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Poorly controlled diabetes mellitus alters placental structure, efficiency, and plasticity

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ABSTRACT

Introduction The hemochorial placenta provides a critical barrier at the maternal–fetal interface to modulate maternal immune tolerance and enable gas and nutrient exchange between mother and conceptus. Pregnancy outcomes are adversely affected by diabetes mellitus; however, the effects of poorly controlled diabetes on placental formation, and subsequently fetal development, are not fully understood.

Research design and methods Streptozotocin was used to induce hyperglycemia in pregnant rats for the purpose of investigating the impact of poorly controlled diabetes on placental formation and fetal development. The experimental paradigm of hypoxia exposure in the pregnant rat was also used to assess properties of placental plasticity. Euglycemic and hyperglycemic rats were exposed to ambient conditions (~21% oxygen) or exposed to low oxygen (0.5% to 1.5%).

Results Diabetes caused placenomegaly and placental malformation, decreasing placental efficiency and fetal size. Elevated glucose disrupted trophectoderm cell differentiation in vitro. Evidence of altered trophoblast differentiation was also observed in vivo, as hyperglycemia affected the junctional zone transcriptome and interfered with intrauterine trophoblast invasion and uterine spiral artery remodeling. When exposed to hypoxia, hyperglycemic rats showed decreased proliferation and ectoplacental cone development on gd 9.5 and complete pregnancy loss by gd 13.5. Furthermore, elevated glucose concentrations inhibited TS cell responses to hypoxia in vitro.

Conclusions Overall, these results indicate that alterations in placental development, efficiency, and plasticity could contribute to the suboptimal fetal outcomes in offspring from pregnancies complicated by poorly controlled diabetes.

INTRODUCTION

The impact of diabetes on pregnancy outcomes has become an increasing health concern.1 During pregnancy maternal glucose can cross the placenta, which results in fetal exposure to hyperglycemia during critical stages of intrauterine development.2 Increasing evidence supports the idea that pregnancy complications and poor fetal outcomes associated with diabetes could be linked to defects in placental development and function.3 Successful pregnancy in the human requires deep trophoblast cell invasion and extensive spiral artery remodeling to facilitate optimal maternal blood delivery to the developing conceptus.4 5 Numerous physiological and environmental factors can affect trophoblast cell development and, thus, the extent of trophoblast cell-mediated spiral artery remodeling.6 7 Insufficient remodeling can then lead to the development of severe pregnancy complications. Thus, investigating placental development in the context of disease is imperative to the discovery of future preventative and therapeutic strategies.

Significance of this study

What is already known about this subject?

► Pregnancy outcomes are adversely affected by diabetes mellitus.
► Successful pregnancy in the human requires deep trophoblast cell invasion and extensive spiral artery remodeling to facilitate optimal maternal blood delivery to the developing conceptus.

What are the new findings?

► Diabetes-induced placental malformation decreases placental efficiency and fetal size.
► Hyperglycemia disrupts rat trophoblast stem cell differentiation into invasive trophoblast.
► Diabetes impairs intrauterine trophoblast invasion.
► Hyperglycemia adversely affects placental adaptations to hypoxia.
► High glucose blunts trophoblast stem cell responses to low oxygen.

How might these results change the focus of research or clinical practice?

► Investigating placental development in the context of disease is imperative to the discovery of future preventative and therapeutic strategies.
In the current study, our goals were to investigate regulatory events mediating the deleterious effects of diabetes on placental development and to evaluate the impact of hyperglycemia on placental plasticity. To facilitate our investigation, we administered streptozotocin (STZ) to pregnant rats on gestation day (gd) 6.5 to induce diabetes during pregnancy. STZ was first described as a model of diabetes mellitus in 1976, due to its ability to induce nearly complete beta cell destruction. STZ is primarily considered a model of type 1 diabetes mellitus because it targets and inactivates beta cells. STZ has also been used in the context of pregnancy to study the consequences of maternal glucose intolerance on fetal outcomes. Using the STZ model, we show that intrauterine exposure to hyperglycemia alters placental morphology, efficiency, and plasticity. Thus, inadequate placental function in mothers with poorly controlled diabetes could contribute to poor fetal outcomes.

**RESEARCH DESIGN AND METHODS**

**Animal care and treatment specifications**

Holtzman Sprague-Dawley rats were housed in an environmentally controlled facility at the University of Kansas Medical Center (KUMC) with lights on from 0600 to 2000 hours and allowed free access to food and water. Female rats (8–10 weeks of age) were mated with adult male rats (>3 months of age). Mating was assessed by inspection of vaginal lavages. The presence of sperm in the vagina was designated gd 0.5. A single dose of either STZ (45 mg/kg body weight) or vehicle control solution (0.1 M citrate buffer, pH 4.2) was administered via tail vein injection on gd 6.5 so as not to negatively impact embryo implantation. In some experiments, STZ and vehicle control-treated pregnant rats were placed in a hypoxic (10.5% (vol/vol) oxygen) gas-regulated chamber (BioSpherix, Lacona, New York, USA) from gd 6.5 to 9.5 or gd 6.5 to 13.5, which represent critical windows in placentation sites, including the ectoplacental cone, labyrinth zone, and metrial gland on gd 13.5 and 18.5, as previously described.

**Blood glucose measurements**

Body weights and blood glucose were determined prior to and after treatment to ensure equal dosage and hyperglycemia, respectively. Blood glucose levels were measured from the tail vein using the OneTouch Ultra Smart blood glucose monitoring system (LifeScan, Milpitas, California, USA).

**Tissue collection**

Rat pancreas, spleen, liver, and placental tissues were collected and weighed on gd 13.5 and gd 18.5. Entire placentation sites, including the ectoplacental cone, were dissected on gd 9.5. Placentation sites were dissected into junctional zone, labyrinth zone, and metrial gland on gd 13.5 and 18.5, as previously described. Selection of gds for analysis represented three critical stages of placentation: (1) the onset of placentation, gd 9.5, (2) establishment of placental zonation, gd 13.5, and (3) robust infiltration of invasive trophoblast cells into the uterine parenchyma, gd 18.5. Placentation sites were frozen in dry-ice cooled heptane for histological and morphometric analyses or dissected placentation site compartments snap-frozen in liquid nitrogen and stored at −80°C until processing for biochemical analyses.

**Rat trophoblast stem (TS) cell culture**

Blastocyst-derived rat TS cells were cultured in basal culture medium (RPMI 1640 (Cellgro), 20% fetal bovine serum (Sigma-Aldrich), 100 μM 2-mercaptoethanol (M7522; Sigma-Aldrich), 1 mM sodium pyruvate (ThermoFisher, 11360-070), 50 μM penicillin (15140122; ThermoFisher), and 50 U/mL streptomycin (15140122; ThermoFisher) supplemented with 70% rat embryonic fibroblast-conditioned medium, fibroblast growth factor 4 (25 ng/mL; Sigma-Aldrich), and heparin (1 μg/mL; Sigma-Aldrich). To model hyperglycemia, 25 mM glucose (Sigma-Aldrich) was added to the culture medium. D-mannitol (Sigma-Aldrich) was added to the culture medium to control for osmolality. Rat TS cells were exposed to ambient or low oxygen (0.5% or 1.5% O2) tensions and 5% CO2 at 37°C in an NAPCO 8000 incubator (ThermoFisher) for 24 hours and then harvested for immunocytochemical or biochemical analyses.

**Immunohistochemistry and immunocytochemistry**

Frozen placental tissues were sectioned at a thickness of 10 μm with a cryostat. Immunohistochemical analysis was performed on placental sections by immunofluorescence detection using a pan-cytokeratin primary antibody (1:250; F3418; Sigma-Aldrich) and Alexa 488-conjugated goat-antimouse secondary antibody (1:200, A11001; ThermoFisher). Nuclei were visualized with 4',6-diamidino-2-phenylindole (Molecular Probes, Eugene, Oregon, USA). Rat TS cells were fixed with 4% paraformaldehyde (Sigma-Aldrich) for 20 minutes at room temperature. Immunocytochemical analysis was performed by immunofluorescence detection using a primary antibody against prolactin family 8, subfamily A, member 5 (PRL8A5, 1:200) followed by Alexa 568-conjugated goat-anti-rabbit secondary antibody (1:200, A11001; ThermoFisher). Images were captured on a Nikon 80i upright microscope (Nikon, Melville, New York, USA) with a Photometrics CoolSNAP-ES monochrome camera (Roper, Sarasota, Florida, USA).

**Morphometric measurements**

Depth and extent of invasion and size of placentation compartments were performed using ImageJ (National Institutes of Health (NIH)) as previously described. Briefly, the depth of intrauterine trophoblast invasion was quantified as an invasion index, defined as the distance of distal-most cytokeratin-positive cell relative to the trophoblast giant cell layer of the placenta divided by the total distance from the giant cell layer to the outer mesometrial uterine surface. The extent of intrauterine trophoblast invasion was determined as the surface area.
within the uterine mesometrial compartment containing cytokeratin-positive trophoblast cells. Junctional zone thickness was estimated as the area intensely staining with antibodies to pan-cytokeratin in placental cross sections. All morphometric measurements were acquired from a histological plane at the center of each placenta site perpendicular to the fetal interface of the placenta.

**RNA isolation, complementary DNA (cDNA) synthesis, and transcript measurements**

Total RNA was isolated from tissues using TRIzol reagent (15596018; ThermoFisher). cDNA was synthesized from 1 µg of total RNA using the High Capacity cDNA Reverse Transcription Kit (4368813; Applied Biosystems, Foster City, California, USA). Quantitative PCR (qPCR) was performed with a reaction mixture (20 µL) containing cDNA diluted five times with water and PowerSYBR Green PCR Master Mix (4367659; Applied Biosystems) using specific primer (250 nM) sequences (online supplementary table 1). Amplification and fluorescence detection were carried out using an ABI 7500 Real Time PCR system (Applied Biosystems) for 40 cycles (95°C, 10 min; 92°C, 15 s; 95°C, 15 s; 60°C, 15 s). Relative transcript expression was calculated by ΔΔCt method and normalized to 18S rRNA.

**RNA sequencing (RNA-seq)**

Transcript profiles for gd 13.5 rat junctional zone tissue isolated from control and hyperglycemic pregnancies were generated by RNA-seq as previously described. cDNA libraries from total RNA samples were prepared with Illumina TruSeq RNA preparation kits (Illumina, San Diego, California, USA). Barcoded cDNA libraries were multiplexed and sequenced with a HiSeq2000 DNA sequencer (100 bp paired-end reads) using a TruSeq 200-cycle SBS kit (Illumina) at the KUMC Genome Sequencing Facility. Reads from *.fastq files were mapped to the rat reference genome (Ensembl Rnor_5.0.78) using CLC Genomics Workbench 12.0 (Qiagen, Redwood City, California, USA). Reads from *.fastq files were mapped to the rat reference genome (Ensembl Rnor_5.0.78) using CLC Genomics Workbench 12.0 (Qiagen, Redwood City, California, USA). Reads from *.fastq files were mapped to the rat reference genome (Ensembl Rnor_5.0.78) using CLC Genomics Workbench 12.0 (Qiagen, Redwood City, California, USA). Reads from *.fastq files were mapped to the rat reference genome (Ensembl Rnor_5.0.78) using CLC Genomics Workbench 12.0 (Qiagen, Redwood City, California, USA). The datasets generated and analyzed during the current study are available in the Gene Expression Omnibus website (https://www.ncbi.nlm.nih.gov/geo/; accession no GSE144276). All data generated and analyzed during this study are included in the published article and the online supplementary files. Resources generated and analyzed during the current study are available from the corresponding authors on reasonable request.

**Western blotting**

Placental tissues were homogenized in radioimmunoprecipitation assay lysis buffer (sc-24948A; Santa Cruz Biotechnology, Dallas, Texas, USA) supplemented with Halt Protease and phosphatase inhibitor cocktail (78443; ThermoFisher). Protein concentrations were determined by the DC protein assay (Bio-Rad, Hercules, California, USA). A total of 50 µg of protein per reaction sample were separated on 4%–20% ExpressPlus PAGE Gels (M42012, M42015; GenScript, Piscataway, New Jersey, USA), transferred to polyvinylidene fluoride blotting membrane (10600023; GE Healthcare). Following transfer, membranes were blocked in 5% non-fat milk in Tris-buffered saline with 0.1% Tween 20, for non-specific binding and subsequently probed with specific primary antibodies to prolactin family 3, subfamily d, member 4 (PRL3D4, 1:50026), PRL8A5 (1:50046), actin, beta (ACTB, 1:4000, A1978; Sigma-Aldrich), and glyeraldehyde-3-phosphate dehydrogenase (1:3000, ab9485; Abcam, Cambridge, Massachusetts, USA). Immunoreactive proteins were visualized by Luminata Crescendo Western HRP substrate (WBLUR0500; Millipore, Billerica, Massachusetts, USA) according to the manufacturer’s protocol.
Figure 1  Diabetes-induced placental malformation decreases placental efficiency. (A) Schematic depicting the experimental design for streptozotocin (STZ) treatment of pregnant rats. Vehicle control solution (0.1 M citrate buffer, pH 4.2) or STZ (45 mg/kg) was injected into pregnant rats on gestation day (gd) 6.5. Rats were subsequently sacrificed for tissue harvest on gd 13.5 or 18.5. (B) Blood glucose was measured on gd 13.5 and 18.5. STZ treatment (gray) significantly increased blood glucose levels compared with vehicle control-treated (black) animals (n=6 per group, p<0.0001). (C) The average number of viable conceptuses was no different between euglycemic and hyperglycemic animals on gd 18.5 (n=6 per group, p>0.05). (D) Placental and fetal weights (g) were measured at the time of sacrifice on gd 18.5. Hyperglycemia significantly increased placental weight (n=15 per group; p<0.001), but significantly decreased fetal weight compared with euglycemia (n=15 per group; p<0.0001). The ratio of fetal to placental weight (g), a measure of placental efficiency, was significantly decreased in animals with hyperglycemia (n=15 per group; p<0.0001). (E) Placental sites collected from euglycemic and hyperglycemic pregnant dams on gd 18.5 were sectioned and stained with a cytokeratin-specific antibody (green) for immunohistochemical analysis and visualization of trophoblast cells and placental compartments. (F) Junctional zone (JZ) and labyrinth zone (LZ) areas (pixels) were quantified using ImageJ (National Institutes of Health). Placentae collected from dams with hyperglycemia had significantly increased JZs (n=10 Eug, n=14 Hyp; p<0.001) and LZs (n=10 Eug, n=14 Hyp; p<0.0001) compared with euglycemic rats.

Increased placental weight (p<0.001) and significantly decreased fetal weight (p<0.0001; figure 1D). Fetal to placental weight ratio, a measure of placental efficiency, was significantly decreased in hyperglycemic rats compared with the euglycemic control (p<0.0001; figure 1D). No significant sex differences were observed in placental weight, fetal weight, or the fetal to placental weight ratio (online supplementary figure 2) on gd 18.5. Taken together, these results suggest that exposure to STZ-induced hyperglycemia,
a hallmark of poorly controlled diabetes, led to placental deficits and, as a result, diminished fetal size.

**Expanded junctional zone contributes to increased placental size**

To further identify the specific placental malformations resulting from diabetes exposure, placentation sites were harvested and their histological organization were assessed using immunofluorescence staining with a pan-cytokeratin antibody. The pan-cytokeratin antibody preferentially detects trophoblast cells within the junctional zone of the placenta (figure 1E). In contrast, trophoblast-associated cytokeratin expression in the labyrinth zone is diminished relative to the junctional zone because of its substantial mesenchymal cell content. Using this immunohistochemical approach, junctional and labyrinth zone areas were quantified. Area quantification confirmed that both junctional and labyrinth zones are significantly expanded in placentas from hyperglycemic pregnancies compared with euglycemic pregnancies (p<0.001, p<0.0001; figure 1F). Expanded regions of low cell density, corresponding to a greater number of glycogen cells, fill the junctional zones of placentation sites from hyperglycemic rats and contribute to the observed increase in junctional zone area.

**Diabetes alters junctional zone gene expression profiles**

To gain insights into molecular mechanisms associated with the expanded junctional zone of placentas from dams with poorly controlled diabetes, we performed RNA-seq to assess transcriptomic profiles from euglycemic and hyperglycemic pregnant rats on gd 13.5 (n=4/group). By gd 13.5, placental zonation is established, but infiltration of invasive trophoblast cells into the uterine parenchyma has yet to occur. Analysis of the RNA-seq data identified 89 significantly upregulated transcripts and 319 significantly downregulated transcripts in the hyperglycemic group compared with the euglycemic group (figure 2A). Pathway analysis encompassing all significantly altered transcripts identified several cell processes implicated in junctional zone changes in response to uncontrolled diabetes. Inflammation, cell death, and glucose metabolism disorder were highly associated with the upregulated transcripts (figure 2A). Conversely, significantly downregulated transcripts in the hyperglycemia group included inflammatory response, cellular growth and proliferation, macronutrient metabolism, cell signaling, as well as cell survival and differentiation. Several significantly upregulated and downregulated genes were validated by reverse transcription quantitative polymerase chain reaction (RT-qPCR) (figure 2B). Among the differentially expressed transcripts were two prolactin family...
members, Prl3d4 and Prl8a5, which are known markers of junctional zone development.\textsuperscript{16} PRL3D4 protein levels were quantified in junctional zone tissue collected from euglycemic and hyperglycemic pregnancies on gd 13.5 and gd 18.5 by western blotting. PRL3D4 was significantly decreased in junctional zone tissues from hyperglycemic pregnancies at both gd 13.5 and 18.5 (gd 13.5, p<0.0001; gd 18.5, p<0.01; online supplementary figure 3). Diminished PRL3D4 expression was also observed within junctional zone compartments of placenta site cross sections from hyperglycemic pregnancies (online supplementary figure 4). Overall, the findings indicate that poorly controlled diabetes induces dysregulation of junctional zone development.

Elevated glucose disrupts rat TS cell differentiation

One of the primary manifestations of diabetes, and a phenomenon we observe in our rat model, is hyperglycemia. To determine if elevated glucose impacts TS cell behavior, we cultured rat TS cells in 5 mM glucose (approximating a euglycemic state) or 25 mM glucose (approximating a hyperglycemic state) and assessed the ability of TS cells to undergo cell differentiation, focusing on junctional zone transcripts affected by hyperglycemia in vivo (figure 2B). Transcript levels were used to determine the extent of differentiation throughout a 16-day differentiation time course. In 5 mM glucose, rat TS cells begin to induce expression of genes indicative of trophoblast cell differentiation (Alox15, Prl3d4, Tpbpa, Taf7b) with increasing transcript levels observed until day 16 (figure 3A). Under TS cell-differentiating conditions in the presence of elevated glucose (25 mM), Alox15 is induced to a significantly greater extent on days 12 and 16 of differentiation compared with culture in 5 mM glucose exposure (figure 3A). In contrast, rat TS cells differentiated in 25 mM glucose exhibit significantly decreased expression of other differentiation-associated transcripts at day 16 of culture, including Prl3d4, Plac1, Tpbpa, Taf71, and Prl8a5 (figure 3A). A similar inhibition of PRL8A5 and PRL3D4 protein expression was also observed in 25 mM glucose-exposed TS cells (figure 3B,C; online supplementary figure 4). Overall, the results indicate that elevated glucose impairs TS cell differentiation and parallel in vivo placental responses to hyperglycemia.

Diabetes impairs intrauterine trophoblast invasion

The junctional zone is prominently altered in our rat model of poorly controlled diabetes (figures 1 and 2) and represents the site of progenitors for invasive trophoblast lineage development within the rat placenta.\textsuperscript{5,21} Therefore, we sought to determine whether hyperglycemia impacts development of the invasive trophoblast lineage in vivo. To assess this, we quantified the depth and extent of cytokeratin-positive trophoblast cell invasion into the uterus and measured transcript levels of invasive trophoblast-specific markers (Prl5a1, Prl7b1, Plac1) in dissected metrial glands, the site of intrauterine trophoblast cell invasion.

Quantitation of the depth of trophoblast invasion by cytokeratin staining in euglycemic and hyperglycemic pregnant rats was assessed on gd 18.5. Endovascular trophoblast invasion depth was similar in euglycemic and hyperglycemic pregnant rats (figure 4B). However, interstitial trophoblast invasion was almost completely absent on gd 18.5 in hyperglycemic rats compared with euglycemic rats (figure 4A). Total area of trophoblast invasion, including endovascular and interstitial, was decreased in hyperglycemic compared with euglycemic pregnant rats (figure 4C). Prl5a1 and Prl7b1 are specific markers of intrauterine invasive trophoblast cells,\textsuperscript{22,23} and Plac1 has recently been implicated as an intrinsic regulator of the invasive trophoblast lineage.\textsuperscript{24} Prl5a1, Prl7b1, and Plac1 transcript levels were significantly decreased within the uterine–placental interface of hyperglycemic versus euglycemic pregnancies (figure 4D). Collectively, the results indicate that hyperglycemia adversely affects development of the invasive trophoblast lineage.

Hyperglycemia and placenta plasticity

A fundamental property of the placenta is its plasticity.\textsuperscript{25} The placenta possesses the ability to adapt to a wide range of environmental challenges that could threaten pregnancy.\textsuperscript{6,25} Adjustments in trophoblast–uterine spiral artery interactions are a key component of placental adaptations to environmental stressors.\textsuperscript{4,26} Hypoxia has proven to be an effective tool to elicit adaptations in the development of the maternal–fetal interface.\textsuperscript{17,27} Since hyperglycemia affected both junctional zone and invasive trophoblast cell lineage development, we sought to determine the impact of hyperglycemia on hypoxia-induced placental plasticity.

We used an experimental paradigm of hypoxia exposure in the pregnant rat to prove that endothelial trophoblast invasion and trophoblast-guided uterine spiral artery remodeling.\textsuperscript{17} Euglycemic and hyperglycemic rats were exposed to ambient conditions (~21% oxygen) or hypoxia (10.5% oxygen) beginning on gd 6.5 and sacrificed on gd 13.5 (figure 5A). The extent of STZ-induced hyperglycemia was similar in ambient and hypoxia exposed pregnant rats (figure 5B). Hypoxia exposure significantly decreased uterine + conceptus weight in hyperglycemic compared with euglycemic pregnant rats at gd 9.5 (p<0.01; figure 5A) but did not adversely affect conceptus viability. Conceptuses from euglycemic pregnant rats exposed to ambient or hypoxic conditions and conceptuses from hyperglycemic pregnant rats exposed to ambient conditions survived to gd 9.5 (figure 5D). However, by gd 13.5 all hyperglycemic rats exposed to hypoxia showed complete pregnancy loss (p<0.0001; figure 5E). These findings prompted examination of conceptus development in each group at an earlier stage of gestation (gd 9.5). Histological analysis of gd 9.5 conceptuses demonstrated that development of the primordial placenta, referred to as the ectoplacental cone, was adversely affected in hyperglycemic pregnancies exposed to hypoxia (figure 5F), which was verified.
Figure 3   High glucose modulates rat trophoblast stem cell (TS) differentiation. (A) Rat TS cells were cultured in differentiation medium containing 5 mM glucose (black) or 25 mM glucose (gray) for 16 days. RNA was isolated from both treatment groups at five time points (days 0, 4, 8, 12, and 16). Significant differences are denoted by asterisks (n=6 per group; *p<0.05, **p<0.01, ****p<0.0001). Western blot depicting PRL8A5 (B) protein levels in rat TS cells cultured in differentiation medium containing 5 mM glucose (black) or 25 mM glucose (gray). Protein was collected from both treatment groups at five time points (days 0, 4, 8, 12, and 16). PRL8A5 protein levels were quantified with ImageJ (National Institutes of Health), normalized to beta actin (ACTIN), and graphed as “Relative Protein Abundance” (y-axis; n=5 per group). (C) Immunocytochemical analysis of PRL8A5 (red) expression in rat TS cells cultured in differentiation medium containing 5 mM glucose or 25 mM glucose for 16 days. 4',6-diamidino-2-phenylindole (DAPI, blue) marks cell nuclei.

by the quantification of ectoplacental cone surface area (p<0.0001; figure 5G) and diminished proliferation, as measured by phosphohistone H3 immunoreactivity (p<0.0001; figure 5H). Thus, hyperglycemia adversely affects placental adaptations to hypoxia.

High glucose blunts TS cell responses to low oxygen

To determine whether the interaction of hyperglycemia and hypoxia was directly altering trophoblast lineage development, we investigated the impact of glucose on TS cell responses to low oxygen in vitro. Transcript levels of known TS cell hypoxia-responsive genes, including IL33 and CD200, were measured by RT-qPCR. \(^\text{27}\) IL33 and CD200 transcript concentrations were similar in TS cells cultured in 5 mM or 25 mM glucose exposed to ambient conditions (p<0.01; online supplementary figure 5). In contrast, exposure to low oxygen significantly induced expression of IL33 and CD200 in 5 mM glucose (p<0.01) but IL33 and CD200 were not induced by hypoxia in 25 mM glucose (p<0.01). We also examined the interactive effects of glucose and hypoxia on TS cell number. TS cell viability was not altered by the culture conditions (figure 5I); however, low oxygen stimulated TS cell proliferation in 5 mM glucose, but not in 25 mM glucose (p<0.0001; figure 5I). These in vitro results support the
Figure 4  Diabetes impairs uterine trophoblast invasion and spiral artery remodeling. (A) Placenta sites collected from euglycemic (Eug) and hyperglycemic (Hyp) rats on gestation day (gd) 18.5 were sectioned and stained with a cytokeratin-specific antibody (green) for immunohistochemical analysis and visualization of trophoblast cells and placental compartments. Brightly labeled, cytokeratin-positive trophoblast cells invade interstitially and endovascularly into the metrial gland compartment. Images were acquired with a 10× objective. The scale bars represent 250 µm. (B) Depth of intrauterine trophoblast cell invasion was quantified in ImageJ (National Institutes of Health (NIH)) as an invasion index, or the distance of distal-most endovascular cytokeratin-positive cell relative to the trophoblast giant cell layer of the chorioallantoic placenta. Invasion depth was similar between Eug and Hyp pregnant rats (n=10 Eug, n=9 Hyp; p>0.05). (C) Total area of trophoblast invasion, including endovascular and interstitial, was quantified in ImageJ (NIH). Hyp significantly reduced interstitial, but not endovascular, trophoblast invasion compared with Eug (n=10 Eug, n=9 Hyp; p<0.001). (D) RNA was isolated from uterine-placental interface tissue dissected from Eug and Hyp pregnant dams on gd 18.5. Prl5a1, Prl7b1, and Plac1 transcripts were significantly downregulated in the uterine-placental interface collected from Hyp animals (gray) compared with Eug animals (black; n=6 per group, *p<0.05, **p<0.01).

in vivo observations that diabetes (hyperglycemia) alters placental cell plasticity through, at least in part, direct actions on trophoblast cell development.

DISCUSSION
Animal models have been effectively used to investigate diabetes and pregnancy. In this study, we examined the impact of poorly controlled diabetes on placental and fetal development in the rat. We provide evidence that placental development in a hyperglycemic state is not optimal. Hyperglycemia leads to the formation of an abnormally large, inefficient placenta that adversely affects fetal growth and that does not adapt well to an adverse maternal environment. Our results suggest that some of the effects of hyperglycemia on placentation are due to the direct actions of high glucose on trophoblast cell behavior.

In this report, we have not attempted to mimic a specific human syndrome or disease state with the rat. Fundamental differences between rat and human pregnancy render such a task highly problematic. Placenta-related events that transpire during rodent gestation best equate to physiological changes that occur during the first trimester of human pregnancy. From a superficial
Pathophysiology/Complications

Figure 5  Hyperglycemia diminishes conceptus viability and ectoplacental cone development (EPC) in pregnant rats exposed to hypoxia. (A) Schematic depicting the experimental design for streptozotocin (STZ) treatment and hypoxia exposure of pregnant rats. Half of the rats were maintained in ambient air housing and the other half were housed in a gas-regulated chamber to mimic hypoxia (10.5% (vol/vol) oxygen) until sacrifice on gestation day (gd) 9.5 or 13.5 for tissue harvest. Photomicrographs of whole uteri collected from euglycemia (Eug) and hyperglycemia (Hyp) animals housed in ambient air or hypoxia. (B) Glucose was measured on gd 13.5 in rats housed in ambient air and hypoxia. Hyp (gray) possessed significantly increased blood glucose levels compared with Eug (black) animals in both ambient air and hypoxia (n=6 for all groups except n=8 for Hyp in hypoxia, p<0.0001). (C) Uterine weight was significantly reduced in Hyp animals compared with Eug animals when exposed to hypoxia n=5 for ambient and n=6 for hypoxia, p<0.01). (D) The average number of viable conceptuses was not different between Eug (black) and Hyp (gray) animals housed in ambient air or hypoxia on gd 9.5 (n=5 for ambient and n=6 for hypoxia, p>0.05). (E) Viable conceptuses were not recovered in Hyp animals housed in hypoxia on gd 13.5, resulting in a significant difference between treatment groups (n=6 per group, p<0.0001). (F) Placentation sites were collected on gd 9.5, sectioned, and stained for cytokeratin (green) and 4’6-diamidino-2-phenylindole (DAPI, blue) from Eug and Hyp rats housed in ambient air or hypoxia (10.5% O2). (G) EPC size, quantified as relative area of cytokeratin-positive cells, is significantly decreased in Hyp animals (gray) compared with Eug animals (black) (n=13 per group, p<0.0001). (H) Hyp significantly decreased EPC cell proliferation, which was measured as the number of phospho-histone H3 positive cells (n=6 per group, p<0.0001). (I) Rat trophoblast stem cells were cultured in medium containing 5 mM glucose (black) or 25 mM glucose (gray) and were maintained in either ambient air or exposed to low oxygen (1.5% O2) for 24 hours and the number of viable cells quantified. Significant differences are denoted by asterisks (n=6 per group, p<0.0001).

Perspective, placentation appears to be different in the rat and human; however, this view is biased by the use of distinct, species-specific nomenclature. In fact, rat and human placentas are highly conserved. Both species possess hemochorial placentas that perform the same critical tasks. Central to those tasks are specialized trophoblast cells that transform the uterine environment, including restructuring the uterine spiral arteries, to regulate delivery of maternal nutrients to the developing fetus. Although the hyperglycemia we observe with our model of STZ-induced beta cell damage may best reflect poorly controlled type 1 diabetes in pregnancy, the goal of the experimentation outlined in this report was to gain broader insights into the impact of glucose dysregulation on the morphogenesis of the placenta.

Disruptions in placental structure are a common feature of hyperglycemia. Our observations of placentomegalia are consistent with earlier reports examining the effects of diabetes mellitus on placentation development in the mouse and rat. An expansion of glycogen cells within the junctional zone compartment has been a common finding. Diabetes in humans is associated with alterations in placental surface area, volume, intervillous space, terminal villi, syncytiotrophoblast, fibrinoid areas, and glycogen deposits. A key element of human placentation is the entry of trophoblast cells into the uterus where they restructure the uterine parenchyma, including remodeling of uterine spiral arteries. This form of deep placenta is also observed in the rat. Two invasive trophoblast cell populations are evident. Interstitial invasive trophoblast cells are situated between the uterine vasculature, whereas endovascular invasive trophoblast cells replace the endothelium of the uterine spiral arterioles.
our experimentation, we observed that hyperglycemia preferentially led to disruptions in the interstitial invasive trophoblast population. Impairments in intrauterine trophoblast cell invasion have also been observed in the biobreeding diabetes-prone rat model.36 There is also some indication that trophoblast cell invasion is adversely affected in humans with diabetes.29 40 Specific mechanisms underlying invasive trophoblast cell reorganization of the uterine parenchyma is not well appreciated. Furthermore, we know little of how hyperglycemia affects differentiation of the invasive trophoblast cell lineage.

Some insights into the effects of hyperglycemia on placentation may be gleaned from the transcript profile of junctional zone placental tissue from euglycemic and hyperglycemic rat pregnancies. In addition to the dysregulation of species-specific members of the prolactin family (Prl3d4, Prl8a5, Prl8a9), transcripts from the pregnancy-specific glycoprotein family (Cgm4), and the cathepsin family (Cts7), were also identified as dysregulated in response to hyperglycemia. Downregulated transcripts, such as Taf7l, Pappa2, and Plac1, could contribute to hyperglycemia-induced placental dysmorphogenesis. Taf7l encodes a member of a family of proteins that associate with the TATA-binding protein (TBP) and is essential for TBP binding to the TATA element and transcription.41 TAF7L has been implicated in sperm development and the regulation of adipocyte differentiation,42 43 but not placentation. Pappa2 encodes a protease targeting insulin-like growth factor (IGF)-binding proteins44 which modulate the activity of IGFs,45 known regulators of placental growth.46 However, disruption of Pappa2 in the mouse has only subtle effects on male reproduction and modest effects on postnatal growth with no reported adverse effects on placental development.47 48 In contrast, placenta specific 1 (Plac1), a highly conserved X-linked gene, possesses pronounced effects on placentaion.49 50 Targeted disruption of Plac1 in the mouse leads to placentomegaly, including an expanded junctional zone.49 50 Downregulation of Plac1 expression in the placenta associated with hyperglycemia fits well with the phenotype of the Plac1 null mouse. Thus, PLAC1 represents a candidate mediator of the effects of hyperglycemia on placentation.

A healthy placenta exhibits plasticity and can adapt to a range of environmental challenges.6 25 Hypoxia is an environmental challenge that can elicit placental adaptations, including the stimulation of trophoblast invasion and spiral artery remodeling.17 Hypoxia is also a physiological regulator of placentation51 and has been identified as a key regulator of invasive/extravillous trophoblast cell differentiation.18 27 54 Dual exposure to hyperglycemia and hypoxia in the rat was not compatible with pregnancy. Pregnancies were terminated due to growth arrest at the ectoplacental cone stage of placentation. Cellular responses to low oxygen were blunted when TS cells were cultured in high glucose implicating direct interactive effects of oxygen tension and glucose on trophoblast cells. High altitude is associated with low oxygen tensions and increased risks for pregnancy complications.52 The combination of high altitude and diabetes may be particularly problematic to the development of an effective hemochorial placenta.

In summary, our data generated using an in vivo STZ model of diabetes and in vitro analyses of TS cells support the hypothesis that exposure to hyperglycemia during gestation induces placentation alterations that result in impaired fetal outcomes. The initiation of hyperglycemia post-embryo implantation was a key element of our experimental design and distinct from other animal models of diabetes.28 Our findings strengthen earlier observations that intrauterine exposure to hyperglycemia alters structural formation and glycogen cell accumulation within the placenta. New insights into the impairment of trophoblast differentiation and invasion in placentation sites from dams with hyperglycemia highlights a potential mechanism by which fetal growth restriction may occur in this model. We also found that placentas from hyperglycemic pregnancies lacked the requisite plasticity to adapt to the stress of hypoxia exposure and led to pregnancy loss. These findings reinforce the damaging effects of poorly controlled diabetes on placental formation and pregnancy outcomes as well as the experimental value of the rat model in the investigation of hemochorial placentation and critical events regulating trophoblast invasion and spiral artery remodeling.

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Competing interests MJS is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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Data availability statement Data are available in a public, open access repository. Data are available on reasonable request. The datasets generated and analyzed during the current study are available in the Gene Expression Omnibus website (https://www.ncbi.nlm.nih.gov/geo/) accession no. GSE414276. All data generated and analyzed during this study are included in the published article and the online supplementary files. Resources generated and analyzed during this study are available from the corresponding authors on reasonable request.

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