Placental nanoparticle gene therapy normalizes gene expression changes in the fetal liver associated with fetal growth restriction in a fetal sex-specific manner

Rebecca L. Wilson¹², Kendal K. Stephens³ and Helen N. Jones¹²

¹Center for Research in Perinatal Outcomes, University of Florida College of Medicine, Gainesville, Florida 32610, USA; ²Department of Physiology and Aging, University of Florida College of Medicine, Gainesville, Florida 32610, USA and ³Department of Obstetrics and Gynecology, University of Cincinnati, Cincinnati, Ohio, 45229, USA

Abstract
Fetal growth restriction (FGR) is associated with increased risk of developing non-communicable diseases. We have a placenta-specific nanoparticle gene therapy protocol that increases placental expression of human insulin-like growth factor 1 (hIGF1), for the treatment of FGR in utero. We aimed to characterize the effects of FGR on hepatic gluconeogenesis pathways during early stages of FGR establishment, and determine whether placental nanoparticle-mediated hIGF1 therapy treatment could resolve differences in the FGR fetus. Female Hartley guinea pigs (dams) were fed either a Control or Maternal Nutrient Restriction (MNR) diet using established protocols. At GD30-33, dams underwent ultrasound guided, transcutaneous, intraplacental injection of hIGF1 nanoparticle or PBS (sham) and were sacrificed 5 days post-injection. Fetal liver tissue was fixed and snap frozen for morphology and gene expression analysis. In female and male fetuses, liver weight as a percentage of body weight was reduced by MNR, and not changed with hIGF1 nanoparticle treatment. In female fetal livers, expression of hypoxia inducible factor 1 (Hif1α) and tumor necrosis factor (Tnfα) were increased in MNR compared to Control, but reduced in MNR + hIGF1 compared to MNR. In male fetal liver, MNR increased expression of Igf1 and decreased expression of Igf2 compared to Control. Igf1 and Igf2 expression was restored to Control levels in the MNR + hIGF1 group. This data provides further insight into the sex-specific mechanistic adaptations seen in FGR fetuses and demonstrates that disruption to fetal developmental mechanisms may be returned to normal by treatment of the placenta.

Introduction
Fetal growth restriction (FGR; estimated fetal weight <10th percentile) occurs in up to 10% of pregnancies in the developed world with suboptimal fetal nutrition and uteroplacental perfusion accounting for 25–30% of cases.¹² As one of the leading causes of stillbirth, miscarriage and infant morbidity, FGR is also associated with an increased risk of developing non-communicable diseases (NCDs; including cardiovascular disease, metabolic disease, central obesity, type 2 diabetes) in later life.³⁵ Currently, FGR is diagnosed after it is established and there is no effective in utero treatment for FGR, thus no clinical intervention that could potentially impact or prevent the development of NCDs in adulthood.

The “developmental origins of health and disease” (DoHAD) hypothesis states that early environmental stressors during critical fetal developmental windows result in permanent, adaptive structural, and physiologic changes that predispose the offspring to metabolic, endocrine, and cardiovascular disease in postnatal life.⁶ Animal models have shown that FGR is associated with increased hepatic gluconeogenic gene expression and glucose production in late pregnancy, in response to reduced placental glucose supply.⁷⁹ Whilst these adaptive responses may be temporarily beneficial in utero, persistent increased liver glucose production beyond birth, when dietary glucose is no longer limited, may have adverse consequences to health. Excess glucose production leads to hyperglycemia and the development of obesity and/or metabolic disease.

Liver gluconeogenesis is the de novo synthesis of glucose from noncarbohydrate precursors and is principally controlled by activities of enzymes such as phosphoenolpyruvate carboxykinase (PKC/PEPCK) and glucose-6-phosphatase (G6PC).¹⁰ There are two isozymes of PKC: the cytosolic isozyme PCK1 and the mitochondrial isozyme PCK2, both of which are involved in hepatic gluconeogenesis.¹¹ During fetal development, gluconeogenesis occurs in late gestation, although activity of gluconeogenic enzymes has been shown at mid-pregnancy.¹² Studies, predominantly performed in sheep models of FGR, and focused on changes in late gestation, have shown fetal hepatic adaptations to support glucose production.⁷¹⁰–¹² FGR is associated with
increased expression of Pck1, Pck2, and G6pc in the fetal liver, and transcriptomic analysis indicates increased amino acid catabolism and cell stress along with decreased mitochondrial activity.\textsuperscript{14} Moreover, increased expression of Pck has been shown to persist into adulthood, at the equivalent human age of 50–60 years.\textsuperscript{16} However, the effects FGR on liver expression of gluconeogenic enzymes around the initiation of FGR are lacking.

Given that fetal hepatic adaptations to glucose production are linked to insufficient placental glucose supply, directly targeting the dysfunctional placenta to increase glucose and other nutrient transport capacities may restore supply and mitigate DoHAD-associated developmental programming. We have developed the use of a polymer-based nanoparticle that facilitates nonviral, transient (does not integrate into the genome), gene delivery specifically to the placenta.\textsuperscript{17,18} Using a biosynthetic HPMA-DMEAMA (N-(2-hydroxypropyl) methacrylamide-2-(dimethylamino)ethyl methacrylate) copolymer, complexed with a plasmid containing the human insulin-like growth factor 1 (hIGF1) gene under the control of trophoblast specific promoters (PLAC1, CYP19A1), we have successfully shown efficient nanoparticle uptake into human syncytiotrophoblast ex vivo\textsuperscript{19} as well as in vivo using animal models.\textsuperscript{17,19-21} Short-term effects of hIGF1 gene therapy results in increased placental expression of glucose and amino acid transporters, maintenance of normal fetal growth under FGR conditions, and increased fetal glucose concentrations.\textsuperscript{17,19-21}

Importantly, our nanoparticle gene therapy is proven to be safe to both mother and fetus and is capable of positively influencing placental function in diverse models of FGR as IGF1 is central to most mechanisms responsible for FGR associated with placental dysfunction, and a major regulator of normal placental and fetal growth and development.\textsuperscript{22}

We have previously shown in the guinea pig MNR model of FGR, efficient uptake of our hIGF1 nanoparticle gene therapy into the guinea pig placenta, and no transfer of nanoparticle or hIGF1 plasmid into fetal circulation.\textsuperscript{21} In the present study, we aimed to characterize the effects of FGR on hepatic insulin-sensing and gluconeogenic enzyme gene expression at the initial stages of FGR establishment in the fetal guinea pig, and determine whether treatment of the placenta with our hIGF1 nanoparticle gene therapy could resolve differences in hepatic insulin-sensing and gluconeogenic enzyme gene expression in the FGR fetus.

Materials and methods
Polymer synthesis and nanoparticle formation

Detailed methods on the synthesis of the (PHPMA\textsubscript{115-b-PDMAEMA\textsubscript{115}}) copolymer, and nanoparticle formation can be found in Wilson et al., 2022.\textsuperscript{21} Briefly, plasmids containing the human IGF1 gene under control of the trophoblast-specific CYP19A1 promoter were mixed with the nonviral PHPMA\textsubscript{115-b-PDMAEMA\textsubscript{115}} copolymer for 1 h at room temperature to form the hIGF1 nanoparticle. Details about physiochemical properties, and cellular safety and efficiency of the PHPMA\textsubscript{115-b-PDMAEMA\textsubscript{115}} nanoparticle has been previously published,\textsuperscript{21} and has been proven safe for both mother and fetus in numerous animal models.\textsuperscript{17,19,20} Finally, it is unlikely clinically that this nanoparticle gene therapy will be provided to a normally growing fetus. However, we have also shown that administration of this nanoparticle gene therapy to the placenta in a normal pregnancy environment results in down-regulation of decidual and placental mTOR signaling and growth factor gene expression in order to maintain placental homeostasis.\textsuperscript{23}

Animal care and transuterine, intraplacental nanoparticle administration

Animal care and usage was approved by the Institutional Animal Care and Use Committees at Cincinnati Children’s Hospital and Medical Center (Protocol number 2017-0065) and University of Florida (Protocol number 202011236). Detailed information on animal care, maternal nutrient restriction (MNR) implementation, and ultrasound-guided transuterine, intra-placental nanoparticle administration can be found in Wilson et al., 2022.\textsuperscript{21} Briefly, female (dams) Dunkin–Hartley guinea pigs (Charles River Laboratories, Wilmington, MA) were purchased and assigned to either a Control diet group (\textit{n} = 7) where food (LabDiet diet 5025: 27% protein, 13.5% fat, and 60% carbohydrate as % of energy) and water was provided ad libitum, or MNR diet group (\textit{n} = 12) where water was provided ad libitum, but food intake was restricted to 70% per kilogram body weight of the Control group from at least 4 weeks preconception through to mid-pregnancy (gestational day (GD) 30), thereafter increasing to 90% until term (GD65-70). Time mating with males was performed as outlined in Wilson et al., 2021,\textsuperscript{21} and pregnancy confirmation ultrasounds performed at GD21 using a Voluson 1 portable ultrasound machine (GE) with a 125 E 12 MHz vascular probe (GE). At GD30–33 dams underwent ultrasound-guided, transuterine, intraplacental injections of either sham (200 μL of PBS; \textit{n} = 7 Control and \textit{n} = 5 MNR) or hIGF1 nanoparticle gene therapy (50 μg plasmid in 200 μL injection; \textit{n} = 7 MNR + hIGF1). Dams were sacrificed 5 days after injection (GD35-38) by carbon dioxide asphyxiation followed by cardiac puncture and exsanguination. Fetuses (Control: \textit{n} = 4 female and 8 male fetuses, MNR: \textit{n} = 7 female and 7 male fetuses, and MNR + hIGF1; \textit{n} = 8 female and 11 male fetuses) and placentas were removed from the gravid uterus and weighed. Glucose concentrations in both maternal and fetal blood was measured using a glucometer. Fetal sex was determined by examination of the gonads and confirmed using PCR as previously published.\textsuperscript{20} Fetal livers were dissected, weighed, and halved to be either fixed in 4% w/v paraformaldehyde (PFA) or snap-frozen in liquid nitrogen and stored at −80°C.

Histology and immunohistochemistry

For assessment of steatosis and fibrosis, 5 μm thick sections of PFA-fixed, paraffin embedded fetal liver tissue were obtained, de-waxed, and rehydrated following standard protocol. Hematoxylin and eosin staining was performed as standard. Immunohistochemistry (IHC) was performed as previously described\textsuperscript{20} to assess nuclear expression of Ki67 (Invitrogen MA5-14520; diluted 1:200), and nuclei were counter stained with hematoxylin. Following IHC staining, slides were imaged using the Axioscan (Zeiss) microscope, and 10 random 40x magnification images obtained using the Zen Blue software (Zeiss). For each image, cells positive for Ki67 (brown) and negative for Ki67 (blue) were counted using the Threshold and Watershed functions in ImageJ software\textsuperscript{24} and averaged across the 10 images per slide to obtain a percentage positive result.

RNA isolations and quantitative PCR (qPCR)

For hepatic gene expression analysis, approximately 50 mg of snap frozen fetal liver tissue was lysed in RLT-lysis buffer
(Qiagen) with homogenization aided by a tissue-lyzer. RNA was extracted using the RNeasy Mini kit (Qiagen), and included DNase treatment following standard manufacturers protocol. 1 μg of RNA was converted to cDNA using the High-capacity cDNA Reverse Transcription kit (Applied Biosystems) and diluted to 1:100. For qPCR, 2.5 μL of cDNA was mixed with 10 μL of PowerUp SYBR green (Applied Biosystems), 1.2 μL of KiCqStart SYBR Green Predesigned Primers (Sigma) at a concentration of 10 nM, and water to make up a total reaction volume of 20 μL. Gene expression was normalized using housekeeping genes β-actin and Rasp20 (See 21). qPCR was performed using the Quant3 Real-Time PCR System (Applied Biosystems), and relative mRNA expression calculated using the comparative CT method with the Design and Analysis 2 v2.6.0 software (Applied Biosystems).

Statistical analysis

All statistical analyses were performed using SPSS Statistics 27 software with female and male fetuses analyzed separately. Due to the small sample size, all data was assumed not normally distributed so generalized estimating equations with gamma log-link were used to determine differences between diet and nanoparticle treatment. Dams were considered the subject, diet, and nanoparticle treatment treated as main effects, maternal environment treated as a random effect and gestational age as a covariate. Litter size was also included as a covariate but removed as there was no significant effect for any of the outcomes. Statistical significance was considered at \( P \leq 0.05 \). For statistically significant results, a Bonferroni post hoc analysis was performed. Results are reported as estimated marginal means ± standard error.

Results

Maternal nutrient restriction reduces fetal liver growth and results in brain-sparing at mid-pregnancy

The MNR diet resulted in decreased fetal weight in both females and males at mid-pregnancy, and there was no effect of hIGF1 nanoparticle treatment after 5 days (Fig. 1a and 1b). Fetal liver weight as a percentage of fetal weight was decreased in MNR and MNR + hIGF1 nanoparticle treatment in female and male fetuses compared to Control (Fig. 1c and 1d). There was evidence of brain-sparing, as indicated by increased fetal brain:liver ratio in female and male fetuses, in the MNR and MNR + hIGF1 nanoparticle treatment fetuses when compared to Control fetuses (Fig. 1e and 1f).

Maternal nutrient restriction reduces liver proliferation in female fetuses and is normalized with hIGF1 nanoparticle treatment

Morphologically, there was no evidence of increased steatosis or fibrosis in the livers of either female or male fetuses with MNR or hIGF1 nanoparticle treatment (Fig. 2). In female fetal livers, MNR reduced proliferation, as evident by the percentage of cells positive for Ki67, of the hepatocytes when compared to Control, however percentage of cells positive for Ki67 in MNR + hIGF1 nanoparticle treated was comparable to Controls (Fig. 3a). In male fetal livers, there was no effect of MNR or hIGF1 nanoparticle treatment on proliferation of hepatocytes (Fig. 3b).

Expression of hypoxia markers is reduced with hIGF1 nanoparticle treatment in MNR female fetal livers

In female fetal livers at mid-pregnancy, there was increased expression of growth factors Tgfβ, Ctgf, and Mmp2 with MNR when compared to Control (Fig. 4a–4c). Expression of Tgfβ, Ctgf, and Mmp2 remained increased in the MNR + hIGF1 nanoparticle treatment compared to Control. On the other hand, expression of markers of hypoxia: Tfia and Hif1α were increased in MNR female fetal livers when compared to Control, but decreased by hIGF1 nanoparticle treatment when compared to MNR, and towards comparable levels with Control female fetal livers (Fig. 4d and 4e). qPCR analysis of growth factors and hypoxia markers in the mid-pregnancy male fetal liver showed no differences in expression with either MNR or hIGF1 nanoparticle treatment (Supplemental Fig. 1).

Placental hIGF1 nanoparticle treatment increased fetal glucose concentrations in male fetuses at mid-pregnancy

We have previously reported no difference in maternal blood glucose levels at mid-pregnancy with either diet or hIGF1 nanoparticle treatment.21 There was no difference in fetal blood glucose concentrations in females and males between MNR and Control (Fig. 5a and 5b). However, hIGF1 nanoparticle treatment increased fetal blood glucose concentrations when compared to Control and MNR, but only in male fetuses.

Fetal liver gene expression of insulin-sensing and gluconeogenesis enzymes is affected by MNR at mid-pregnancy, and normalized with placental hIGF1 nanoparticle treatment in male fetuses only

It has been hypothesized that developmental programming in the fetal liver that predisposes offspring to obesity and metabolic diseases is due to increased gluconeogenesis in the fetal liver that persists after birth.15,25 qPCR analysis of insulin-sensing and gluconeogenesis enzymes in the mid-pregnancy female fetal liver showed very few differences (Supplemental Fig. 2), with only Igbp3 increased in MNR compared to Control, and returned to Control expression levels in the MNR + hIGF1 group (Estimated marginal mean ± SEM of Relative Expression: Control = 1.15 ± 0.22 vs. MNR = 2.10 ± 0.06 vs. MNR + hIGF1 = 1.31 ± 0.22; \( P \)-value Diet = 0.002, \( P \)-value Treatment = 0.005). In mid-pregnancy males, MNR increased expression of Igf1, and reduced expression of Igf2, G6pc and Pck1 when compared to Control (Fig. 6a–6d, respectively). Furthermore, MNR + hIGF1 nanoparticle treatment returned mid-pregnancy liver expression of Igf1, Igf2, G6pc, and Pck1 to that, or towards that, of Control. Additionally, hIGF1 nanoparticle treatment increased male fetal liver expression of Igbp1 and GcgR compared to Control and MNR (Fig. 6e and 6f, respectively).

Discussion

Currently, there is no effective in utero treatment for FGR and thus, no clinical intervention that could potentially impact or prevent the increased risk of developing an NCD. In the present study, we show that MNR affects different physiological pathways depending on fetal sex; in females fetuses growth mechanisms are impacted compared to pathways involved in glucose production which are impacted in male fetuses. Moreover, we show short-term treatment of the placenta with hIGF1 nanoparticle gene therapy is capable of normalizing changes to fetal hepatic gene
expression. Overall, this data provides further mechanistic understanding of how MNR and FGR affect hepatic gene expression and development and highlights the importance of understanding sex-specific risk windows during fetal development and potential considerations when developing pregnancy therapeutics.

The influence of fetal sex as a biological variable is well established. In human pregnancies, data consistently shows that risk of complications such as preterm delivery is higher in males compared to females.26 In this study, fetal weight, and liver weight were decreased in female and male fetuses with MNR when compared to Control, an outcome routinely found in late-pregnancy in the guinea pig MNR model of FGR.27-29 However, at mid-pregnancy, and during the initial stages of FGR establishment, it was only in female fetal livers where MNR reduced liver hepatocyte proliferation, and increased the expression of growth factors and markers of

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**Fig. 1.** Effect of maternal nutrient restriction (MNR) and hIGF1 nanoparticle treatment on mid-pregnancy fetal growth parameters. 

- **A.** MNR reduced mid-pregnancy fetal weight of female fetuses, and was not different with hIGF1 nanoparticle treatment.
- **B.** MNR reduced mid-pregnancy fetal weight of male fetuses, and was not different with hIGF1 nanoparticle treatment.
- **C.** MNR reduced mid-pregnancy fetal liver weight, as a percentage of fetal weight in female fetuses, and was not different with hIGF1 nanoparticle treatment.
- **D.** MNR reduced mid-pregnancy fetal liver weight, as a percentage of fetal weight in male fetuses, and was not different with hIGF1 nanoparticle treatment.
- **E.** MNR increased mid-pregnancy brain:liver weight ratio in female fetuses, and was not different with hIGF1 nanoparticle treatment.
- **F.** MNR increased mid-pregnancy brain:liver weight ratio in male fetuses, and was not different with hIGF1 nanoparticle treatment.

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**Fig. 2.** Representative images of hematoxylin and eosin (H & E) stained fetal livers at mid-pregnancy with maternal nutrient restriction (MNR) and hIGF1 nanoparticle treatment. There was no evidence of increased steatosis or fibrosis in the livers of either female or male fetuses with MNR or hIGF1 nanoparticle treatment. H & E images are taken at 40x magnification, scale bar = 10 μm. n = 4 Control dams (4 female and 4 male fetuses), 5 MNR dams (5 female and 5 male fetuses), and 7 MNR + hIGF1 nanoparticle dams (7 female and 7 male fetuses).
Fig. 3. Effect of maternal nutrient restriction (MNR) and hIGF1 nanoparticle treatment on mid-pregnancy fetal liver hepatocyte proliferation. A. In female fetal livers, MNR reduced the percentage of hepatocytes positive for Ki67 (brown nuclei) when compared to Control. However, percentage of hepatocytes positive for Ki67 in MNR + hIGF1 nanoparticle treated female fetal livers was comparable to Control. B. There was no effect of either MNR or hIGF1 nanoparticle treatment on percentage of hepatocytes positive for Ki67 in male fetal livers. C. Representative images of Ki67 immunohistochemistry stained fetal livers, taken at 40x magnification; scale bar = 10 μm. n = 4 Control dams (4 female and 4 male fetuses), 5 MNR dams (5 female and 5 male fetuses), and 7 MNR + hIGF1 nanoparticle dams (7 female and 7 male fetuses). Data are estimated marginal means ± 95% confidence interval. P values calculated using generalized estimating equations with Bonferroni post hoc analysis. *P < 0.05.

Fig. 4. Effect of maternal nutrient restriction (MNR) and hIGF1 nanoparticle treatment on mid-pregnancy female fetal liver growth factor and markers of hypoxia gene expression. MNR increased expression of transforming growth factor beta (Tgfβ; A), connective tissue growth factor (Ctgf; B), matrix metalloproteinase 2 (Mmp2; C) in female fetal livers compared to Control female fetal livers. hIGF1 nanoparticle treatment did not affect expression of Tgfβ, Ctgf, and Mmp2 which remained increased when compared to Control. MNR increased expression of tumor necrosis factor alpha (Tnfa; D) and hypoxia inducible factor 1 alpha (Hif1α; E) when compared to Control. hIGF1 nanoparticle treatment decreased expression in male MNR fetal liver tissue compared to MNR. n = 7 Control dams (4 female fetuses), 5 MNR dams (7 female fetuses), and 7 MNR + hIGF1 nanoparticle dams (8 female fetuses). Data are estimated marginal means ± 95% confidence interval. P values calculated using generalized estimating equations with Bonferroni post hoc analysis. *P < 0.05; **P < 0.01; ***P < 0.001.
hypoxia; expression remained comparable to Control in male livers. It has previously been reported in young adult male FGR offspring, that mRNA expression of Tgfβ, Ctgf, and Mmp2 were increased in liver tissue,30 however, females were not assessed. Traditionally, increased expression of these growth factors has been associated with the promotion of liver fibrosis in adulthood,31 and increased expression would be considered detrimental to liver physiology. However, in mid-pregnancy fetal development, the pro-proliferative effects32 are likely positive for organ development, particularly given reduced liver weight as a percentage of fetal weight. Overall, our results suggest in the early stages of FGR induction, growth pathways are more disrupted than pathways relating to insulin-sensing and gluconeogenic enzyme expression in the livers of female fetuses, indicating that female fetuses prioritize supporting liver growth.

Under normal growth conditions, endogenous fetal glucose production is negligible because glucose supply from the placenta is sufficient. However, in cases of FGR the fetal liver increases gluconeogenesis, and thus glucose production, in order to maintain vital glucose supply to the developing organs.15,25 Others have shown increased hepatic gene expression of gluconeogenesis enzymes in growth restricted fetuses compared to normal fetuses in fetal sheep and rats in late-pregnancy, but did not separate fetal sex.7,8,15 In our study, at the time of FGR establishment, there was reduced expression of gluconeogenic enzymes G6pc and Pck1, which may represent a compensation trigger for the increased expression seen in late-pregnancy in other studies. However, the decrease in G6pc and Pck1 expression was only observed in male fetuses. Adequate liver development and functionality ensures glycogen deposition and gluconeogenic ability, both of which are essential during the first stages of postnatal life.33 In the guinea pig fetal liver between GD30 and 40, the glycolytic rate is thought to be high, but there is little formation of glycogen and glucose, despite the expression of gluconeogenic enzymes.12

Fig. 5. Effect of maternal nutrient restriction (MNR) and hIGF1 nanoparticle treatment on mid-pregnancy fetal blood glucose levels. A. There was no difference in blood glucose concentrations in female fetuses with either MNR or hIGF1 nanoparticle treatment. B. In male fetuses, there was no difference in blood glucose concentrations between Control and MNR, but hIGF1 nanoparticle treatment increased blood glucose concentrations compared to sham. n = 7 Control dams (4 female and 8 male fetuses), 5 MNR dams (7 female and 7 male fetuses), and 7 MNR + hIGF1 nanoparticle dams (8 female and 11 male fetuses). Data are estimated marginal means ± 95% confidence interval. P values calculated using generalized estimating equations with Bonferroni post hoc analysis. *P < 0.05.

Fig. 6. Effect of maternal nutrient restriction (MNR) and hIGF1 nanoparticle treatment on mid-pregnancy male fetal liver insulin sensing and gluconeogenesis enzyme gene expression. MNR increased expression of insulin-like growth factor 1 (Igf1; A), and decreased expression of Igf2 (B), glucose-6-phosphatase (G6pc; C), and phosphoenolpyruvate carboxykinase 1 (Pck1; D) in male fetal livers compared to Control male fetal livers. hIGF1 nanoparticle treatment restored expression of Igf1, Igf2, G6pc, and Pck1, toward or back to normal. hIGF1 nanoparticle treatment increased expression of Igf Binding Protein 1 (IgfBP1; E) and Glucagon Receptor (GcgR; F) compared to Control and MNR male fetuses. n = 7 Control dams (8 male fetuses), 5 MNR dams (7 male fetuses), and 7 MNR + hIGF1 nanoparticle dams (11 male fetuses). Data are estimated marginal means ± 95% confidence interval. P values calculated using generalized estimating equations with Bonferroni post hoc analysis. *P < 0.05; **P < 0.01; ***P < 0.001.
in liver development to adverse in utero environments, including blood flow and gene expression, whilst important to ensure short-term survival, may have longer-term detrimental consequences in the face of an enriched postnatal diet. Evidence from experimental models demonstrate responses to adverse in utero environments, with males affected to a greater extent than females, putting male offspring at higher risk of cardiovascular and metabolic disease. Overall, our data suggests that during the early stages of FGR establishment in males, insulin signaling and gluconeogenic enzyme expression is affected disproportionately compared to growth patterning and may be a key contributor as to why males are at higher risk of developing obesity diabetes in adulthood.

At time of FGR establishment in both female and male fetuses, changes in liver gene expression of some genes were normalized with short-term placenta hIGF1 nanoparticle treatment. We have previously shown the inability for both nanoparticle and plasmid to cross the placenta and enter fetal circulation, thus any changes in fetal liver gene expression with placental hIGF1 nanoparticle treatment are indirect. During fetal development, there is multidirectional communication between mother, placenta, and fetus. Nutrients, oxygen and signaling factors like hormones, are transferred across the placenta, through the umbilical cord to the liver. Thus, the liver is the first organ which receives nutrient and oxygen rich blood from the placenta. Presented here, there was increased expression of gluconeogenic enzymes, towards normal, in male fetal livers with placenta hIGF1 treatment when compared to sham. Analysis of the placental response to hIGF1 nanoparticle treatment shows increased expression of glucose and amino acid transporters, likely resulting in increased glucose transport across the placenta, presenting a potential mechanism by which fetal liver gene expression of these enzymes is changed. Furthermore, placental hIGF1 nanoparticle treatment resulted in reduced expression of hypoxia markers in MNR female fetal livers. The molecular and physiological mechanisms behind the normalization of Hif1α and Tnfa are yet to be determined, however suggest the ability for the hIGF1 nanoparticle treatment of the placenta to result in reduced hypoxia in fetal livers. Our analysis of placental morphology with hIGF1 nanoparticle treatment indicates reduced interhaemal distance between maternal and fetal circulation likely resulting in increased oxygen diffusion, and a possible mechanism by which expression of Hif1α and Tnfa reduced.

The aim of this study was to assess the immediate, short-term impacts of placenta hIGF1 nanoparticle treatment on developmental programming in the fetal liver at the initial stages of FGR establishment. We have shown that secondary to placental treatment normalization of gene expression changes associated with MNR/FGR occurs. However, given the short time period, there was no significant improvement in fetal weight. Therefore, we are focusing our future research on performing multiple placental hIGF1 nanoparticle treatments over a longer time period in mid-late pregnancy. Overall, this data shows a potential method by which an in utero treatment of the placenta can impact developmental programming and may prevent increased risk of diseases like cardiovascular disease, metabolic disease, obesity, and diabetes in later life.

Acknowledgments. We would like to thank Drs Craig Duvall and Mukesh Gupta for providing the co-polymer and Mrs Kristin Lampre for her assistance with the animal experiments.

Author Contributions. RLW conceived the study, performed experiments, analyzed data, and wrote manuscript. KKS performed experiments, analyzed data, and edited manuscript. HNJ obtained funding, conceived the study, and edited manuscript. All authors approve final version of manuscript.

Funding. This study was funded by Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD) award R01HD090657 (HNJ).

Competing Interests. The authors have declared that no competing interest exists.

Ethics approval. Animal care and usage was approved by the Institutional Animal Care and Use Committees at Cincinnati Children’s Hospital and Medical Center (Protocol number 2017-0065).

Supplementary materials. For supplementary material for this article, please visit https://doi.org/10.1017/S2040174423000016

Data availability. All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials.

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