Fetal exposure to a diabetic intrauterine environment resulted in a failure of cord blood endothelial progenitor cell adaptation against chronic hypoxia

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Abstract: Gestational diabetes mellitus (GDM) has long-term health consequences, and fetal exposure to a diabetic intrauterine environment increases cardiovascular risk for her adult offspring. Some part of this could be related to their endothelial progenitor cells (EPCs). Understanding the vessel-forming ability of human umbilical cord blood (HUCB)-derived endothelial colony-forming cells (ECFCs) against pathological stress such as GDM response to hypoxia could generate new therapeutic strategies. This study aims to investigate the role of chronic hypoxia in EPCs functional and vessel-forming ability in GDM subjects. Each ECFC was expressed in endothelial and pro-angiogenic specific markers, namely endothelial nitric oxide synthase (eNOS), platelet (PECAM-1) endothelial cell adhesion molecule 1, vascular endothelial-cadherin CdH5 (Ca-dependent cell adhesion molecule), vascular endothelial growth factor A, (VEGFA) and insulin-like growth factor 1 (IGF1). Chronic hypoxia did not affect CdH5, but PECAM1 MRNA expressions were increased in control and GDM subjects. Control hypoxic and GDM normoxic VEGFA MRNA expressions and hypoxia-inducible factor 1-alpha (HIF1a) protein expressions were significantly increased in HUCB ECFCs. GDM resulted in most failure of HUCB ECFC adaptation and eNOS protein expressions against chronic hypoxia. Chronic hypoxia resulted in an overall decline in HUCB ECFCs’ proliferative ability due to reduction of clonogenic capacity and diminished vessel formation. Furthermore, GDM also resulted in most failure of cord blood ECFC adaptation against chronic hypoxic environment.

Keywords: endothelial progenitor cells, gestational diabetes mellitus, chronic hypoxia, human cord blood

Introduction
Gestational diabetes mellitus (GDM) has long-term health consequences for her adult offspring.1,2 GDM demonstrates higher prevalence of obesity, insulin resistance, type 2 diabetes mellitus (DM) (during adolescence), development of metabolic syndrome (in early childhood), cardiovascular disease, and dyslipidemia in her offspring.1,2 Some of them could be related to decreased endothelial cells (ECs) and their progenitors.

Until 1997, it was believed that the differentiation of mesodermal cells into angioblasts and endothelial differentiation occur during embryonic development.3,4 When Asahara et al isolated endothelial progenitor cells (EPCs) from human peripheral blood, they started a predominant paradigm about new vessel formation in adult subjects.4 After they isolated
hematopoietic progenitor cells from adults, they demonstrated that those progenitor cells can actually differentiate into ECs.\textsuperscript{5,6}

The role of the endothelium is to receive, integrate, and respond against systemic and local environments.\textsuperscript{7} ECs and their progenitors, which form the vascular endothelium, play a key role in sensing and responding to different physiological and pathological stresses such as hypoxia.\textsuperscript{4,9} This can be described as a compensatory mechanism. During this period, a larger amount of EPCs are released and mobilized into the peripheral blood and cause vasodilation and increased vessel formation. On the other hand, chronic hypoxia mostly exhibits apoptosis, with the cells unable to survive.

Adult peripheral blood and cord blood contain EPCs, which represent a minor subpopulation of blood mononuclear cells (MNCs). EPCs also contain different subpopulations such as colony-forming unit ECs (CFU-ECs) and endothelial colony-forming cells (ECFCs). They exhibit different functional properties.\textsuperscript{6,8} Decrease in CFU-EC colonies is oppositely correlated with hypercholesterolemia, hypertension, and type 2 DM.\textsuperscript{6,9} On the other hand, ECFCs are associated with angiogenesis and apoptosis.\textsuperscript{6,8} The number of ECFCs and their functional capacity determine de novo vessel formation and coordinate the physiological and pathological vessel-forming ability, including cell survival, proliferation, migration, and capillary-like structure formation in ECs. ECFCs directly contribute to the re-endothelization process, which plays an important role in the maintenance of the endothelium.\textsuperscript{10–12} Importantly, they have the capacity to repair the injured endothelium and the ability to promote vessel formation.\textsuperscript{11–13} Especially during acute hypoxia, a large number of EPCs mobilize into peripheral blood and they directly contribute to the revascularization process by a series of reactions.\textsuperscript{14–16}

This project mainly designed to investigate the adaptation ability and/or deficiency of EPCs against long-term hypoxia and a fetal diabetic environment. It clarifies whether fetal exposure to a diabetic intrauterine environment would result in EPCs’ adaptation deficiency against a hypoxic environment. Collectively, it illuminates the role of hypoxia on the development of apoptosis, and gives beneficial information related to future therapeutic modalities.

Research design and methods

Clinical data, cord blood collection, and human blood samples

This work was undertaken according to the Declaration of Helsinki based on the primary ethical obligations of subjects, especially to collect human cord blood samples. Pregnant subjects from uncomplicated pregnancies were identified during outpatient clinics and/or delivery (previously approved by the Institutional Review Board (IRB), Department of Pediatrics, Indiana University School of Medicine).

Maternal data

GDM subjects demonstrated symptoms of diabetes during their pregnancy plus high blood glucose (BG) concentration (BG ≥11.1 mM [200 mg/dL], or fasting plasma glucose ≥7.0 mM [126 mg/dL], or 2-hour plasma glucose ≥11.1 mM [200 mg/dL] during an oral glucose tolerance test). HgbA1Cs 39 mmol/mol 117 mg/dL estimated average glucose [eAG]).

ECFC isolation

Around 50 mL blood was collected from the cord blood (from IRB-approved patients) of freshly delivered placentas by a research nurse or trained research personnel. Mononuclear cells (MNCs) were isolated as described previously.\textsuperscript{8,17} MNCs were washed three times with EBM-2 medium (Cambrex, Walkersville, MD, USA) supplemented with 10% fetal bovine serum (FBS; Hyclone, Logan, UT, USA), 2% penicillin/streptomycin, 5 mM dextrose, and 0.25 µg/mL amphotericin (Thermo Fisher Scientific, Waltham, MA, USA) (complete EBM-2 medium) and seeded onto six-well tissue culture plates precoated with collagen. After 24 hours, the adherent cells were washed and the medium changed daily until first ECFC colonies appeared between 5 days and 11 days, as previously described.\textsuperscript{8,17} The number of colonies was counted for each cord blood sample. The colonies were trypsinized and replated in a flask precoated with collagen. The cells were passaged upon reaching 80%–90% confluence. Early passage (<5) ECFC cell lines were used for the present study. For that purpose, five controls and five GDM human umbilical cord blood (HUCB) ECFCs were selected.

Long-term hypoxia

Early passage (<5) HUCB ECFCs obtained from GDM (n=5) and control (n=5) subjects were cultured with plates precoated with collagen under hypoxic condition (5% CO\textsubscript{2}/94% N\textsubscript{2} and 1% O\textsubscript{2} in a triple-gas incubator)\textsuperscript{7,18} as well as under normoxic condition\textsuperscript{17} for 7 days using EGM, Medium with 10% FBS.

Characterization of HUCB ECFCs and real-time polymerase chain reaction

HUCB ECFCs were identified with previously described classical cell culture methods.\textsuperscript{8,17} Endothelial and
pro-angiogenic-specific primers were designed to identify their endothelial origin and angiogenic capacity for real-time polymerase chain reaction (PCR): endothelial specific markers: platelet endothelial cell adhesion molecule (PECAM, CD31), vascular endothelial-cadherin (Cdh5), Ca-dependent cell adhesion molecule (CdH5), endothelial nitric oxide synthase (eNOS), pro-angiogenic specific markers: vascular endothelial growth factor A (VEGFA) and insulin-like growth factor 1 (IGF-1).

Total ribonucleic acid (RNA) was extracted according to manufacturer’s instructions (SV total RNA isolation system [Promega Corporation, Fitchburg, WI, USA]). At the end of the isolation, RNA samples were dissolved in nuclease-free water (pH 7.5). RNA samples were used as templates for the synthesis of first-strand cDNAs, as described previously.19,20 The total RNA quantity and quality were determined using the Experion™ semiautomated electrophoresis system (Bio-Rad Laboratories Inc., Hercules, CA, USA). In the Experion priming station, the microfluidic-based LabChip sample wells (RNA StdSens Chips, Experion, Bio-Rad Laboratories Inc.) were filled with a polymer-sieving matrix containing a fluorescent dye and 1 µL of denatured total RNA.21 The concentration of total RNA was determined by using the ratio of the sample RNA area to the Experion RNA ladder area. RNA samples with distinct 18S and 28S ribosomal RNA fragments in the electropherogram were used to assess the RNA quality.21 Selected good-quality RNA samples were used as templates for the synthesis of first-strand cDNAs.21

Real-time PCR reactions were performed for PECAM1, VE-cadherin, eNOS, VEGFA, IGF-1, and β-actin duplicated with a custom-designed SYBR green mix [2.4 µL of 25 mM MgCl₂, 5 µL of 1:10,000 dilution SYBR Green I (Molecular Probes), and 5 µL of 1 mM fluorescein calibration dye (1 mM/L in DMSO, Bio-Rad Laboratories Inc.)] in 50 µL of total reaction using Taq DNA polymerase (Promega).19,20 Primers were designed using the Genomatix and the National Center for Biotechnology Information database. The primer sequences used for transcripts were as follows: PECAM1 (CD31) sense GAGATGAGTGATGGCTGGA and antisense TGCTGCAATGAAGTGTTCT (accession number AH002931); VE-cadherin (CDH5) sense CCTACCAGCCCAAGTTGT and antisense GAGATGCCACGGGTAGAA (accession number NM_001795); and eNOS sense ACCCTACCGCTACACATC and antisense GCTATTCTCCAGGTGCTTC (accession number NM_000603). VEGFA primers (sense CCAACTTCTGGGCTTTTCTC and antisense CCCCTCCTCCTCTTCTTCTC) were designed with the GENOMATIX program based on previously published accession numbers (BC011177, NM_001025366, NM_003376, NM_001025367, NM_001025368, NM_001025369, NM_001025370, NM_001033756). IGF-1 primers (sense CATGTCCCTCGCATCCTC and antisense ATACCCTGTGGCTGTGTTGA) were designed with the Genomatix program based on previously published accession numbers (NM_00618, NM_00111283, NM_001111284, NM_001111285). β-Actin primers (sense GGACTTCGAGCAAGAGATGG and antisense AGCAGCTGTGGCGCTGAC) were designed for integral control of the experiment (accession number NM_001101).

Amplification was carried out with the iCycler iQ multicolor real-time PCR detection system (Bio-Rad Laboratories Inc.) as follows: 30-seconds denaturation (95°C) followed by 45-seconds annealing and 1-minute extension (72°C), repeated for a total of 33 cycles. β-Actin was amplified in each set of PCR reactions and served as an internal reference during quantitation to correct for operator and/or experimental variations. The data were analyzed in duplicate using the modified 2–∆∆CT equation:19–22

$$\Delta\Delta C_T = (C_T - C_{T-\beta-actin})_{\text{target}} - (C_T - C_{T-\beta-actin})_{\text{control}}$$

The mean threshold cycle (Cₜ) values for target and internal control (β-actin) genes were determined in each sample. On the other hand, the formula was further modified for the control and GDM samples under hypoxic and normoxic conditions using the specific gene primers as well as internal control β-actin primers in each duplicated set (Table 1).

**Western blot analysis eNOS, P-eNOS, and hypoxia-inducible factor 1-alpha (HIF1α)**

For 7 days, hypoxic and normoxic HUCB ECFCs were obtained from GDM (n=4) and control (n=4). Proteins were isolated and homogenized with lysis buffer containing protease inhibitors (50 mmol/L Tris-HCl, 1% NP-40, 150 mmol/L NaCl, 1 mmol/L EDTA, 1 mmol/L phenylmethylsulfonyl fluoride, 1 mg/mL aprotinin, 1 mg/mL leupeptin, and 1 mg/mL pepstatin). The supernatants were collected and used for analysis. Protein quantity and quality were determined using the Experion semiautomated electrophoresis system (Bio-Rad Laboratories Inc.). Basically, 4 µL of denatured protein for each sample was...
loaded into the Pro260 chip (Experion, Bio-Rad Laboratories Inc.). The Pro260 ladder used in the chip was based on protein gel electrophoresis, and contains nine proteins ranging from 1 kDa to 260 kDa as well as 1.2 kDa lower markers. Electropherograms were obtained for each group. Equivalent amounts of protein were separated by gel electrophoresis, and contains nine proteins ranging from 1 kDa to 60 kDa as well as 60 kDa lower markers. Electropherograms were obtained for each group. Equivalent amounts of protein were separated by gel electrophoresis, and contains nine proteins ranging from 1 kDa to 60 kDa as well as 60 kDa lower markers. Electropherograms were obtained for each group. Equivalent amounts of protein were separated by gel electrophoresis, and contains nine proteins ranging from 1 kDa to 60 kDa as well as 60 kDa lower markers. Electropherograms were obtained for each group. 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Senescence assay

Acidic β-galactosidase (β-gal) assay was conducted to investigate the functional capacity of EPCs against long-term hypoxia. Two thousand cells were seeded and triplicated into a well of six-well plates for each experimental group (human umbilical cord blood ECFCs obtained from control pregnancies and plated under normoxic condition [C-HUCB ECFC-N], human umbilical cord blood ECFCs obtained from control pregnancies and plated under hypoxic condition [C-HUCB ECFC-H], human umbilical cord blood ECFCs obtained from GDM subjects and plated under normoxic condition [GDM-HUCB ECFC-N], and human umbilical cord blood ECFCs obtained from GDM subjects and plated under hypoxic condition [GDM-HUCB ECFC-H]) and exposed to hypoxia (5% CO₂/94% N₂ and 1% O₂) in a triple gas incubator. After 7 days, the triple gas hypoxia cells were fixed with 2% formaldehyde and 0.2% gluteraldehyde combination. Then cells were stained with β-galactosidase solution (X-gal + β-gal; 1 mg/mL final concentration) and incubated with normoxic cell culture overnight. The next day, the blue stained cells were counted under a microscope.

Increased stained cells with β-gal contributed to the reduction of clonogenic capacity of the hypoxic samples.

Matrigel assay

After exposure to chronic hypoxia (5% CO₂/94% N₂ and 1% O₂ in a triple gas incubator for 7 days), ECFCs from each experimental group (C-HUCB ECFC-N, C-HUCB ECFC-H, GDM-HUCB ECFC-N, and GDM-HUCB ECFC-H) were seeded and triplicated at 6,000 cells/well in 96-well tissue culture plates coated with 35 μL of matrigel (BD Biosciences,

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### Table 1 The primer sequences used for real-time PCR

<table>
<thead>
<tr>
<th>(A)</th>
<th>Primers</th>
<th>Sense</th>
<th>Antisense</th>
</tr>
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<tr>
<td>eNOS</td>
<td>ACCCTCACGGCTACAACATC</td>
<td>GCTCATTTCCAGGTGCTTC</td>
<td></td>
</tr>
<tr>
<td>VE-Cadherin (CDH5)</td>
<td>CCTACAGCCCAAGAGTTGG</td>
<td>GAGATGACACCGGGTGAAGA</td>
<td></td>
</tr>
<tr>
<td>PECAM1 (CD31)</td>
<td>GAGATGAGTTAGGGCTGGA</td>
<td>TGTCGATCAGGGGTGTTTC</td>
<td></td>
</tr>
<tr>
<td>VEFA</td>
<td>CCAACTCCCTCCCTCTTCCTC</td>
<td>CCCCCTCTCTTCCTCTTC</td>
<td></td>
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<tr>
<td>IGF-I</td>
<td>CATGGTCCCTACTGACTTCTC</td>
<td>ATACCCCTGAGGCTTGTTGGA</td>
<td></td>
</tr>
<tr>
<td>beta-actin</td>
<td>GGACCTCGAGAAGAGATGG</td>
<td>AGCAGTGTGTTGGCGTACAG</td>
<td></td>
</tr>
</tbody>
</table>

**2-ΔΔCt Equation**

\[
\Delta \Delta Ct = \Delta Ct_{GDM \text{ nor}} - \Delta Ct_{\text{control nor}} = (Ct_{\text{gdm nor}} - Ct_{\text{gdm nor BA}}) - (Ct_{\text{cont nor}} - Ct_{\text{cont nor BA}})
\]

**Notes:**
- Ct<sub>gdm nor</sub>, Ct<sub>gdm nor BA</sub>: GDM samples under normoxic condition using interested gene primers; Ct<sub>cont nor</sub>, Ct<sub>cont nor BA</sub>: control samples under normoxic condition using beta-actin primers;
- Ct<sub>gdm hypx</sub>, Ct<sub>gdm hypx BA</sub>: GDM samples under hypoxic condition using interested gene primers; Ct<sub>cont hypx</sub>, Ct<sub>cont hypx BA</sub>: control samples under hypoxic condition using beta-actin primers.

**Abbreviations:**
- eNOS, endothelial nitric oxide synthase; VE-cadherin, vascular endothelial-cadherin; PeCAM1, platelet endothelial cell adhesion molecule 1; VEFGA, vascular endothelial growth factor A; IGF-1, insulin-like growth factor 1; GDM, gestational diabetes mellitus.
San Jose, CA, USA). After 6–10 hours, the vessel-forming capacity (number of closed network units) was quantified with visual microscopy (Zeiss Axiolov 2 inverted microscope with a 5x CP-Achromat/0.12 NA objective). The images were collected for each experimental group with a SPOT RT camera.

Data and statistical analysis
Differences among group values were evaluated by one-way analysis of variance followed by a Newman–Keuls test. Data are presented as means ± SE (Standard Error).

Results
Characterization of HUCB ECFCs using cell culture and real-time PCR
The HUCB MNCs were plated onto collagen-coated plates in endothelial-specific media, as previously described.6,23 When nonadherent cells were discarded, cells appeared in cobblestone formation. They proved to be phenotypically indistinguishable from cultured ECs and possessed de novo vessel-forming ability.8,9,24 Flow cytometry demonstrated the endothelial origin of human hematopoietic progenitor cells.17,25 Cells were obtained from IRB-approved (under Declaration of Helsinki) ready HUCB ECFCs from the EPC Bank, The Herman B Wells Pediatric Research Center, Department of Pediatrics, Riley Pediatric Foundation and University Pediatric Association, Indiana University School of Medicine, Indianapolis, IN, USA, previously isolated and declared as ECFC (early passage <5).

Real-time PCR results demonstrated that each HUCB ECFC expressed eNOS, PECAM1, VE-cadherin, VEGFA, and IGF-1 messenger RNA (mRNA) (Figures 1–3). All control and GDM HUCB ECFCs expressed eNOS, PECAM1 (CD31), VE-cadherin, VEGFA, and IGF-1 mRNAs under normoxic as well as hypoxic conditions.

HUCB ECFCs obtained from GDM and C subjects demonstrated increased PECAM1 (CD31) mRNA expressions with chronic hypoxic condition (Figure 1). PECAM1 (CD31) was only expressed in ECs. This indicates, first, that all samples have endothelial origin, and, second, type 1 transmembrane protein PECAM1 mRNA expressions were stimulated with chronic hypoxia. Using real-time PCR, significant differences were not detected in eNOS mRNA expression in HUCB ECFCs; on the other hand, it tended to increase in Control and GDM hypoxic samples. Significant differences were also not detected in VE-cadherin (CdH5) mRNA expressions in HUCB ECFCs with hypoxia. VE-cadherin is a Ca++-dependent cell adhesion molecule, and it indicated that chronic hypoxia did not affect Ca++-induced cell adhesion.

According to real-time PCR, C hypoxic and GDM normoxic HUCB ECFCs, VEGFA mRNA expressions were significantly increased compared to control HUCB ECFCs, (Figure 2). Chronic hypoxia triggered VEGFA mRNA expressions in HUCB ECFCs obtained from control pregnancies but did not affect the GDM samples. On the other hand, GDM itself resulted in increased VEGFA mRNA expressions. According to real-time PCR, significant differences were also detected in IGF-1 mRNA expression in HUCB ECFCs in GDM hypoxic samples when compared with control hypoxic HUCB ECFC mRNAs (Figure 3). Hypoxia alone did not cause any changes, but if long-term hypoxia was applied after diabetic intrauterine environment, IGF-1 mRNA expressions significantly increased.

Western blot analysis
eNOS, P-eNOS, and HIF 1α protein expressions were obtained from HUCB ECFCs with normoxic as well as hypoxic condition (Figure 4A–C). eNOS protein expressions were significantly decreased in HUCB ECFCs in control as well as GDM subjects plated under hypoxic condition if compared with those plated under normoxic condition. Their quantitative assessments demonstrated that exposure to a gestational diabetic environment resulted in the most significant increase in eNOS protein expressions when plated under normoxic condition (Figure 4I and IIA). On the other hand, chronic hypoxia was significantly increased in P-eNOS protein expressions obtained from control and GDM HUCB-ECFCs when compared with their normoxic samples (Figure 4I and IIB). When combined with mRNA expression data, this means that the reduction of eNOS protein expressions against chronic hypoxic environment results from posttranslational modifications of those proteins.

Chronic hypoxia resulted in increased HIF 1α protein expression in HUCB ECFCs obtained from control subjects, but decreased expression of GDM samples.

Gestational diabetic environment resulted in increased HIF 1α protein expression with regular condition when compared with control subjects. On the other hand, when chronic hypoxia was applied to GDM samples, the increase was not as high as in control subjects. The most significant increase was...
seen in control subjects (C-HUCB ECFC-N) that were plated under chronic hypoxia (Figure 4I and 4IIC).

**Senescence assay**

The photomicrographs in Figure 5 represent senescence assays, and when positively stained with senescence assay (SA) β-gal of HUCB ECFCs indicates the clonogenic capacity of those cells. HUCB ECFCs obtained from GDM and control subjects exhibited significant increase in senescence in vitro exposure to chronic hypoxia (5% CO<sub>2</sub>/94% N<sub>2</sub> and 1% O<sub>2</sub> in a triple gas incubator).<sup>7,18</sup> It shows the reduction of clonogenic capacity against a chronic hypoxic environment (Figures 5 and 7A). The most significant enhanced senescence was observed in ECFCs

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**Figure 1 (Continued)**
Figure 1 Real-time PCR reactions performed to characterize and compare hUCB eCFCs.

Notes: PECAM (CD31), VE-cadherin (CDH5), eNOS and beta-actin primers were used for real-time PCR with a duplicated, custom-designed SYBR green mix, as previously described.19,20 The PCR melt curve (I–I) and amp/cycle (I–II), (I–A I and II) are eNOS, (I–B I and II) are VE-cadherin, (I–C I and II) are PECAM 1. Real time PCR results shown in I–II; A is eNOS, B is VE-cadherin and C is PECAM 1. Each mean gene fold change is normalized with the internal control β-actin. Real-time PCR results shown in (II); C-HUCB ECFC-N represents HUCB-derived ECFCs obtained from control pregnancies and plated under normoxic condition; C-HUCB ECFC-H represents HUCB-derived ECFCs obtained from control pregnancies and plated under hypoxic condition; GDM-HUCB ECFC-N represents HUCB-derived ECFCs obtained from GDM subjects and plated under normoxic condition; GDM-HUCB ECFC-H represents HUCB-derived ECFCs obtained from GDM subjects and plated under hypoxic condition. β-Actin was amplified in each set of PCR reactions and served as an internal reference during quantitation to correct for operator and/or experimental variations. The data were analyzed in duplicate using the modified $2^{-\Delta\Delta CT}$ equation.21 (II) demonstrates that, using real-time PCR, only significant differences were detected in PECAM1 (CD31) mRNA expression in HUCB ECFCs in C hypoxic and GDM hypoxic samples when compared with C normoxic and GDM normoxic mRNAs. *C-HUCB ECFC-N is significantly different from C-hUCB eCFC-h and GDM-hUCB eCFC-h.

Abbreviations: PCR, polymerase chain reaction; HUCB, human umbilical cord blood; ECFCs, endothelial colony-forming cells; platelet endothelial cell adhesion molecule (PECAM1, CD31); VE-cadherin, (CDH5); eNOS, endothelial nitric oxide synthase; GDM, gestational diabetes mellitus.
obtained from GDM exposed to a chronic hypoxic condition (Figure 7A).

Matrigel assay

The photomicrographs represent matrigel assay, which represents the vessel-forming ability (Figure 6). A significant reduction in tube formation was observed in control and GDM HUCB ECFCs when cultured under chronic hypoxic conditions (Figure 7B). Their quantitative assessments demonstrated that exposure to long-term hypoxia exhibited impairment in HUCB ECFCs’ tube-forming ability. The most significant decrease in capillary tube formation was observed in ECFCs obtained from GDM mothers exposed to a chronic hypoxic condition.

Discussion

GDM is defined as any degree of glucose intolerance with onset or first recognition during pregnancy, and complicates around 2%–7% of all pregnancies in the US (more than 200,000 pregnancies annually). Major risk factors for developing GDM are family history of diabetes, increased pre-gravid body mass index, and a history of prior GDM. Women with multiple gestations, obesity, and bigger placental mass are at higher risk for GDM. On the other hand, GDM itself generates insulin resistance, and reduced sensitivity to insulin action. Furthermore, referer to GDM increases the release of human placental lactogen, growth, and corticotropin-releasing hormones from the placenta. The destiny of GDM is related to uncompensated insulin

Figure 2. Real-time PCR reactions were performed with endothelial growth factor A (VEGFA) mRNA for HUCB ECFCs obtained from gestational and control mothers under normoxic as well as chronic hypoxic condition. (A) The real-time PCR melt curve, (B) The real-time PCR AMP/Cycle, (C) Quantitative assessment of real-time PCR.

Notes: The effects of chronic hypoxia and gestational diabetes were investigated using the pro-angiogenic specific marker VEGFA. Real-time PCR experiments were performed in duplicate in the custom-designed SYBR green mix, as previously described. Each mean gene fold change was normalized to internal control β-actin. Control normoxic (C-HUCB ECFC-N) represents HUCB ECFCs obtained from control pregnancies and plated under normoxic condition; control hypoxic (C-HUCB ECFC-H) represents HUCB ECFCs obtained from control pregnancies and plated under hypoxic condition; GDM normoxic (GDM-HUCB ECFC-N) represents HUCB ECFCs obtained from GDM subjects and plated under normoxic condition; GDM hypoxic (GDM-HUCB ECFC-H) represents HUCB ECFCs obtained from GDM subjects and plated under hypoxic condition. C-HUCB ECFC-H and GDM-HUCB ECFC-N significantly different from C-HUCB ECFC-N.

Abbreviations: PCR, polymerase chain reaction; VEGFA, vascular endothelial growth factor A; mRNA, messenger ribonucleic acid; HUCB, human umbilical cord blood; ECFCs, endothelial colony-forming cells; GDM, gestational diabetes mellitus.
sensitivity or type II DM, whether only appearing during the gestational period or the condition persisting after pregnancy. The definition applies regardless of whether insulin or only diet modification is used during pregnancy for their treatment if the subject demonstrates symptoms of diabetes or high blood glucose (BG) concentration.27,29

GDM has serious long-term health consequences for her adult offspring, by inducing the development of metabolic syndrome, hypertension, type II DM, and premature cardiovascular disease.26,27,29 Development of metabolic syndrome in children with increasing age is correlated with maternal GDM and maternal obesity.1,2

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**Figure 3** Real-time PCR reactions were performed to identify the angiogenesis capability of HUCB eCFCs. (A) The real time PCR melt curve, (B) The real time PCR AMP/Cycle, (C) Quantitative assessment of real time PCR.

**Notes:** For that purpose, pro-angiogenic specific primers IGF-1 were run for real-time PCR with duplicated custom-designed SYBR green mix, as previously described.19,20 Each mean gene fold change was normalized to internal control β-actin. Control normoxic (C-HUCB eCFC-N) represents HUCB-derived EFCFs obtained from control pregnancies and plated under normoxic condition; control hypoxic (CHUCB ECFC-H) represents HUCB-derived EFCFs obtained from control pregnancies and plated under hypoxic condition; GDM normoxic (GDM-HUCB ECFC-N) represents HUCB-derived EFCFs obtained from GDM subjects and plated under normoxic condition; GDM hypoxic (GDM-HUCB ECFC-H) represents HUCB-derived EFCFs obtained from GDM subjects and plated under hypoxic condition. *GDM-HUCB ECFC-H significantly different from C-HUCB ECFC-H.

**Abbreviations:** PCR, polymerase chain reaction; HUCB, human umbilical cord blood; ECFCs, endothelial colony-forming cells; IGF-1, insulin-like growth factor 1; GDM, gestational diabetes mellitus; VEGFA, vascular endothelial growth factor A.
EPCs are believed to differentiate into ECs, apart from angiogenesis. This belief has introduced new therapeutic strategies, and may be a potential therapeutic approach for a variety of vascular diseases. Previous studies have demonstrated that the number and migratory activity of EPCs are inversely correlated with the risk factors for cardiovascular disease and hypertension. The number and function of EPCs sometimes is accepted as a better predictor than the Framingham risk factor score to understand cardiovascular function. In adults, circulating EPCs directly participate in

![Figure 4](image-url)
Chronic hypoxia and human endothelial progenitor cell

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Figure 5 Photomicrographs representing senescence assay against chronic hypoxia. Notes: ECFCs derived from C (n=4) and GDM (n=4) subjects, which were stained positively for SA-β-gal. C-HUCB ECFC-N represents HUCB-derived ECFCs obtained from control pregnancies and plated under normoxic condition; C-HUCB ECFC-H represents HUCB-derived ECFCs obtained from control pregnancies and plated under hypoxic condition; GDM-HUCB ECFC-N represents HUCB-derived ECFCs obtained from GDM subjects and plated under normoxic condition; and GDM-HUCB ECFC-H represents HUCB-derived ECFCs obtained from GDM subjects and plated under hypoxic condition. Abbreviations: ECFC, endothelial colony-forming cell; HUCB, human umbilical cord blood; GDM, gestational diabetes mellitus.

Figure 6 Photomicrographs representing matrigel assay against chronic hypoxia, which represents the vessel-forming ability. Notes: C-HUCB ECFC-N represents human umbilical cord-derived blood ECFCs obtained from control pregnancies and plated under normoxic condition; C-HUCB ECFC-H represents HUCB-derived ECFCs obtained from control pregnancies and plated under hypoxic condition; GDM-HUCB ECFC-N represents HUCB-derived ECFCs obtained from GDM subjects and plated under normoxic condition; and GDM-HUCB ECFC-H represents HUCB-derived ECFCs obtained from GDM subjects and plated under hypoxic condition. Abbreviations: HUCB, human umbilical cord blood; ECFC, endothelial colony-forming cell; GDM, gestational diabetes mellitus.

EPCs represent a minor subpopulation of blood MNCs. In vitro cell culture methods have been developed to identify and expand the EPC population from blood MNCs. ECFCs have proven to be phenotypically indistinguishable from cultured ECs and possess de novo vessel-forming ability. For that reason, ECFCs were selected in all experimental procedures. Gene-specific primers were also specifically designed to reveal their endothelial origin. All control and GDM HUCB ECFCs expressed eNOS, PECAM1 (CD31), VE-cadherin, VEGFA, and IGF-1 mRNAs. The “A type 1 trans-membrane protein” PECAM1 is expressed only in ECs. PECAM1 expressions were stimulated with chronic hypoxia in both control and GDM subjects.

Significant differences were not detected in eNOS mRNA expressions in hypoxic C and GDM HUCB ECFCs but they tended to increase. Chronic hypoxia resulted in significantly decreased eNOS and increased P-eNOS protein expressions obtained from C and GDM HUCB ECFCs. Their quantitative assessments demonstrated that exposure to a gestational
diabetic environment resulted in the most significant decrease in eNOS and increase in P-eNOS protein expressions. If we combine these data, it can be concluded that the reduction of eNOS protein expression against chronic hypoxic environment results from posttranslational modification and phosphorylation of that protein. Diabetic intrauterine environment also negatively affects eNOS protein expression in HUCB ECFCs.

The Ca\(^{++}\)-dependent cell adhesion molecule VE-cadherin (CdH5) is expressed in all cells, but chronic hypoxia did not affect Ca\(^{++}\)-dependent cell adhesion molecule. This indicates that all that cells were able to express Ca\(^{++}\)-regulated adhesion molecule in the hypoxic cell culture environment. Chronic hypoxia also demonstrated increased VEGFA mRNA expressions but not IGF-1. Even GDM-generated insulin resistance and the reduced sensitivity to insulin action of IGF-1 were not affected by hypoxia, though significant differences were detected in the GDM-hypoxic samples when compared to C-hypoxic samples. VEGFA expressions were also significantly increased in GDM-normoxic and C-hypoxic subjects but did not change in GDM-hypoxic samples. This indicates that GDM leads to the failure of ECFCs’ adaptation ability against a chronic hypoxic environment.

Vascular ECs exhibit adaptation to a hypoxic environment using complex compensatory mechanisms, and their progenitors play a key role in sensing and responding to insufficient O\(_2\) perfusion. Under hypoxia, ECs produce different metabolites such as nitric oxide, reactive oxygen species, and lactate. Hypoxia is described as a deficiency in oxygen and distinguishes three major cases according to the quantity of oxygen: moderate hypoxia (5% O\(_2\) triple gas vs normoxia 20% O\(_2\)); severe hypoxia (1% O\(_2\) triple gas); and anoxia. The difference between physiological and pathological vessel formation under different hypoxic conditions for a certain amount of time determines the fate of the cells. The knowledge of how endothelial progenitors act under an existing hypoxic environment could lead to new therapeutic strategies. Furthermore, deficiency in EPCs’ adaptation ability against hypoxia obtained from GDM subjects demonstrates increased cardiovascular risk during adulthood. If we include their HbA\(_{1c}\) level in GDM subjects, which were all normal and basically reflect a short duration of diabetes, we get another perspective. The question, “how did GDM affect their EPCs in a short period” also needs further investigation. The real difference between a clinical and an experimental setting is that we continuously treat the patients and their blood glucose (BG) is always under control except for some fluctuations. Behind that clinical reality, HUCB ECFCs demonstrated that GDM caused reduced proliferation capacity and vessel-forming ability under normoxic as well as hypoxic conditions.

**Figure 7** (A) Senescence assay. Enhanced senescence of HUCB ECFCs (n=4 C and n=4 GDM) exposed to 7 days of hypoxic condition. SA-β-gal assays demonstrated that C- and GDM-HUCB ECFCs exhibited significantly increased in senescence during in vitro exposure to chronic hypoxia which contributed to the reduction in clonogenic capacity and correlated with reduced proliferative ability. C-HUCB ECFC-H and GDM-HUCB ECFC-H are significantly different from C-HUCB ECFC-N and GDM-HUCB ECFC-N; C-HUCB ECFC-N significant difference from GDM-HUCB ECFC-N, and GDM-HUCB ECFC-H; C-HUCB ECFC-H significant difference from GDM-HUCB ECFC-N; C-HUCB ECFC-H significant difference from GDM-HUCB ECFC-H. (B) Matrigel assay. Quantitative assessment of tube formation was conducted by scoring the number of closed capillary tube networks per well. Y-axis represents the percentage of C-normoxic.
HIF 1 activates the transcription of a large group of genes that encode many proteins in angiogenesis against response to hypoxia.\textsuperscript{39} It mainly regulates some genes that are involved in vasodilatation, increased vascular permeability, extracellular matrix remodeling, and EC proliferation. It plays an important role in developmental and pathological vessel formation under hypoxic conditions via \(O_2\)-dependent and/or \(O_2\)-independent pathways.\textsuperscript{7,16,40} It coordinates the proliferation and migration of ECs via increases in eNOS, VEGF, fibroblast growth factor (FGF), platelet-derived growth factor (PDGF), C-X-C chemokine receptor type 4 (CXCR-4), and stromal cell-derived factor 1 (SDF-1) expressions under hypoxic conditions.\textsuperscript{7,16,40} These physiological responses provide either increased \(O_2\) delivery or cell survival with hypoxia, which activates alternative metabolic pathways or reduces reactive oxygen species. On the other hand, HIF-1\(\alpha\) (\(O_2\)-regulated subunit) is a heterodimeric DNA-binding complex that continuously synthesizes and degrades during that period.\textsuperscript{15,16,40} Similarly, chronic hypoxia resulted in the highest increase in HIF 1\(\alpha\) protein expression in HUCB ECFCs obtained from control subjects.

However, when GDM exists as a pathological condition, it demonstrated decreased expression of HIF 1\(\alpha\) transcription factor compared to control subjects. It demonstrates that GDM results in lower vessel-forming ability compared to controls in response to hypoxia.

HUCB ECFCs obtained from GDM and control subjects demonstrated reduced proliferation capacity and vessel-forming ability against chronic hypoxia. They exhibited significantly increased senescence in vitro when exposed to a chronic hypoxia, which contributes to the reduction of their clonogenic capacity. The most significant decrease in capillary tube formation was observed in ECFCs obtained from GDM subjects when exposed to a chronic hypoxic condition. These data suggest that chronic hypoxia results in an overall decline in the proliferative capacity of ECFCs due to premature senescence and diminished vessel-forming ability, as seen from control and GDM subjects. If we combine these data with increased HIF 1\(\alpha\) protein expressions, it mainly indicates that further degradation of that protein or chronic hypoxia-induced HIF 1\(\alpha\) pathway does not affect the vessel-forming ability. The underlying mechanisms need further investigations.

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Disclosure
The author reports no conflicts of interest in this work.

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