; NaHCO3, 2.5; CaCl2, 2; glucose, 11.1) at 4 °C and gassed with a carbogenic mixture (95% O2 and 5% CO2). Vascular rings were suspended on triangular-shaped stainless steel hooks in warmed (37 °C) and aerated (95% O2, 5% CO2) K–H solution using Automatic organ bath (Panlab, Harvard Apparatus, USA). A computerized force transducer (Powerlab System, AD Instruments, Australia) was used to record the isometric tension of the aortic and mesenteric arteries rings. After 60 min of equilibration, each ring was challenged with KCl (60 mM). Phenylephrine (PE) (Sigma-Aldrich, P1240000) was added cumulatively (10−6–3 × 10−6 M) to the bath to establish a concentration–response contraction curve, and the contraction was expressed as a percentage of the contraction produced by 60 mM KCl for each ring. Acetylcholine (ACh) (Sigma-Aldrich, A6626) or sodium nitroprusside (SNP) (Sigma-Aldrich, 1614501) with cumulative concentrations (10−10–3 × 10−6 M) were added to endothelium-intact or de-endothelialized aortic rings, respectively, to establish a concentration–response relaxation curve. Relaxation response was expressed as a percentage of the maximal contractile response induced by PE (3 × 10−6 M). Concentration–response curves were analyzed by linear regression analysis (fitted to the Hill equation) using Prism (GraphPad Software, CA, USA), and expressed as the values of pD2 (−log EC50). To determine the role of PI3K-Akt in vascular function, PI3K inhibitor (LY294002, 20 μM) (Sigma-Aldrich, 440202) [21] was added 30 min before vascular reactivity assessment and stayed in the Automatic organ bath during the whole vascular reactivity assessment.

MeDIP-seq

Four samples from both the F2 control and F2 LPS-P groups were used for methylated DNA immunoprecipitation sequencing (MeDIP-seq). Sample preparation and MeDIP-Seq were accomplished by Kangcheng Bio-tech, Shanghai, China. Briefly, high-quality genomic DNA (gDNA) isolated from thoracic aorta tissues (3.5 – 4.5 mg) was fragmented to a size range of 200–700 bp with a Diagenex Bioruptor, and a portion of the denatured DNA fragments was taken as input DNA. About 1 μg fragmented DNA was prepared for Illumina HiSeq 2000 sequencing by using the following steps: (1) end repair of DNA samples with T4 DNA polymerase, Klenow DNA polymerase, and T4 polynucleotide kinase (T4 PNK); (2) a single “A” base was added to the 3’ ends with Klenow (exo minus) polymerase; (3) ligation of Illumina’s genomic adapters to DNA fragments; (4) MeDIP to the enrich methylated DNA by using anti-5-methylcytosine antibodies (Abcam, ab10805); (5) PCR amplification of the enriched precipitated DNA fragments; and (6) gel purification to extract the DNA fragments with length of 300–800 bp. The completed libraries were quantified by Agilent 2100 Bioanalyzer. The libraries were then denatured with 0.1 M NaOH (Sigma-Aldrich, S8045) to generate single-stranded DNA molecules, captured on Illumina flow cell and amplified in situ. The libraries were then sequenced on the Illumina HiSeq 2000 following the TruSeq SBS Kit v5 protocol.

Raw files were processed by FastQC for quality control. Quality and adapter trimming was performed using Cutadapt. Reads were mapped to the respective reference genomes (rn6) using BOWTIE software (V2.1.0). The average alignment rate of MeDIP-seq reads from all samples was 96.83%, and on average 76.61% of reads were uniquely mapped to the reference genome (Supplementary Fig. S1a). Average enrichment score was 1.25 in all tested samples (Supplementary Fig. S1b). MEDIPS saturation analysis indicated that Pearson’s correlation values of all samples were close to one, and the read coverage was sufficient for next downstream analysis (Supplementary Fig. S1c). The CpG coverage plots also illustrated similar shape between the two groups and indicated an effective MeDIP enrichment. On average, 12.45% of mapped reads did not cover a pattern (Supplementary Fig. S1d). The data showed that methylated DNAs were properly captured by MeDIP-seq used in this study. Differentially methylated regions (DMRs) were identified using MEDIPS package version 1.12.0 with the condition that log2 (fold-change) ≥ 2 and FDR <0.05. Then we used ChiPSeeker to annotate DMRs using m6A known gene table from UCSC. The reads were visualized and individual genes were examined using WashU Epigenome Browser.

To conduct enrichment analysis, the package cluster Profiler (version 3.2.14) of R (version 3.3.3) was used for analyzing the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways and Gene Ontology (GO) processes based on the DMR associated genes. The Ingenuity Pathway Analysis (IPA) (https://www.qiagen.com/us/products/informatics-and-data/interpretation-content-databases/ingenuity-pathway-analysis/#orderinginformation) was also used for analyzing the canonical pathways and interaction networks. P value for the pathway enrichment is calculated by using Fisher’s exact test, and P < 0.05 is considered statistically significant. Then the results were imported to Cytoscape (version 3.4.0) to construct gene-pathway network.

BSP

gDNA isolated from both F2 thoracic aortas and F1 sperm was carried out by using QIAamp DNA Mini Kit (QIAGEN, 51304, Hilden, Germany). Bisulfate treatment of gDNA was performed by using EZ DNA Methylation-Gold Kit™ (ZYMO Research, D5005, Irvine, CA, USA), according to the manufacturer’s instructions. PCR primers were designed with Meth Primer (http://www.urogene.org/methprimer/), andbisulfate-treated DNA was amplified with Epicq Tq polymerase (Takara, R110Q, Beijing, China).

The PCR products were separated by using agarose gel electrophoresis and purified by using the Wizard® SV Gel and PCR Clean-Up System (Promega, A9281, Madison, WI, USA). Purified DNA fragments were cloned into pMD T-19 (Takara, No. 6013Code) and ten clones from each PCR assay were subjected to next direct sequencing. The final sequence results were analyzed by online QUMA (quantification tool for methylation analysis) software [22]. Primer sets used in bisulfate modified PCR are shown in Supplementary Table S1.

Real-time RT-PCR

Total RNA was extracted from thoracic aortas by using TRIzol Reagent method (Invitrogen, 15596-026) and then reverse-transcribed into cDNA by using a Bestar™ quantitative real-time polymerase chain reaction (qPCR) RT Kit (DBI Bioscience, DBI-2220, Ludwigshafen, Germany), as described previously [23]. qPCR was performed with Bestar™ Sybr Green qPCR Master Mix (DBI Bioscience, DBI-2043) according to the following conditions: 2 min at 95 °C; 40 cycles at 95 °C for 10 s, 60 °C for 12 s, 72 °C for 20 s; 5 min at 72 °C, using ABI Prism 7700 sequence detection system (Applied Biosystems). Primer sequences for qPCR were designed by Pubmed Primer-BLAST and listed in Supplementary Table S2. Relative mRNA expression was calculated by normalizing the circulating threshold with β-actin and then normalizing the relative gene expression to the control group.

Immunoblotting

Protein expression in thoracic aortas was determined by immunoblotting, as described previously [23]. Briefly, thoracic aortas were homogenized with T-PER tissue protein extraction reagent (Thermo-Pierce, 78510, Rockford, IL, USA) containing protease inhibitor cocktails (Sigma-Aldrich, P8340) and protein concentrations were determined by BCA Protein Assay Kit (Bio-Rad, 5000001, Hercules, CA, USA). Proteins from each sample (20
PIE and multi-generational inherited hypertension
X Guan et al.

μg) were separated by 8%–12% Tris SDS-PAGE gel electrophoresis and transferred to polyvinylidene difluoride membranes (Bio-Rad Laboratories, Inc., 1620256, CA, USA). After blocking with 5% non-fat dry milk, membranes were incubated with anti-p-Akt (Ser473), anti-p-S6 Ribosomal Protein (Ser235/236), anti-Akt (C73H10), anti-S6 Ribosomal Protein (Cell Signaling Technology, 4058; 4856; 2938; 2217, Beverly, MA, USA), or anti-β-actin (AC-15) (Sigma-Aldrich, A5441) antibodies overnight at 4 °C, and incubated with the secondary antibodies (Santa Cruz, sc-2004; sc-2005, Santa Cruz, CA, USA) at room temperature for 1 h. Proteins were visualized with enhanced chemiluminescence reagents (Thermo-Pierce, Rockford, IL, USA) and the blots were exposed to hyperfilm. Results were quantified by using Quantity-one software (Bio-Rad).

Statistical analysis
Results are expressed as mean ± SEM. Statistical analysis was performed by using an unpaired Student’s t test (two unpaired groups) or two-way ANOVA followed by Sidak’s multiple comparison test (multiple groups comparison) by using SPSS 18.0 software (SPSS Inc., Chicago, IL, USA). All of the statistical tests were two-tailed, and P < 0.05 was considered to represent a significant difference.

RESULTS
PIE-programmed hypertension is inherited by multi-generation in rat offspring
The Bisulfite sequencing PCR (BSP) of F1 PIE male offspring, including F2 LPS-P, F2 LPS-T, F2 Poly (I:C)-P, F2 Poly (I:C)-T, was significantly higher than that in F1 control offspring (Supplementary Fig. S2a). To interrogate whether the increased blood pressure could be inherited by multiple generations, we first determine the SBP of F2 PIE offspring. The SBP of F2 PIE offspring from whose fathers experiencing transient or persistent treatment with LPS or Poly (I:C) prenatally was all significantly elevated from 8 weeks old onwards than that of F2 control offspring (Fig. 1a–c), which is similar to those found in F1 PIE progenies [12, 13]. Compared with F2 control offspring, more F2 PIE offspring were distributed at higher blood pressure values, while the maximum value of blood pressure in F2 PIE offspring was also greater than that of control F2 offspring, irrespective of F1 fathers experiencing prenatal LPS or Poly (I:C) exposure (Fig. 1d, e). Further analysis revealed that the SBP of both male and female F2 PIE offspring showed statistical difference with F2 control offspring, indicative of no sexual difference (Supplementary Fig. S2b–e). We further selected F2 LPS-P model to verify the transgenerational inheritance of increased SBP in F3 and F4 offspring. The SBP in both F3 and F4 offspring from F1 fathers experiencing prenatal persistent treatment with LPS (F3 LPS-P and F4 LPS-P) showed similar increased levels as seen in F2 LPS-P offspring (Fig. 1f–i).

Histological image of vessel wall from F2 control thoracic aorta clearly displayed the tunica intima-media- adventitia layering, and the media consist of regularly organized musculo-elastic complexes. However, the morphologic observation revealed that F2 LPS-P offspring (whose fathers experiencing three times of LPS stimulation on E8.5, 10.5, and 12.5 prenatally) showed obvious vascular remodeling in both thoracic aortas and mesenteric arteries, such as increased arterial vascular wall thickness, wall thickness/lumen diameter ratio, and VSMCs width (Fig. 2a, b). Some of the endothelial cells of thoracic aortas from F2 LPS-P offspring were seen to be necrotic and detached from the subendothelial layer. Fragmented and unclear internal elastic fibers were also exhibited in thoracic aorta of F2 LPS-P offspring (Fig. 2a, b). In addition, hypertrophic and disarranged VSMCs were observed migrating and breaking through the internal elastic lamina (Fig. 2a, b). In addition, we also determined the changes of endothelial structure by immunofluorescence staining of eNOS, which is a critical marker for endothelium integrity in the thoracic aorta of both F2 control and F2 LPS-P offspring. The fluorescently labeled eNOS in the endothelium are shown in red as indicated by the arrow. The fluorescent labeling of eNOS in F2 control showed smooth and intact vascular endothelium, while the endothelium was shrunken, broken, and had increased gaps in F2 LPS-P offspring. This result suggests that the endothelium of thoracic aorta in F2 LPS-P offspring has obvious tissue structure abnormalities (Supplementary Fig. S3a).

The vascular reactivity assay also revealed that both the thoracic aorta and mesenteric artery from F2 LPS-P offspring showed dramatically higher contractile responses to PE without affecting the ACH-induced vasodilation (Fig. 3a–d). As renin-angiotensin system (RAS) is critical to vascular contraction [1, 18], we next determined the mRNA levels of genes in RAS family, such as angiotensin-converting enzyme (ACE), angiotensin II receptor type 1 (AT1R), angiotensinogen (Agt) in both the thoracic aorta and SMA of both F2 control and F2 LPS-P offspring. The data revealed that in the thoracic aorta and SMA of F2 LPS-P, the mRNA expression of ACE, which is responsible for catalyzing the conversion of angiotensin I to angiotensin II, was significantly higher than F2 control offspring (Supplementary Fig. S3b).

Therefore, our data revealed that PIE induced multi-generational effect specifically on hypertension.

PIE alters overall methylation patterns in the aortic tissue of F2 offspring
Epigenetic mechanisms, referring to chromatin-based mechanisms without affecting DNA sequence, have emerged as a major goal in researching on hypertension, including DNA methylation, histone modification, microRNAs, and long noncoding RNAs [24]. DNA methylation pattern could better reflect the cumulative effects of environmental and lifestyle factors than expression profiles that are subject to transient challenges [16, 25]. It is also implicated in the transmission of environmentally acquired phenotypes [26]. Therefore, we next determined the DNA methyleome distribution of the thoracic aorta from F2 PIE offspring by MeDIP-seq, and followed by the verification in PIE F2 sperm samples. To produce the most robust phenotype, PIE F2 LPS-P offspring at the age of 16 weeks from whose fathers experiencing prenatal persistent LPS stimulation were chosen. Our first set of data demonstrated 1111 hypomethylated DMRs and 1696 hypermethylated DMRs (loci with a methylation change >1.5-fold and an edge P value <1 × 10−5) (Supplementary Fig. S4a, b and Supplementary Data 1 and 2) in thoracic aortas between F2 control and F2 LPS-P offspring, indicative of a global changed DNA methylation occurring in PIE F2 offspring. BSP validation of three hypomethylated (Cfhr1, Ndrg2, and Cox7a2l) and three hypermethylated (Fgf2, Pfdn1, and Srpk2) DMRs within the promoter region or with the greatest fold-change between F2 control and F2 LPS-P offspring was shown to be remarkably consistent with the findings derived from MeDIP-seq data (Supplementary Fig. S5).

Annotation with genomic features illustrated that the majority of both hypomethylated and hypermethylated DMRs were located in intron, extron, 5′UTR, and 3′UTR in intragenic element-associated regions. Interestingly, hypomethylated DMRs were distributed more at the proximal promoter region of ~1 kb upstream and ~500 bp downstream of the transcriptional start site (TSS) (labeled as Promoter-TSS) and the region of the transcriptional termination site than those in hypermethylated DMRs (Supplementary Fig. S4c). David GO Biological Process enrichment and KEGG analysis demonstrated that genes with hypomethylated DMRs were mainly enriched in positive regulation of RNA transcription and gene expression, endothelial cell migration, NF-κB transcription factor activity, MAPK activation, Ras, and Jak-STAT signaling pathway (Fig. 4a, b), all of which were implicated in the cardiovascular pathological changes and
activation of vascular remodeling signaling pathways [27, 28]. In addition, genes with hypermethylated DMRs were mainly enriched in brain and nervous system development, synapse assembly, cAMP, and TGF-β signaling pathway (Fig. 4c, d). These suggested that genes with hypomethylated DMRs were mainly involved in the development of multi-generational inherited hypertension in PIE descendants, which was consistent with the findings in previous human epigenetic contribution to hypertension [29, 30]. This is also consistent with recent studies that heritable hypomethylated chromosomal segments could be propagated for eight or more generations [31].

IPA software package to identify the enriched canonical pathways and IPA interaction networks demonstrated that there were several significantly enriched signaling pathways in genes...
with hypomethylated DMRs associated with vascular remodeling and the pathogenesis of hypertension, such as Gβγ signaling, JAK/STAT signaling, renin-angiotensin signaling, LPS-stimulated MAPK signaling, p53 signaling, p70S6K signaling, GPCR signaling, and ERK/MAPK signaling pathways (Fig. 4f and Supplementary Data 3) [27, 28, 32]. Gβγ signaling is the fourth most significant canonical pathway enriched in genes with hypomethylated DMRs (P = 4.81 × 10^-4, overlap = 12.5%) (Fig. 4e). The IPA network interaction analysis suggested that Gβγ signaling was closely correlated with GPCR, JAK/STAT, LPS-stimulated MAPK, and ERK/MAPK signaling (at least four genes shared by both signaling pathways) (Fig. 4f). This suggested that gene clusters in Gβγ signaling with hypomethylated DMRs might play a core role in the vascular pathological changes and development of hypertension in PIE F2 offspring.

Validation of hypomethylated DMRs of genes clustered in Gβγ signaling in both the F2 thoracic aorta and F1 sperm of PIE LPS-P offspring

As stated above, we next validated the mRNA expression levels and DNA methylation status within the promoter region of the genes in Gβγ signaling pathways by qPCR and BSP, respectively. The data demonstrated that the mRNA expression of the genes in Gβγ signaling showed an increased trend in thoracic aortas of PIE F2 LPS-P offspring. Among these genes, the mRNA expression levels of Sos1, Sos2, Gng10, Prkacb, and Pik3c3 showed statistical
Fig. 3  Prenatal LPS exposure leads to vasoconstrictor dysfunction in both the thoracic aorta and superior mesenteric artery (SMA) of adult F2 offspring. a–d Vascular reactivity of both the thoracic aorta (a and c) and SMA (b and d) rings from 16-week-old F2 offspring. PE-induced cumulative concentration–response curves (a, b) and ACh-induced relaxation–response curves (c, d) were measured in endothelial-intact thoracic aorta and SMA rings from 16-week-old F2 offspring. Concentration–response curves are expressed as the percentage of contractions caused by KCl (60 mM), and relaxation–response curves are expressed as the percentage of the maximal PE-induced contractions. Data are presented as means ± SEM. n = 8 from 4 litters for each group for a–d. **P < 0.01 denotes the statistical comparison between two marked groups (two-way ANOVA followed Sidak’s multiple comparisons test for a–d). The pD2 (-log EC50) values were calculated for each group and shown on the top left of each panel.

significance (Fig. 5a). BSP data revealed that the ratio of methylated DNA within the promoter region of Sos1, Sos2, Gng10, and Prkacb was statistically reduced in thoracic aortas of F2 LPS-P offspring as compared to those in F2 control offspring (Fig. 5b and Supplementary Fig. S6).

The most important evidence supporting multi-generational inheritance came from the cytosine methylation patterns of Gβγ signaling pathway in the sperm of F1 offspring. Highly pure (>99%) sperms isolated from the caudal epididymis of males from both F1 control and F1 LPS-P offspring were isolated for BSP validation of the ratio of methylated DNA within the promoter region of genes clustered in Gβγ signaling. The data demonstrated that the cytosine methylation patterns of Sos1, Sos2, Gng10, and Prkacb in the sperm of PIE F1 LPS-P offspring were similar to those found in the thoracic aorta of F2 LPS-P offspring (Fig. 5c and Supplementary Fig. S7). These data suggested that transgenerational inherited reduced frequency of DNA methylation in the promoter region of genes clustered in Gβγ signaling might play a critical role in the development of multi-generational inherited hypertension.

Activation of PI3K-Akt signaling, downstream of Gβγ signaling, is implicated in the enhanced vasoconstrictor response in F2 LPS-P offspring. Gβγ signaling pathway belongs to G protein signal transduction system that plays critical role in participating hypertension development [33, 34]. As reported, PI3KB could be activated by direct interaction of Gβγ with p110β [35, 36], a critical substrate for PI3K activation. Activation of PI3K-Akt signaling plays critical roles in the regulation of VSMC proliferation and vascular tone response (Fig. 6a) [37]. Overactivation of PI3K-Akt signaling is usually associated with vascular remodeling and pathological changes [32]. Therefore, we next determined the activity of the critical downstream effector molecules of PI3K-Akt signaling, such as phosphorylated Akt at Ser473 site (p-AktSer473) and phosphorylated S6 at Ser235/236 site (p-S6Ser235/236), which were also the distinctive substrates for mTORC1 and mTORC2, respectively [38]. The immunoblotting data showed that there was significant increased phosphorylation level of Akt and S6, but not the total protein level of Akt and S6, in thoracic aortas of F2 LPS-P offspring, as compared to those in F2 control offspring (Fig. 6b). This suggested that the activity of PI3K-Akt signaling was enhanced in PIE F2 offspring.

To further assess the role of PI3K-Akt activation in the enhanced arterial contractility of the thoracic aorta of F2 LPS-P offspring, thoracic aortic rings from F2 LPS-P and control rats were isolated and vascular reactivity was tested in the presence of PI3K inhibitor, LY294002 [21]. LY294002 treatment significantly reversed the enhanced PE-induced contractile responses in the thoracic aorta of F2 LPS-P offspring, which even showed no statistical significance with that of F2 control offspring treated with LY294002 (Fig. 6c). In addition, there was no difference of ACh-induced vasodilation of thoracic aortas between both F2 control and F2 LPS-P offspring with LY294002 treatment (Fig. 6d). These data demonstrated that PI3K-Akt signaling was critical for the enhanced PE-induced contractile responses of thoracic aortas in F2 PIE offspring.

DISCUSSION

Most of common genetic variants discovered by genomic studies affecting blood pressure account for less than 4% of the between-person variation in trait [24], indicating that environmental factors are also closely relevant in influencing the inheritance of disease risk [10]. Transgenerational epigenetic inheritance refers to that the transmission of a specific phenotype from parent to the next generation via epigenetic marks in sperm [39]. Our findings
Fig. 4  Characteristic of genomic differentially methylated regions (DMRs) in the thoracic aorta of F2 LPS-P offspring. a, b Significantly enriched Gene Ontology (GO) terms in the biologic process (BP) associated with hypomethylated DMRs (a) and hypermethylated DMRs (b) identified in aortic MeDIP-Seq data of F2 LPS-P offspring relative to control offspring. Each circle represents one gene (left) and the description of each GO item are shown (right). c, d Canonical pathway enrichment of Kyoto Encyclopedia of Genes and Genomes (KEGG) terms from David of hypomethylated DMRs (c) and hypermethylated DMRs (d). The color shade and size of each circle reflect $-\log (P$ value) and the number of genes enriched in each KEGG pathway, respectively. $n = 4$ from 4 separate litters for each group for a-d. e, f DMRs functional analysis by Ingenuity pathway analysis (IPA). Top five enriched IPA canonical pathways (e) and the most significantly enriched IPA interaction networks (f) of hypomethylated DMRs identified in aortic MeDIP-Seq data of F2 LPS-P offspring relative to control offspring. The color shade of bar reflects $-\log (P$ value) and the horizontal axe represents the number of genes enriched in each canonical pathway, respectively. Red indicates the enriched canonical pathways; blue indicates the nearest genes associated with DMRs identified in aortic MeDIP-Seq data of F2 LPS-P offspring relative to control offspring (f).
vascular tissue (unpublished data). Previous study has revealed that offspring also showed a similar overactivation of PI3K-Akt signaling in axis in participating the enhanced arterial contractility of thoracic programming of multi-generational inherited hypertension.

methylated DNA methylation might play critical roles in the fetal higher transcription of Gβγ

Application of PI3K inhibitor in the test of vascular reactivity in the offspring and the thoracic aorta of F2 PIE LPS-P descendants, thoracic aorta further con

Notably, our data showed that genes with hypomethylated DMRs were mainly enriched in pathways associated with vascular remodeling and the pathogenesis of hypertension, especially Gβγ signaling pathways in the core of IPA interaction networks. In rat model of maternal low protein diet, previous study found a significantly undermethylated proximal promoter of the Agtr1b gene, along with upregulated mRNA and protein expression of Agtr1b, in the adrenal of F1 rat offspring [42]. The above evidence suggested that undermethylated DNA methylation might play critical roles in the fetal programming of multi-generational inherited hypertension.

Interestingly, our further experimental studies confirmed the close association of DNA hypomethylation in the promoter region and downstream PI3K-Akt signaling, accompanied by overactivation of downstream PI3K-Akt signaling. Application of PI3K inhibitor in the test of vascular reactivity in the thoracic aorta further confirmed the role of enhanced Gβγ-PI3K-Akt axis in participating the enhanced arterial contractility of thoracic aorta from F2 LPS-P offspring. Our preliminary studies in F1 PIE offspring also showed a similar overactivation of PI3K-Akt signaling in vascular tissue (unpublished data). Previous study has revealed that the activation of PI3K-Akt signaling enhances the spontaneous and agonist-induced contraction in aorta of deoxycorticosterone acetate-salt hypertensive rats [43]. PI3K-Akt signaling has also been reported to mediate the sympathetic and arterial pressure effects of leptin [44]. Therefore, these data demonstrated that PIE-induced multi-generational inherited hypomethylation of genes clustered in Gβγ signaling, via activation of PI3K-Akt signaling, played a critical role in the development of hypertension in PIE descendants. Of note, we could not still exclude other potential fraction of DNA methylation locus that contribute to the development of F2 hypertension, as we do not carry out a rescue experiment of Gβγ in our study. We could not also exclude the factors that is contributed by PIE female offspring, because we only used the male offspring in this current study.

PIE could be caused not only by “hot” inflammation after pathogen infections (virus, bacterial, and parasites infection), but also by “cold” inflammation in several chronic inflammatory diseases [3, 45]. It is of great challenge to avoid infection-induced inflammation or chronic inflammatory states during pregnancy.

Therefore, exploration of the potential mechanisms related to the altered DNA methylation with tissue specificity in PIE progenies from the aspect of maternal-fetal interface or adolescent F1 offspring would be of immense research benefit. By doing this, it will provide potential specific diagnostic biomarkers and universal therapeutic strategies for the prevention of PIE-programmed transgenerational hypertension.

Conclusively, we have revealed that PIE induces a multi-generational effect on the development of hypertension, which is due to the altered DNA methylene in F2 vascular tissue. In particular, PIE-induced DNA hypomethylation of the Gβγ signaling pathway in both the F1 sperm and the thoracic aorta of F2 offspring may be responsible for the vascular remodeling and dysregulated vascular tone via affecting PI3K/Akt signaling in PIE F2 offspring. Our findings broaden the current knowledge regarding the multi-generation effect...
of the cumulative early life environmental factors on the development of hypertension. Further studies are warranted to find similar evidence in human hypertension.

ACKNOWLEDGEMENTS
This study was supported by National Natural Science Foundation of China (81773742, 81520108029), National Key Research and Development Project (2019YFA0111200, 2020YFA0113500), and Chongqing Science and Technology Commission of China (cstc2017jcyjBX0058).

AUTHOR CONTRIBUTIONS
XG, BBC, YCD, and XHL designed the project and analyzed the data. XG, WNS, and YCD wrote the manuscript. GRD, WJL, FJW, MM, BHM, YJ, PH, TTY, YFD, LS, and WG performed the animal experiments. YY performed the canonical pathways analyzing. XG and GRD performed the qRT-PCR assays and western blots. PY, JHY, YG, and WNS...
