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Poor fetal growth, also known as intrauterine growth restriction (IUGR), is a worldwide health concern. IUGR is commonly associated with both an increased risk in perinatal mortality and a higher prevalence of developing chronic metabolic diseases later in life. Obesity, type 2 diabetes or metabolic syndrome could result from noxious "metabolic programming." In order to better understand early alterations involved in metabolic programming, we modeled IUGR rat pups through either prenatal exposure to synthetic glucocorticoid (dams infused with dexamethasone 100 µg/kg/day, DEX) or prenatal undernutrition (dams feeding restricted to 30% of ad libitum intake, UN). Physiological (glucose and insulin tolerance), morphometric (automated tissue image analysis) and transcriptomic (quantitative PCR) approaches were combined during early life of these IUGR pups with a special focus on their endocrine pancreas and adipose tissue development. In the absence of catch-up growth before weaning, DEX and UN IUGR pups both presented basal hyperglycaemia, decreased glucose tolerance, and pancreatic islet atrophy. Other early metabolic [...]
Early Metabolic Defects in Dexamethasone-Exposed and Undernourished Intrauterine Growth Restricted Rats

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Abstract

Poor fetal growth, also known as intrauterine growth restriction (IUGR), is a worldwide health concern. IUGR is commonly associated with both an increased risk in perinatal mortality and a higher prevalence of developing chronic metabolic diseases later in life. Obesity, type 2 diabetes or metabolic syndrome could result from noxious “metabolic programming.” In order to better understand early alterations involved in metabolic programming, we modeled IUGR rat pups through either prenatal exposure to synthetic glucocorticoid (dams infused with dexamethasone 100 µg/kg/day, DEX) or prenatal undernutrition (dams feeding restricted to 30% of ad libitum intake, UN). Physiological (glucose and insulin tolerance), morphometric (automated tissue image analysis) and transcriptomic (quantitative PCR) approaches were combined during early life of these IUGR pups with a special focus on their endocrine pancreas and adipose tissue development. In the absence of catch-up growth before weaning, DEX and UN IUGR pups both presented basal hyperglycaemia, decreased glucose tolerance, and pancreatic islet atrophy. Other early metabolic defects were model-specific: DEX pups presented decreased insulin sensitivity whereas UN pups exhibited lowered glucose-induced insulin secretion and more marked alterations in gene expression of pancreatic islet and adipose tissue development regulators. In conclusion, these results show that before any catch-up growth, IUGR rats present early physiologic, morphologic and transcriptomic defects, which can be considered as initial mechanistic basis of metabolic programming.

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Introduction

Adverse fetal environment alters organ development and leads to poor fetal growth, a phenomenon also called intrauterine growth restriction (IUGR). Clinically, IUGR is commonly associated with an increased incidence in perinatal mortality but also with an elevated risk to develop chronic metabolic diseases (such as obesity and type 2 diabetes) later in life, reflecting noxious “metabolic programming” [1]. To elucidate the mechanisms involved in this deleterious process, IUGR can be experimentally induced in rodents by placental blood supply restriction [2], hypoxia [3], maternal diabetes [4] and more commonly by different manipulations of the maternal diet such as protein or caloric restriction [5–10]. Special focus has been placed on the impact of the IUGR modeling regarding endocrine pancreas development due to its central function in the control of glucose homeostasis [11]. Maternal low caloric diet decreases beta-cell neogenesis in relation to elevated glucocorticoid levels [12]. Indeed, excessive fetal exposure to glucocorticoids appears to be a common mechanism by which inadequate intrauterine environment leads to deleterious pancreatic islet development resulting in later metabolic disturbances. In vitro treatment of pancreatic buds with dexamethasone, a synthetic glucocorticoid, favors pancreatic differentiation into exocrine tissue and represses beta-cell differentiation [13], [14]. In vivo, dexamethasone exposure during fetal life, mimicking maternal stress, as well as pharmacological inhibition of the placental 11 beta-hydroxysteroid dehydrogenase type 2 (11β-HSD2), the enzyme protecting the fetus from maternal glucocorticoids, result in IUGR, decreased beta-cell pool and impaired glucose homeostasis [15], [16]. Inversely, mice harbouring a deletion of the glucocorticoid receptor in Pdx-1-expressing precursor cells present an elevation of beta-cell mass [13], [17]. Numerous studies dealing with IUGR and programming of metabolic diseases have focused on late-onset metabolic syndrome in IUGR pups [5–7], [18–20]. Less knowledge is currently available about pre-weaning metabolic defects in IUGR pups. Moreover, the causal link between early catch-up growth and metabolic programming remains elusive. In consequence, precious metabolic investigations would allow to better understand the genesis of late onset metabolic syndrome. To this aim, we combined physiological, morphometric and transcriptomic approaches, focusing on endocrine pancreas and adipose tissue development during the early postnatal period. Rat IUGR pups were modeled through prenatal exposure to synthetic glucocorticoid (dams infused with dexamethasone, DEX) or through prenatal undernutrition (restricted pair-fed dams, UN). This set
of experiments should allow better understanding precocious physiopathological processes involved in metabolic programming.

Methods

Animals and experimental design

All experimental protocols were approved by the “State of Geneva Veterinary Office” (Geneva, Switzerland). Female and male (275–325 g) Sprague-Dawley OEA rats (Charles River Laboratories, France) were used for breeding and housed under specific pathogen-free conditions in an environmentally-controlled clean room at the School of Medicine animal facilities (Medical Center University, University of Geneva). The animals were time mated (the day that sperm-positive smears were obtained was declared day 0 of gestation) and then caged individually before being assigned to one experimental group. In parallel a control group (CON) and two maternal models aimed to induce IUGR in progeny were used: the first model (DEX) consisted in dexamethasone (Sigma Aldrich, Buchs, Switzerland) exposure during the last week of gestation (infusion through an osmotic Alzet minipump delivering 100 μg/kg/day) and the second model (UN) consisted in maternal severe food restriction (gestational time-matched pair-feeding to 30% of normal food intake during the entire gestation). Body weights of gestating rats were monitored daily. At parturition (postnatal day 0, PND0), all pups were sexed, weighed and then nourished by their own mothers fed ad libitum. No reduction of litter size was performed, only excluding litters with extreme number of pups (<10 and >15). Post-delivery dam weights and food intake were monitored twice weekly and pup weights were monitored at PND7, 14 and 21. To avoid any sex confusing effect and interference in hormonal status, all investigations were performed on male pups only. One cohort of pups was sacrificed at PND7 (for hepatic gene expression and protein levels, pancreatic morphometry and histology, pancreatic islet gene expression analysis, pancreatic insulin content, corticosterone levels) and another cohort was sacrificed at PND21 (for white and brown adipose tissue investigations and leptin levels). Another independent cohort of animals was used at PND21 for a glucose tolerance test (GTT) and the same cohort was used after one week of recovery at PND28 for an insulin tolerance test (ITT). For physiological tests (GTT, ITT, insulin content), morphometric analyses (pancreatic islet size and number) and transcriptomic measures (gene expression in liver/pancreatic islets/adipose tissue), we studied pups originating from three independent litters.

Histology and morphometry of pancreas

At PND7, pancreases of 5 to 6 rats in each group were fixed in Bouin’s solution, embedded in paraffin and cut (5 μm sections). For morphometry, three sections of each pancreas were stained with aldehyde fuchsine before being analysed with a personalized program in MetaMorph® Imaging System (Meta Imaging Software, Molecular Devices Corporation, Pennsylvania, U.S.A.). This program determines the total area of pancreatic tissue, the total area of Langerhans islets, the number of islets and their size. For number and size determination, only islets larger than 300 μm² were used for calculations. For the immunohistochemical staining of insulin and glucagon positive cells, the sections were deparaffinized with xylene and rehydrated through graded alcohols before permeabilisation (PBS-Triton 0.1%) and antigen blockade (PBS-BSA 5%). Sections were then incubated 2 h at room temperature with a solution containing a 1:500 mouse monoclonal insulin antibody (Sigma Aldrich, Buchs, Switzerland) and a 1:500 rabbit polyclonal glucagon antibody (Millipore, Billerica, MA, U.S.A). Sections were washed and incubated 45 min with secondary fluorescent antibodies AlexaFluor 488 goat anti-mouse and 555 goat anti-rabbit (respectively for insulin and glucagon). Images were taken using the Mirax system or the axicam system (Carl Zeiss MicroImaging GmbH, Germany).

Islet isolation

At PND7, pancreases were immediately excised and cut into small pieces in Hanks solution before transfer into 5 mL of Hanks-collagenase type V solution (Sigma Aldrich, Buchs, Switzerland) incubated in a 37°C water bath for 7 minutes. Digestion was stopped by addition of Hanks/BSA on ice and the pancreatic tissue was centrifuged several times before undigested fragments were carefully removed. After washing with Hanks solution, the tissue was concentrated into a pellet which was resuspended on a Histopaque® 1077 gradient (Sigma Aldrich, Buchs, Switzerland). After centrifugation at 2500 rpm for 20 minutes, islets were harvested from the interface between the layers, washed and finally concentrated into pellets and immediately used for RNA extraction.

Gene expression

Total RNA from PND7 rat islets was extracted using the RNeasy Mini Kit® according to the manufacturer’s protocol (Qiagen, Basel, Switzerland) and total RNA from PND7 liver, PND21 epididymal white adipose tissue (eWAT) and PND21 interscapular brown adipose tissue (iBAT) were prepared using the Trizol reagent (Invitrogen, Basel, Switzerland), according to manufacturer’s instructions. One microgram of pancreatic islets total RNA and five micrograms of liver, eWAT and iBAT total RNA were reverse-transcribed using 800 units of Moloney murine leukemia virus reverse transcriptase (Invitrogen, Basel, Switzerland), in the presence of 0.3 units/μL RNAsin (Promega Corp, Madison, WI), 7.5 μM of random primers (oligo(dN)6), 1.2 mM dNTP and 12 μM of DTT. The expression of the cDNAs of interest were determined by quantitative real-time PCR using an ABI 7000 Sequence Detection System (Applied Europe, Rotkreuz, Switzerland) and were normalized using the housekeeping genes Ppia (cyclophilin) and Rp1p (36B4). PCR products were quantified using the SYBR Green Core Reagent kit (Applied Europe, Rotkreuz, Switzerland) and results were expressed in arbitrary units (A.U) relative to the mean value of the control group. Primers sequences designed using the Primer Express software (Applied Europe, Rotkreuz, Switzerland) are listed in Table 1.

Western blot

Liver proteins were extracted from four rats in each group at 7 days of age and were subjected to reducing SDS-PAGE using 10% Tris gels. Proteins were then electroblotted from the gels on Nitrocellulose membranes (Amersham Hybond, GE Healthcare, Glattbrugg) and probed with a PEPCK2 primary antibody (1:1000; Cell Signaling, Beverly, MA, USA) or a G6Pase primary antibody (1:6000, as previously reported [21]), followed by rabbit anti-rat horseradish peroxidase-conjugated secondary antibody. ECL mediated by horseradish peroxidase was developed with the ECL kit and detected with Amersham hyperfilm (Amersham ECL Plus Western Blotting Detection Reagents, GE Healthcare, Glattbrugg).
Pancreatic insulin content

At PND7, total pancreases were removed, frozen in liquid nitrogen and stored at −80°C. Tissues were later pulverized, resuspended in cold acid ethanol, and left at 4°C for 40 h, with sonication every 24 h. Insulin content in the acid ethanol supernatant were determined with a rat insulin RIA kit (Linco, Labodia, Yens, Switzerland).

Corticosterone, leptin, insulin and C-peptide levels were recuperated and frozen for later dosage. Corticosterone and leptin levels were assessed in gestating dams respectively by immunoassay (Corticosterone AC-4F1, IDS, Boldon, UK), and radioimmunoassay (Linco Research, St Charles, MO, U.S.A.). Insulin and C-peptide levels were assessed in pups using the 10–1250 rat insulin or 10–1251 rat ultrasensitive insulin and 10–1172 rat C-peptide Elisa assays (Merckodia, Uppsala, Sweden).

Statistics

Results are expressed as mean ± SEM. The unpaired Student’s t-test and repeated-measures one-way analysis of variance (ANOVA) were used for comparison between groups of rats when appropriate. These tests were performed with SYSTAT 10.01 (SPSS, Chicago, IL). A p value of <0.05 was considered statistically significant.

Results

Maternal weights, leptin and corticosterone levels

Gestation of control dams (CON) was monitored in parallel to those of dams exposed to dexamethasone (DEX) or undernutrition (UN). Total weight gain during gestation of CON dams was more than 100 g (Fig. 1A). In contrast, DEX dams exhibit an arrest in those of dams exposed to dexamethasone (DEX) or undernutrition (UN). Total weight gain during gestation of CON dams was more than 100 g (Fig. 1A). In contrast, DEX dams exhibit an arrest in weight. Blood samples were collected by tail puncture for immediate glycaemia measurement using a glucometer (Actrapid, Novo Nordisk, Bagsvaerd, Denmark) per g of body weight. For the insulin tolerance test 7 days later, at PND28, the same rats were fasted for 3 h before receiving an i.p. injection containing 2 mg glucose (Sigma Aldrich, Buchs, Switzerland).

Table 1. PCR primers for quantitative real-time PCR.

| Tissue | Gene | Forward | Reverse |
|--------|------|---------|---------|
| Liver  | Pck1 | 5′-ACAACTGGTGCTTGGCTCTCA-3′ | 5′-GGTAATGTAGCGCCGTTGCTTGC-3′ |
| Liver  | G6pc | 5′-ATCTACCTTGCGGCTCTTCTT-3′ | 5′-SGGGCACAGCCAGGATCAAGTGC-3′ |
| Islets | Foxa2 | 5′-CCCGGATCCGCTCCTCTAC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Islets | Pdx1 | 5′-CCGGTGGCTGCTTCCCTTTT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Islets | Onecut1 | 5′-TCGGCGTGCCCTTTAG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Islets | Neurog3 | 5′-GAAGCTAGGATGCGGCTCAT-3′ | 5′-CCGGCAGACAGGATCAAGTGC-3′ |
| Islets | Nkx6-1 | 5′-CCCGGATCCGCTCCTCTAC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Islets | Pax6 | 5′-CCGACCACCTACATACCGGTGT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Islets | Mafa | 5′-CTTCTCCTTGACAGGTCCCG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Islets | Ins1 | 5′-CCGACCACCTACATACCGGTGT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Islets | Gcg | 5′-CCGCTGCTCTCAAGATTGT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Gata2 | 5′-AATCGGCGCTCTCAACAG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Dlk1 | 5′-CTGCACTGACCACCCCCATTCGTC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Gata2 | 5′-AATCGGCGCTCTCAACAG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Egr2 | 5′-TGGCTGGAGAGATGCTAT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Klf5 | 5′-GTCCGATACACACAGGAAGTAC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Cebpa | 5′-AGGCTGATAGHACAGCCACG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Pparg | 5′-CTGCACTGACCACCCCCATTCGTC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Srebfl | 5′-CATGCACATACCTGCTTCACA-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Lep | 5′-ATTTTCCACACACGACCCTCGGTAT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Adipoq | 5′-AAGGGAGACGCCAGGATGCTT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Tnf | 5′-GACCCCTACATCAGATCCTCTTC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Il6 | 5′-ATAGTTTCCTACAGGGATCGTCCTG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| WAT | Il1r1 | 5′-CACGGCTAGGAGGATGATATC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| BAT | Ucp1 | 5′-TGAGCCGCTAGGCTCTTGCATT-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| BAT | Ppargc1a | 5′-CTGCACTGACCACCCCCATTCGTC-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| BAT | ADRB3 | 5′-GGCGTTGAGCCAAGAGGAG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Housekeeping | Ppia | 5′-TCACATCTCAGCAGTGGGAG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |
| Housekeeping | Rplp0 | 5′-TCCTCACTTGCGCTGAAAGTG-3′ | 5′-CTGCTGCGGGCAGAAGTGT-3′ |

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Glucose tolerance test and insulin tolerance test

For the glucose tolerance test, 21-day-old rats of each group were removed from their mother 5 h before receiving an i.p. injection containing 2 mg glucose (Sigma Aldrich, Buchs, Switzerland) per g of body weight. For the insulin tolerance test 7 days later, at PND28, the same rats were fasted for 3 h before receiving an i.p. injection containing 0.5 mL human recombinant insulin (Actrapid, Novo Nordisk, Bagsvaerd, Denmark) per g of body weight. Blood samples were collected by tail puncture for immediate glycaemia measurement using a glucometer (Glucotrend Premium, Roche Diagnostics, Rotkreuz, Switzerland).
Perinatal body weight (b.w.) in IUGR pups

Neither litter size (13.0 ± 1.2 pups/litter in CON group; 14.2 ± 0.7 in DEX group, p = 0.42, 12.5 ± 0.6 in UN group, p = 0.73) nor sex ratio of born pups (47 ± 5% of males in CON group; 54 ± 3% in DEX group, p = 0.28, 47 ± 4% in UN group, p = 0.98) were significantly different between the IUGR groups and the control group in this set of experiments (data not shown).

As expected, birth b.w. (postnatal day 0, PND0) of pups exposed in utero to maternal dexamethasone or undernutrition were significantly reduced. In fact, both male (4.92 ± 0.07 g, p < 0.0001) and female (4.69 ± 0.08 g, p < 0.0001) DEX pups, as well as male (4.75 ± 0.06 g, p < 0.0001) and female (4.41 ± 0.07 g, p < 0.0001) UN pups were lighter compared to male (5.64 ± 0.10 g) and female (5.42 ± 0.11 g) CON pups (Fig. 2A), respectively. In consequence, without taking the slight sex difference into consideration, maternal dexamethasone and undernutrition exposure induced approximately 13% and 17% limitation of intrauterine growth, respectively. No major catch-up was observed during the first week of life, since at PND7, DEX and UN pup remained 13% (p < 0.005) and 24% (p < 0.0001) significantly lighter than CON pups (Fig. 2B), respectively. At PND7, circulating corticosterone levels of DEX pups (6.9 ± 1.5 ng/ml, p = 0.77) were not different to those of CON pups (6.3 ± 0.7 ng/ml) whereas corticosterone levels of UN pups were increased (14.6 ± 2.7 ng/ml, p < 0.02) (data not shown). Finally, at the end of the suckling period (PND21), DEX and UN pups were 10% (p < 0.01) and 13% (p < 0.001) lighter than CON pups (Fig. 2C), respectively.

These results indicate that both DEX and UN IUGR pups failed to recover a normal b.w. at weaning, excluding spontaneous early postnatal catch-up. To avoid any sex and hormonal confounding interference, all the following results were obtained with male pups only.

Early glucose homeostasis in IUGR pups

The hepatic gene expression of phosphoenolpyruvate carboxykinase (Pck1) and glucose-6-phosphatase (G6pc) were increased by 46 ± 19% (p < 0.05) and 101 ± 55% (p < 0.05) in DEX pups and by 53 ± 7% (p < 0.05) and 85 ± 20% (p < 0.05) in UN pups, compared to CON pups at PND7 (Fig. 3A). The corresponding protein levels were also increased; PEPCK by 59 ± 7% (p < 0.002) in UN pups and G6Pase by 24 ± 3% (p < 0.05) in DEX and by
precocious elevation in gluconeogenesis and liver glucose output in DEX pups compared to CON pups as soon as PND21. This observation was confirmed by increased area under the curve (AUC) for the entire glycaemia excursion in DEX pups (860±27 mmol/L·min, p<0.0001) and UN pups (870±26 mmol/L·min, p<0.0001) compared to CON pups (693±16 mmol/L·min) (Fig. 3D, insert panel). Fasting insulin levels were undetectable in almost all pups from the three experimental groups at PND21 (Fig. 3E, first time point). Insulin secretion was decreased 15 minutes after glucose administration in UN (0.51±0.09 ng/mL, p<0.02) compared to CON pups (0.89±0.11 ng/mL) and increased 30 minutes after glucose administration in DEX pups (Fig. 3E). Fasting C-peptide levels were unchanged in DEX (260±40 pmol/L, p = 0.23) and increased in UN (511±40 pmol/L, p<0.001) compared to CON pups (203±23 pmol/L). Fold of basal C-peptide secretion was reduced 15 and 30 minutes following glucose administration in UN but not in DEX pups (Fig. 3F). To assess the global sensitivity to insulin action, an insulin tolerance test (ITT) was performed on the same animal groups, one week later. At PND20, after 7 days of recovery and after a 3h-fast, the drops of glycaemia (Fig. 3G) in response to insulin administration were attenuated in DEX pups compared to UN and CON pups, suggesting a reduction in insulin sensitivity in relation to prenatal dexamethasone but not to undernutrition.

These results demonstrate that both DEX and UN pups present an early increase in gluconeogenesis and glucose intolerance. UN pups exhibit a decrease in glucose-induced insulin secretion whereas DEX pups exhibit a precocious decrease in insulin sensitivity.

Early endocrine pancreatic morphology in IUGR pups

In order to evaluate the impact of IUGR modelling on endocrine pancreas development, morphometric pancreatic analysis were performed in DEX and UN pups at PND7 (Fig. 4). The pancreatic endocrine area was not modified in IUGR pups when reported to b.w. (data not shown) but the ratio of endocrine to whole pancreas area was significantly reduced both in DEX (1.33±0.08%, p<0.03) and UN (1.18±0.11%, p<0.005) pups compared to CON pups (1.66±0.10%) (Fig. 4A). The reduction of endocrine pancreas development was related to a concomitant decrease in the number of islets (62.2±5 islets/section, p<0.01 for DEX and 52±4, p<0.0001 for UN pups compared to 82±4 for CON pups) (Fig. 4B) and in the mean islet size (1462±66 μm², p<0.0001 for DEX and 1612.79 μm², p<0.02 for UN compared to 1917±80 μm² for CON pups) (Fig. 4C). Quantitatively, the number of small islets (300–5000 μm²), medium islets (5000–10000 μm²) and large islets (>10000 μm²) per pancreatic section was reduced in DEX and UN pups compared to CON pups in each size category (Fig. 4D). Likewise, when expressed relatively to pancreatic surface unit, the number of small islets (6.88±0.37/mm², p = 0.36 for DEX and 5.62±0.19/mm², p<0.001 for UN pups), medium islets (5.30±0.06/mm², p<0.03 for DEX and 0.27±0.03/mm², p<0.01 for UN pups) and large islets (0.05±0.01/mm², p<0.02 for DEX and 0.80±0.03/mm², p = 0.06 for UN pups) remained globally lower in IUGR pups compared to CON pups (7.37±0.3 small islets/mm², 0.33±0.06 medium islets/mm² and 0.21±0.06 large islets/mm²) (data not shown).

These results demonstrate that both DEX and UN pups presented an altered endocrine pancreatic development due to a concomitant reduction in the size and number of Langerhans islets.

Figure 2. Postnatal body weight in IUGR pups. Body weight at birth (postnatal day 0, PND0) (A), at PND7 (B) and at PND21 (C) of male and female control pups (CON), prenatally dexamethasone-exposed pups (DEX) and prenatally undernourished pups (UN). Number of pups considered is indicated below each corresponding bar. * p<0.0001 for DEX vs. CON group and # p<0.0001 for UN vs. CON group (panel A), * p<0.005 for DEX vs. CON group and # p<0.0001 for UN vs. CON group (panel B), * p<0.01 for DEX vs. CON group and # p<0.001 for UN vs. CON group (panel C). doi:10.1371/journal.pone.0050131.g002

38±3% (p<0.01) in UN pups (Fig. 3B and 3C), suggesting a precocious elevation in gluconeogenesis and liver glucose output in these two IUGR rat models. This early defect in the control of glucose homeostasis was substantiated by elevated basal glycaemia (observed before weaning, at PND21, suckling-out), in the DEX (6.9±0.1 mmol/L, p<0.005) and the UN pups (6.8±0.1 mmol/L, p<0.02) when compared to the CON pups (6.2±0.2 mmol/L) (Fig. 3D, first time point). Moreover, the glycaemia during a glucose tolerance test (GTT) was also increased at the different time points in DEX and UN pups compared to CON pups (Fig. 3D), suggesting an attenuated glucose tolerance in IUGR pups as soon as PND21.

Early Metabolic Defects in IUGR Rats
Early pancreatic hormone production in IUGR pups

To extend these morphological investigations, we first studied the transcriptional changes in pancreatic islets isolated from DEX and UN pups at PND7 with a special focus on transcription factors known to orchestrate the beta- and alpha-cell differentiation process.

Whereas no change in HNF-3β (Foxa2) gene expression was observed in pancreatic islets of IUGR pups (Fig. 5A), Pdx1, Nkx6-1, Pax6 and Mafa gene expression were all specifically decreased in UN islets compared to CON islets (Fig. 5A). A trend to increased gene expression for Neurogenin3 (Neurog3) and HNF6 (Onecut1), its upstream regulator, was also observed in UN islets compared to CON islets (Fig. 5A), suggesting a selectivity in these transcriptional modifications. In accordance with the gene repression of the main endocrine transcription factor, we measured significant decrease in both insulin (Ins1) and glucagon (Gcg) mRNA levels in islets from UN pups, indicating an alteration in alpha- and beta-cell activity following prenatal undernutrition exposure. The mRNA levels of the glucose transporter 2 (Glut2), Kir6.2, a subunit of the K<sub>ATP</sub> channel, and the enzyme glucokinase, all classical beta-cell marker genes, were also reduced by 60% to 65% in UN islets (data not shown).

Transcriptional changes were less pronounced in islets from DEX pups (Fig. 5A), in line with unaltered secretory capacity. Immunolabelling of insulin and glucagon positive cells (Fig. 5B) showed no visual evidence of disorganization in IUGR islets (Fig. 5B). Glucagon positive cells represented 21 ± 6% of the islet area in both CON and DEX pups and 26 ± 1% in UN pups (p = 0.10 vs CON), suggesting a downward trend in beta-cell/alpha-cell proportion in UN pups. Whole pancreas insulin content (12.6 ± 1.0 mg for DEX, 13.7 ± 1.2 mg for UN, p < 0.01, compared to 20.3 ± 1.5 mg for CON pups) (Fig. 5C) corroborates the morphological alterations previously depicted in Fig. 4.

Taken together, these results demonstrate a perinatal developmental defect of the pancreatic endocrine cells in IUGR pups.

Early adipose tissue development in IUGR pups

The early suckling period constitutes the most active period for adipose tissue development in rats [24]. We investigated adipose tissue deposition at PND21 in DEX and UN IUGR pups. As shown in Fig. 6A, DEX pups present the same amount of epididymal white adipose tissue (eWAT) (77 ± 10 mg, p = 0.65) as CON pups (71 ± 7 mg) whereas UN pups show reduced eWAT accretion (44 ± 5 mg, p < 0.01) (Fig. 6A). These observations

![Figure 3. Glucose homeostasis in IUGR pups. Phosphoenolpyruvate carboxykinase (Pck1) and glucose-6-phosphatase (G6pc) mRNA (A) and protein (B and C) levels in liver at PND7, circulating glucose (D), insulin (E) and C-peptide (F) levels during a GTT at PND21, relative glycaemia during an ITT at PND28 (G) of male control pups (CON), prenatally dexamethasone-exposed pups (DEX) and prenatally undernourished pups (UN). N = 6 male rats per group for panel A, N = 4 male rats per group for panel B and C, N = 12-15 male rats per group for panel D and G, N = 10–12 male rats per group for panel E and F. For all panel, * p ≤ 0.05 or indicated value for DEX vs. CON group, # p ≤ 0.05 for UN vs. CON group and $ p ≤ 0.05 for DEX vs. UN group. doi:10.1371/journal.pone.0050131.g003

Early metabolic defects in IUGR rats
remained true when eWAT mass was reported to b.w. (0.18±0.01%, p = 0.86 for DEX and 0.11±0.01%, p < 0.0001 for UN compared to 0.18±0.01% for CON pups). No major change was observed in DEX and UN white adipocyte size and morphology (data not shown) but the analysis of gene expression revealed several transcriptional modifications, mainly in UN pups. Among preadipocyte markers, Pref-1 (Dlk1) was over-expressed in eWAT from DEX and UN pups (Fig. 6B). Gene expression of Krox-20 (Egr2) and KLF5 (Klf5), early stage adipogenesis regulators, were both reduced in eWAT from UN pups compared to CON pups (Fig. 6B). Inversely, gene expression of C/EBP-α (Cebpα), PPAR-γ (Pparg) and SREBP-1c (Srebfl), more late stage adipogenesis regulators, were all increased in eWAT from UN compared to eWAT from CON pups (Fig. 6B). Among adipokines, gene expression of leptin (Lep) and adiponectin (Adipoq) were elevated in eWAT from DEX and UN pups whereas gene expression of TNF-α (Tnf) and IL-6 (Il6) remained unchanged and gene expression of IL-1Ra (Il1rn) was decreased in eWAT from UN pups (Fig. 6C). No statistical change in the amount of interscapular brown adipose tissue (iBAT) was observed in DEX (169±15 mg, p = 0.46) and UN (185±10 mg, p = 0.09) IUGR pups compared to CON pups (153±15 mg) (Fig. 6D).

Discussion

Our present study attempted to better describe and understand the early metabolic defects in IUGR rat pups, modeled either by prenatal exposure to synthetic glucocorticoid or through prenatal undernutrition. As expected, birth b.w. of male and female pups from DEX and UN dams were severely lowered (13 to 17%), mimicking the human situation in which IUGR is defined as a fetal weight below the 10th percentile for gestational age. Early postnatal accelerated weight gain is clinically observed in IUGR newborns during infancy [25], [26]. This catch-up growth is considered as one major mechanism contributing to their predisposition to later develop metabolic disorders [27–30]. Thus, rodent models could be useful to dissociate the noxious effects of low birth weight per se to those of a subsequent catch-up growth. In our present study, both DEX and UN IUGR pups remained lighter than control pups until PND21 (weaning). Our post-weaning observations, outside the field of the present study, showed however that DEX rats remained permanently lighter than control rats, in line with a previous report [20], whereas UN
rats recovered a normal b.w after 4–6 weeks of chow diet consumption without any hyperphagia (data not shown). The absence of early catch-up in our UN pups could be linked to 1) the stringency of prenatal maternal undernutrition used (moderate food restriction allows pups to recover normal b.w at weaning [31], [32] contrary to severe food restriction [8], [19]), 2) to poor nutrition provided by remaining underweight postpartum DEX and UN mothers nevertheless eating the same amount of food as CON and 3) to the strain of rats used (Sprague-Dawley rats exhibit a slightly delayed catch-up growth phase compared to Wistar rats [33]).

Therefore, we used a combination of physiological, morphometric and transcriptomic approaches to study pancreatic islet and adipose tissue development during the early postnatal period to confirm that DEX and UN IUGR rats remain prone to early metabolic defects independently of any catch-up growth.

We initially report that both DEX and UN pups present early elevation in circulating glucose levels at PND21, just before weaning. This basal hyperglycaemia is substantiated by an increase in gluconeogenesis demonstrated by hepatic overexpression and increased protein content of G6Pase and PEPCK in DEX and UN pups as soon as PND7. This precocious elevation in gluconeogenesis seems permanent since it was also reported in adult offspring prenatally exposed to dexamethasone [15]. Dexamethasone-induced programming of later hyperglycaemia was also previously demonstrated to be fetus-focused [34], sex-dependent [35] and transgenerational [36]. Mechanistically, this programming process involved elevated Hnf4a gene expression due to a shift from its fetal to adult promoter utilization occurring in the liver periportal zone [37]. We also presently report that DEX and UN pups show an early intolerance to glucose administration observable as soon as PND21. Once again, this early defect mirrors the decreased glucose utilization observed in adult offspring prenatally exposed to dexamethasone [35] or undernutrition [32], [38].

Additional causes for the early impaired glucose utilization, beyond shared elevation of gluconeogenesis, appeared to be model-specific for DEX and UN pups. In fact, DEX pups present early attenuated insulin sensitivity, assessed by ITT, compared to UN pups. ITT did not allow to decipher tissular origin of insulin resistance and is less sensitive than standard euglycaemic hyperinsulinemic clamps used in older IUGR rodents [2], [39], [40] but are technically unsuitable in frail IUGR pups. Nevertheless, our early observations corroborate data from older animals demonstrating both insulin resistance in prenatal dexamethasone exposed rats [41] and improved insulin sensitivity at adulthood in perinatally malnourished animals devoid of catch-up growth [42]. Taken together, our physiological measurements demonstrate that impaired glucose homeostasis, characteristic of metabolic pro-

Figure 5. Endocrine pancreas hormone production in IUGR pups. mRNA levels of transcription factors and hormones in pancreatic islets (A), pancreatic immunolabelling of insulin (green) and glucagon (red) (B), pancreatic insulin content (C) of male control pups (CON), prenatally dexamethasone-exposed pups (DEX) and prenatally undernourished pups (UN) at PND7. N = 3–8 pools of 3 male rats per group (panel A), N = 5–6 male rats per group (panel C). * p<0.05 for DEX vs. CON group, # p<0.05 for UN vs. CON group for all panels. Dotted white lines indicate the delimitation of the pancreatic tissue in panel B. Note the lower density and reduced size of islets from DEX and UN pups in panel B. doi:10.1371/journal.pone.0050131.g005
gramming, is already present around weaning in IUGR pups, before any sign of catch-up growth. At tissular level, development of the endocrine pancreas in UN pups seems differently affected than in DEX pups. In fact, if a reduction in number and size of pancreatic islets as well as in insulin content is observed in both models, in agreement with previous observations [9], [12], [43], [44], transcriptomic alterations of hormones and endocrine transcription factors were more marked in UN pups in line with their selective defects in glucose-induced insulin secretory capacity. Exacerbated glucocorticoid signaling during pancreas development, extensively investigated [13], [14], [17], could be only one of the cause of IUGR pancreatic islet impairment. Differences in IUGR pup islet development could also be linked to the prenatal availability of essential nutrients. In this way, maternal low protein supply induces an increase in beta-cell apoptosis in progeny and a decrease in beta-cell proliferation whereas low caloric diet decreases beta-cell neogenesis [12], in agreement with the transcriptional alterations in regulators of pancreatic endocrine cell lineages that we observed in UN pups. A developmental delay in maturation of beta-cell precursors could be a possible etiology for reduced beta-cell mass in UN rat pups as substantiated by downregulation of Mafa and mature markers of beta-cell lineage in UN islets.

We finally investigated the impact of IUGR modeling regarding pup adipose development at PND21 since the suckling period is critical concerning the ontogeny of this metabolic tissue in rodents [24]. First, we observed no change in eWAT deposition in DEX pups but a specific decrease in eWAT mass (even reported to b.w.) in UN pups compared to control pups. Interestingly, this reduction in adipose tissue deposition was accompanied by an increased gene expression of the preadipocyte marker Pref-1 (Dlk1) in eWAT from UN pups. This over-expression could reflect either an increase in the pool of preadipocytes already present in the eWAT from UN pups at weaning, providing a cellular basis to the obesity–prone status of UN pups at adulthood, or a developmental delay in a fraction of the preadipocyte pool. Early programming of
a mature adipose tissue mass due to an adverse prenatal environment is still debated. Some reports refute a change in adipocyte proliferation [45] or glucose uptake [5] in offspring prenatally malnourished, suggesting that alterations in fat accretion is secondary to other metabolic defects involving neuroendocrine or pancreatic functions. Conversely, other studies relate primary alterations in WAT of IUGR offspring. In fact, a global up-regulation of genes involved in nutrient metabolism and adipocyte differentiation [46], a higher precocious proliferative capacity [47] and a higher glucose uptake [48] were observed in adipocytes isolated from IUGR pups, Over-expression of C/EBP-α (Cebpa), PPAR-γ (Pparg), SREBP-1C (Sreb1), later stage adipogenesis regulators detected in eWAT from UN pups at PND21 are in line with up-regulation of PPAR-γ2 observed as soon as PND1 and upregulation of SREBP-1C observed at adulthood in male IUGR rats [49]. These results explain that primary changes in the adipogenesis program, prior to the development of obesity in IUGR pups, could be initiating mechanisms involved in the so-called thrifty phenotype programming.

Adipose tissue could play an important role in linking poor fetal growth to later onset of adult diseases not only through its own over-development, but also through its secretion of numerous hormones (adipokines) modulating the metabolism. Clinical data suggest that differential regulation of adipokines in IUGR children may influence the risk for development of chronic diseases later in life [50]. In our UN model, we observed that despite eWAT depot hypotrophy, circulating leptin levels were not altered which could be explained by the increased leptin gene expression observed. A previous study has shown that IUGR pups present increased plasma leptin levels at weaning [10], suggesting that disproportionation production of leptin relative to fat mass could play a role in the etiology of metabolic programming, as previously suggested by Vickers et al [19]. Among other adipokines investigated, the expression of adiponectin (Adipoq) was also elevated in eWAT from DEX and UN pups, possibly to enhance fatty acid use. Gene expression of pro-inflammatory cytokines such as TNF-α (Tnf) and IL-6 (Il6) remained unchanged, demonstrating that adipose transcriptional alterations in UN pups at PND21 were specific and not associated with an early inflammatory state. This observation was also confirmed by the down-expression in eWAT from UN pups of IL-1 receptor antagonist (Il1rn), an acute-phase adipokine intervening in the counterregulation of the inflammatory processes [51] and characterized as a good marker of adiposity [52], [53]. Finally, the slight relative increase in brown adipose tissue deposition observed in UN pups at weaning, associated with increased gene expression of PGC-1α (Ppargc1a) and β3-adrenoreceptor (Adrb3) could contribute to elevate thermogenesis and maintain body temperature more efficiently in these frail pups, in line with a recent report describing an over-expression of UCPI1 associated with a white adipose tissue brown-like phenotype in perinatally food restricted pups [54].

In conclusion, we modeled IUGR rat pups through maternal dexamethasone or undernutrition exposure. In the absence of catch-up growth before weaning, DEX and UN pups presented some similar early metabolic defects, such as basal hyperglycaemia and decreased tolerance to glucose. Model-specific early metabolic defects were also observed, since DEX pups presented decreased insulin sensitivity whereas UN pups exhibited more marked defects in glucose-induced insulin secretion and gene expression of regulators of pancreatic islet and adipose tissue development which can be considered as mechanistic basis of the metabolic programming. Finally, the metabolic syndrome can be detected early in IUGR pups, independently of any catch-up growth, and with model-specific features.

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Author Contributions

Conceived and designed the experiments: ES DMV MLA PSH VMS. Performed the experiments: ES DMV AG AT PCR PK PM. Analyzed the data: ES DMV VMS. Wrote the paper: ES.

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