Acute nutritional stress during pregnancy affects placental efficiency, fetal growth and adult glucose homeostasis

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ABSTRACT

Exposure to maternal malnutrition impairs postnatal health. Acute nutritional stress is less clearly implicated in intrauterine programming.

We studied the effects of stressing pregnant mothers on perinatal growth and adult glucose homeostasis. We compared one group ("stressed", mothers fasted for 16 hours) with controls ("unstressed"). We found that fasting stress had adverse effects on the weight of the fetuses conceived (p<0.005) and the placental efficiency (p<0.001) in stressed compared to unstressed offspring. Placental weight was increased (p<0.001) presumably in compensation.

Stress affected the glucose homeostasis of the offspring when they became adults (p<0.005) when analysed as individuals.

We previously linked nutritional stress throughout pregnancy with a mitochondrial stress response. We modelled placenta with cultured human trophoblast cells (BeWos) and fetal tissues with mouse embryonic fibroblasts (MEFs). High throughput imaging showed that the mitochondria of both cell types underwent a similar sequence of changes in morphology, induced by nutritional stresses.

The contrasting stress responses on fetal and placental weight were not captured by the cellular models. The stress of maternal fasting may be an important determinant of perinatal outcome in the mouse and might be relevant to nutritional stress in human pregnancy.

INTRODUCTION

Fetal growth is a major determinant of perinatal mortality and postnatal health [1]. The Dutch famine [2] and subsequent studies [3, 4] have shown that aspects of intrauterine nutrition may reduce placental “efficiency” (the ratio of birth weight to placental weight) [5] and cause adverse phenotypes in postnatal life, such as metabolic syndrome. The effect of other types of stress on human offspring has not been easy to assess because it is frequently confounded by other factors such as maternal diet, smoking and social deprivation. Second-trimester maternal psychological stress increases the risk of infants being born small for gestational age [6]. In rodent studies, the effects of repeated stress on fetal growth may similarly be confounded with the effects of the maternal weight loss that it causes. Nevertheless, animal experiments demonstrate that in utero stress can permanently reprogram the hypothalamic-pituitary axis (HPA), and this can have significant post-natal neuro-behavioural sequelae [7].

Effective fetal growth requires adequate maternal nutrition coupled to active transport of nutrients across the placenta, which, in turn requires ATP much of which is generated by mitochondria. We previously developed a mouse model to explore the links between mitochondrial...
function and maternal protein intake in programming fetal growth and post-natal glucose homeostasis [8]. In order to emulate maternally inherited mitochondrial dysfunction affecting both mother and offspring, we used the anti-AIDS drug, zidovudine (AZT) that inhibits mitochondrial DNA (mtDNA) replication and hence causes mtDNA depletion [8]. We also administered a low protein diet (LPD), which has been linked to insulin resistance in later life. We compared pregnant dams exposed to one of three treatments: 1) LPD 2) AZT on a normal protein diet (so NPDAZT) 3) AZT and LPD (LPDAZT) with each other and with controls (NPD). All three treatments reduced neonatal weight. Further, AZT increased postnatal fasting glucose and the mean beta-cell area/islet was reduced in the LPD + AZT group compared with controls. Our major conclusion was that mitochondrial dysfunction exacerbates the effect of LPD on reduced neonatal weight, impairing islet development and postnatal glucose homeostasis [8].

In a later publication we linked maternal low protein diet throughout pregnancy with a stress response in placental mitochondria [9]. The low protein diet was sufficient to reduce individual fetal fresh weights and placental dry weight. The least successful litters had low placental efficiency but raised placental ATP and mtDNA content. We suggested that these changes implicated a stress response known as stress-induced mitochondrial hyperfusion (SIMH) and involves mitochondrial elongation [10]. It is associated with reduced mitophagy (recycling of damaged mitochondrial fragments) [11] and hyperpolarisation of mitochondria. It could thus underline the increased cellular ATP that we documented in placenta of mothers on a LPD [10]. Further we previously postulated that while this SIMH may confer an epigenetic benefit acutely, it is likely to disadvantage mitochondria by impairing the quality of mtDNA in the longer term. This likely accounts for the additive effects of LPD (causing SIMH) and AZT (causing mtDNA depletion and potentially increased point mutations) on neonatal weight (reduced) and glucose homeostasis (increased fasting glucose) [8].

Knowing that fibroblast cultures respond to 8-24 hours culture in glucose-free media with this SIMH response, we sought to model this in vivo. A response such as SIMH might be able to compensate for short-term maternal nutrient deprivation in mid pregnancy but might exacerbate the adverse effects in our model. We therefore fasted mothers for 16 hours between days E12 and E13. We anticipated that SIMH would already be present in the offspring of mothers already on LPD and their compensatory response to the stress of maternal fasting might thus be affected. We assessed the effect of maternal diet and mitochondrial (dys) function on fetal and placental weights in mice at embryonic day E18 and on live offspring from parallel litters on fasting glucose aged 20 weeks. All treatments were continued throughout postnatal life, as previously [8]. We report a major effect of the stress of maternal fasting (henceforth “stress”) on fetal weights and placental efficiency as well as glucose homeostasis in the adult (20 week) offspring, comparing stressed litters to the same experiment carried out over the preceding months in which mothers were not fasted (unstressed). Stressed mothers had dramatically reduced birth rate and fetal weights, compared to controls. Furthermore, placental efficiency and glucose homeostasis in the offspring were impaired.

We previously used trophoblast-derived cultured cells to model the effects of LPD on placenta [9]. We found that amino acid deprivation lengthened trophoblast mitochondria, consistent with a mitochondrial stress response that may underlie the low dry weight, raised ATP and impaired function of placenta in this mouse model. In that study, we documented changes in placental weight and water content that were reflected by, but less extreme, in fetuses. This contrasts with the present study where the short term nutritional stress changed fetal and placental weight in opposite directions. Hence we considered the possibility that mitochondrial stress responses might differ in fetal and placental tissues. We therefore went on to use cell culture models to compare mitochondrial stress responses in cell types representative of placenta and fetus to explore potential differences between them.

RESULTS

Pregnancy rate and litter size

Mothers were fasted overnight once during pregnancy on day E12-13. Overall 9/44 (20.4%) litters reached term, which is similar to the rate for unfasted (unstressed) mothers (41/177 (23%)). Stress had no significant effect on litter size.

The average litter size was however reduced by AZT (p=0.02, Figure 1).

Fetal and placental weights

Figure 2 shows that maternal fasting increased placental weight (Figure 2A, p<0.001) and reduced fetal weight (Figure 2B, p<0.005) in the offspring of the fasted mothers (averaged over litters, so that the dam is the unit of experimentation) in all 4 conditions (see Supplementary Figures 1 and 2 for raw data). Hence we then considered placental efficiency (which we define as fetal weight divided by placental weight). In our previous study we found that summed placental and fetal weights were greatest in large litters in which individual placental and fetal weights were reduced. We found that using the ratio of placental to fetal weight partly adjusts for the effect of litter size. Figure 2C shows that placental efficiency was significantly reduced by maternal stress (p<0.001, see Supplementary Figure 3 for raw data). These effects were not explained by reduced litter size, nor by the effects of LPD and AZT.
Fasting glucose in the offspring

Postnatal adult glucose homeostasis, determined by fasting glucose was affected (Figure 3 \( p=0.002 \). ANOVA) in stressed compared to unstressed offspring, using individual mice as units (see Supplementary Figure 4 for raw data). LPD, significantly reduced and AZT increased postnatal fasting glucose, as previously \( p<0.05 \) and 0.001 respectively. As expected, the effect of gender was also significant (\( p<0.01 \), not shown), fasting glucose being higher in males. When sexes were analysed separately, the effect of stress on fasting glucose remained significant in females but not males (\( p=0.008 \) and NS respectively by ANOVA). When the analysis was repeated with litters as the unit of experimentation LPD and AZT remained significant, but stress and sex did not.

Mitochondrial response to nutritional stress in tissue culture models

Having previously linked mitochondrial stress responses with effects on placental weight and efficiency we sought to investigate this in cultured cells, representative of placenta and fetus. We modelled placenta with cultured human trophoblast cells (BeWo) and fetal tissues with MEFs in order to explore potential differences between these tissues. Both cell types were cultured in a variety of media (media deficient in glucose, amino acids or fetal calf serum for up to 3 days) and mitochondrial morphology assessed by high throughput imaging [12] (Figure 4). Neither cell type thrived in media in which amino acids had been removed, and BeWo were the more sensitive.

As previously reported [9], amino acid starvation elongated mitochondria as did low glucose, low fetal calf serum (5%) and media enriched in glutamine. Where glucose was removed from the medium and galactose used as the carbon source, mitochondria appeared fragmented compared with regular 25mM glucose. More detailed analysis showed that fragmentation at 3 days was preceded by elongation at 24-48h. Inspection of Figure 4 and the Supplementary Table 1 shows that though the starting morphology was different (BeWo mitochondria being longer) the direction of change in mitochondrial morphology was similar for both cell types. Each experiment was carried out in triplicate and a representative run is shown in Figure 4.

We were unable to detect differences in the mitochondrial response to nutritional stress over 0-3 days that might reflect the differences documented in response of placental and fetal weights to stress.

DISCUSSION

We found that the stress engendered by a 16 hour fast on E12-13 during pregnancy adversely affected fetal weight and placental efficiency, that is, ratio of fetal to placental weight (\( p<0.005 \) and \( p<0.001 \) respectively, Figure 2). Placental weight, which is known to increase in compensation for hypoxia [13], was increased (\( p<0.001 \)). It also increased fasting glucose in the offspring (\( p=0.002 \), Figure 3). These effects were not explained by reduced litter size, nor by the significant effects of LPD and AZT. While the effects of nutritional stresses on trophoblast and fetal mitochondria in tissue culture model were grossly similar (Figure 4), the effects of maternal fasting stress on placental and fetal weights were in opposite directions.

To our knowledge, this is the first study linking stress in pregnancy with all three parameters that are central to fetal programming: fetal weight, placental efficiency and adult glucose homeostasis [14]. Fielder et al [15] showed that fasting for 24 and 48 hours resulted in a significant increase in mouse placental lactogen-II on day 15 of pregnancy, and that fetal weights were significantly reduced after a 48-hour maternal fast. Animal experiments show that both repeated stress and impaired maternal nutrition are inter-related factors that can permanently affect the fetus. Stress affects maternal glucocorticoid

Figure 1: Litter size at E18 and maternal fasting. Stress did not have a simple effect on litter size. It interacted significantly with the effects of LPD and AZT on litter size. Error bars are 1SE. Data is from 5, 5, 5 and 5 unstressed and 12, 9, 12 and 14 stressed litters corresponding to NPD controls, LPD fed on low protein diet, NPDAZT treated with AZT but on normal diet, LPDAZT fed on low protein diet and treated with AZT respectively.
Figure 2: Placental weight increased and fetal weight and placental efficiency are reduced at E18 by maternal fasting. (A) ANOVA showed that placental weight is increased by maternal stress ($p<0.001$) in all 4 conditions. (B) Fetal weight is reduced by maternal stress ($p=0.005$) in all 4 conditions (averaged across the average for each litter). (C) The ratio of fetal to placenta weight (a surrogate for the efficiency of placental transport) was reduced by maternal stress ($p<0.001$). This ratio partly adjusts for the interacting effects of litter size in such studies. N=44, 14, 27 and 26 in unstressed and 34, 31, 31 and 35 in stressed offspring from 5, 5, 5 and 5 unstressed and 12, 9, 12 and 14 stressed litters corresponding to control, LPD, NPDAZT and LPDAZT respectively. NPD controls, LPD fed on low protein diet, NPDAZT treated with AZT but on normal diet, LPDAZT fed on low protein diet and treated with AZT. Error bars are 1SE. Calipers represent ANOVA F test $p$-value for effect of maternal stress, ***$p<0.001$, **$p<0.01$. 
metabolism [7] and blood pressure [16], and this in turn affects the weight and health of the offspring [17]. In our model of mitochondrial intra-uterine programming, maternal stress, caused by fasting, reduced placental efficiency. While it is clear that glucocorticoids play a key role [18, 19], there has been scanty direct evidence that a single stressful episode in pregnancy can impair placental efficiency.

Previous studies have shown that mild stresses, such as exposure to strobe light for 2 hours in mid pregnancy, can re-programme the HPA [20]. Nutritional stress may affect placental physiology [21] through this system, and this is potentially important both in animal models [16] and human disease [17]. Nevertheless maternal fasting for 16 hours on day E12-13 exerted pronounced effects on the offspring. In particular, rodents appear to have an increased appetite, doubling their food intake during pregnancy [22]. Further studies are needed to determine the critical period for the effect that we observed. Other investigators have implicated the HPA in the response to stress both in mid-gestation [23] and later, when the fetal HPA begins to release its own adrenocorticotrophic hormone (ACTH) and corticosterone [24]. In addition to mechanisms potentially involving glucocorticoids, we speculate that perfusion could be affected, as major changes in murine placental vasculature and fetal nutrition occur between day E10.5 and 12.5 [25] (corresponding to around week 11 in humans [26]).

In our previous studies we found that mitochondrial dysfunction appeared to exacerbate some of the effects of low protein diet, though not the relationship with postnatal fasting glucose. In this previous work we found that LPD decreased litter size and neonatal weight [8]. Furthermore, early exposure to AZT interacted with LPD to impair fetal development and post-natal glucose homeostasis [8]. Glucose homeostasis was impaired in both the NPDAZT, and LPDAZT groups, and beta-cell area/islet was reduced [8]. We also implicated SIMH in some of the effects of LPD on placenta (increased ATP and impaired function) [9], and conversely excessive mitochondrial fragmentation can be detrimental [27].

Our results, based on mice, are supportive of, but have substantial advantages over, studies of the effects of stress on fetal growth in humans, where stresses are inevitably confounded by other factors [29]. For instance, studies that have found strong causative associations between stress and birth weight have rarely been able to exclude effects of poor nutrition and smoking. Repeated stresses may have effects on both brain structure and
Figure 4: Similarities in the mitochondrial response of cultured cells derived from trophoblast and embryo to nutritional stresses. The figure shows stacked bar chart illustrating the distribution of mitochondrial length for 7 different media (media deficient in glucose, amino acids or fetal calf serum) in MEFs (top) and BeWo cells (bottom). In the case of amino acid free media, exposure was for 24 and 48 hours duration for BeWos and MEFs respectively, as longer exposures were poorly tolerated. All other exposures were 3 days. For each condition >1100 cells were imaged and the mean mitochondrial length for each calculated. For means of each of these distributions see Supplementary Table 1.
postnatal behaviour. In our study we were able to link 

a single episode of maternal stress at a critical stage of 

pregnancy both to perinatal parameters and potentially to 

adult health. Fasting for 16 hours is clearly very stressful 

for a mouse, much more so than an overnight fast in a 

pregnant woman. Nevertheless, our findings may be 

relevant for women who inadvertently undergo significant 

fasting in human pregnancy.

MATERIALS AND METHODS

Animals

Mice were housed in conventional wire-top polycarbonate cages, with a 12:12 light:dark cycle at 
temperatures between 19-23°C and relative humidity 55±10%. Food and water were offered ad libitum. The 
facility is free of MHV, EDIM, MVM, MPV, PVM, Sendai, TMEV, ectomelia, LCMV, Mad 1 and 2, MCMV, 
reovirus 3, Citrobacter rodentium, Clostridium piliforme, Corynebacterium kutscheri, Mycoplasma, Pasteurellaceae, 
Salmonella, beta-hemolytic streptococci, Streptococcus pneumoniae, Streptobacillus moniliformis, endoparasites 
and ectoparasites. Helicobacter and MNV are present in this facility. All animals were housed and managed 
according to the United Kingdom’s Home Office protocols, covered by the Animals (Scientific Procedures) 
Act 1986. The protocol was approved by the Oxford 

University Committee on Animal Care and Ethical 
Review, University of Oxford Medical Sciences division 
(Project licences 3001526 and 3002208). Seven-week-old 
female mice from the inbred strain C57BL/6J/OxJR (bred 
in house but originally obtained from Harlan/Envigo) were 
acclimatized over a one-week period by handling and 

recording weights. In order to assess the effects of nutritional stress 

and impaired mitochondrial function, the offspring were 
exposed to a low protein diet (8% as opposed to 20% acid 
casein based protein corresponding to LPD and NPD) and/or the mitochondrial inhibitor, AZT, in utero and for 

the rest of their postnatal lives (AZT rapidly crosses the 
human placenta [30]). Following acclimatisation, female 

mice were maintained on a regime of one of the following: 1) normal (20% protein) diet (NPD); 2) low protein diet 
-(LPD); 3) NPD and 0.15 mg/ml AZT-(NPDAZT); or 4) LPD and 0.15 mg/ml AZT (LPDAZT), from the age of 
8 weeks for 2 weeks prior to mating. Both 8% and 20% 
protein diets were obtained from Arie Blok, (Netherlands, 
catalogue numbers 4400.00 and 4400.01 respectively). 
AZT (Zidovudine, Glaxo Smithkline, UK) was mixed into the drinking water at a concentration of 0.15 mg/ml and 
this was changed twice weekly. Food and water were given ad libitum. Day 0.5 of gestation was determined 
by detection of a vaginal plug. At embryonic day 18 
some animals were sacrificed by cervical dislocation and 

the concepti collected. In an earlier publication [9] we 
previously presented data on the stressed NPD and LPD 
litters (of that publication Figure 1 shows the placental and 
fetal weights in relation to water content, and Figure 5 the 
placental mtDNA content). In work carried out previous to this study, dams were fasted overnight 3 times during pregnancy, at embryonic days 6, 12 and 18. However, only 3/44 (6.8%) litters reached term. This is significantly lower than 41/177 (23%), the rate for pregnancies in which the 
mothers were not fasted (p =0.05). Because of the effect on fecundity, the data therefore presented in this study are from mothers fasted once during pregnancy prior to 
blood sampling from a tail vein (fasting overnight on E12, 

deciforce “stressed”) or no fasting (“unstressed”). This is a critical time in pregnancy in the mouse, being soon after the onset of maternal placental vascular perfusion at day 10.5 and coinciding with substantial changes in three marker proteins for reactive oxygen species damage [31]. Baseline food and water consumption was measured just prior to mating by weighing the food and water/AZT in a cage of on average four females, and weighing again 
24 hours later. Litters were culled to 4-5 pups by 4 days 
of age. At 20 weeks, the remaining offspring were fasted 
overnight prior to sampling under terminal anaesthesia (sodium pentobarbital (Sagatal, Rhone Merieux, Harlow, 
UK) 60 mg/kg (i.p.)) from the tail vein for a fasting glucose measurement.

Cell biology methods

BeWo cells [32] and mouse embryonic fibroblasts 
[27] in which mitochondria were fluorescent (DS-red 
targeted to mitochondria [32]) were cultured in a 96-well 
plate and treated for 6 hours in the indicated conditions 
before fixation with 4% paraformaldehyde (PFA). After 
DAPI staining, the plate was imaged using the InCell1000 
analyser (GE healthcare life sciences, 500 cells per 
well). Raw images were binarised and mitochondrial 
morphological characteristics were quantified, notably the degree of branching or mitochondrial form factor (FF) and the mean mitochondria length (in μm). FF is defined as (Pm²)/(4πAm), where Pm is the length of mitochondrial 
outline and Am is the area of mitochondrion [33].

Statistical analysis

We used SPSS version 22, with the help of the 
University of Oxford Statistics Department. Analysis of 
perinatal parameters (placental and fetal weights) was 
carried out using ANOVA, with dams as the unit of 
experimentation. In line with previous investigators in this 
field, analysis of postnatal fasting glucose was carried out 
on an individual basis.
Abbreviations
AZT: zidovudine; HPA: hypothalamic-pituitary axis; LPD: low protein diet; MEFs: mouse embryonic fibroblasts; MtDNA: mitochondrial DNA; NPD: normal protein diet; SIMH: stress-induced mitochondrial hyperfusion.

Author contributions
S Malik, A Diot, K Morten and E Dombi carried out laboratory work. M Vatish helped draft the paper, CAR Boyd and J Poulton raised the grant funding, designed the study, supervised the laboratory work, analysed the and drafted the paper.

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CONFLICTS OF INTEREST
The authors declare no conflicts of interest.

REFERENCES
1. Armitage JA, Taylor PD, Poston L. Experimental models of developmental programming: consequences of exposure to an energy rich diet during development. J Physiol. 2005; 565:3-8.
2. Roseboom TJ, van der Meulen JH, Ravelli AC, Osmond C, Barker DJ, Bleker OP. Effects of prenatal exposure to the Dutch famine on adult disease in later life: an overview. Mol Cell Endocrinol. 2001; 185:93-8.
3. Hales CN, Barker DJ. Type 2 (non-insulin-dependent) diabetes mellitus: the thrifty phenotype hypothesis. Diabetologia. 1992; 35:595-601.
4. Fleming TP, Velazquez MA, Eckert JJ. Embryos, DOHaD and David Barker. J Dev Orig Health Dis. 2015; 6:377-83.
5. Lumey LH. Compensatory placental growth after restricted maternal nutrition in early pregnancy. Placenta. 1998; 19:105-11.
6. Khaskan AS, Everard C, McCowan LM, Dekker G, Moss-Morris R, Baker PN, Poston L, Walker JJ, Kenny LC. Second-trimester maternal distress increases the risk of small for gestational age. Psychol Med. 2014; 44:2799-810.
7. Seckl JR, Meaney MJ. Glucocorticoid programming. Ann N Y Acad Sci. 2004; 1032:63-84.
8. Morten K, Field P, Ashley N, Williams KA, Harris D, Hartley M, Clark A, Poulton J. Fetal and neonatal exposure to AZT and low-protein diet affects glucose homeostasis: a model with implications for AIDS prevention. Am J Physiol Endocrinol Metab. 2005; 289:1115-8.
9. Chiaratti MR, Malik S, Diot A, Rapa E, Macleod L, Morten K, Vatish M, Boyd R, Poulton J. Is placental mitochondrial function a regulator that matches fetal and placental growth to maternal nutrient intake in the mouse? PLoS One. 2015; 10:e0130631.
10. Tondera D, Grandemange S, Jourdain A, Karbowski M, Mattenberger Y, Herzig S, Da Cruz S, Clerp P, Raschke I, Merkwirth C, Ehses S, Krause F, Chan DC, et al. SLP-2 is required for stress-induced mitochondrial hyperfusion. EMBO J. 2009; 28:1589-600.
11. Gomes LC, Di Benedetto G, Scorrano L. During autophagy mitochondria elongate, are spared from degradation and sustain cell viability. Nat Cell Biol. 2011; 13:589-98.
12. Diot A, Hinks-Roberts A, Lodge T, Liao C, Dombi E, Morten K, Brady S, Fratter C, Carver J, Muir R, Davis R, Green CJ, Johnston I, et al. A novel quantitative assay of mitophagy: combining high content analysis fluorescence with mitochondrial DNA mutant load to identify novel pharmacological modulators of mitophagy. Pharmacol Res. 2015; 100:24-35.
13. Matheson H, Veerbeek JH, Charnock-Jones DS, Burton GJ, Yung HW. Morphological and molecular changes in the murine placenta exposed to normobaric hypoxia throughout pregnancy. J Physiol. 2016; 594:1371-88.
14. Fowden AL, Sferruzzi-Perri AN, Coan PM, Constancia M, Burton GJ. Placental efficiency and adaptation: endocrine regulation. J Physiol. 2009; 587:3459-72.
15. Fielder PJ, Ogren L, Edwards D, Talamanes F. Effects of fasting on serum lactogenic hormone concentrations during mid- and late pregnancy in mice. Am J Physiol. 1987; 253:E40-4.
16. Igosheva N, Klimova O, Anishchenko T, Glover V. Prenatal stress alters cardiovascular responses in adult rats. J Physiol. 2004; 557:273-85.
17. Benediktsson R, Lindsay RS, Noble J, Seckl JR, Edwards CR. Glucocorticoid exposure in utero: new model for adult hypertension. Lancet. 1993; 341:339-41.
18. Vaughan OR, Fisher HM, Dionelis KN, Jefferies EC, Higgins JS, Musial B, Sferruzzi-Perri AN, Fowden AL. Corticosterone alters materno-fetal glucose partitioning.
and insulin signalling in pregnant mice. J Physiol. 2015; 593:1307-21.

19. Vaughan OR, Phillips HM, Everden AJ, Sferruzzi-Perri AN, Fowden AL. Dexamethasone treatment of pregnant F0 mice leads to parent of origin-specific changes in placental function of the F2 generation. Reprod Fertil Dev. 2015; 27:704-11.

20. Kapoor A, Matthews SG. Prenatal stress modifies behavior and hypothalamic-pituitary-adrenal function in female guinea pig offspring: effects of timing of prenatal stress and stage of reproductive cycle. Endocrinology. 2008; 149:6406-15.

21. Mairesse J, Lesage J, Breton C, Breant B, Hahn T, Darnaudery M, Dickson SL, Seckl J, Blondeau B, Vieau D, Maccari S, Viltart O. Maternal stress alters endocrine function of the feto-placental unit in rats. Am J Physiol Endocrinol Metab. 2007; 292:E1526-33.

22. Makarova EN, Kochubei ED, Bazhan NM. Regulation of food consumption during pregnancy and lactation in mice. Neurosci Behav Physiol. 2010; 40:263-7.

23. Mueller BR, Bale TL. Impact of prenatal stress on long term body weight is dependent on timing and maternal sensitivity. Physiol Behav. 2006; 88:605-14.

24. Boudouresque F, Guillaume V, Grino M, Strbak V, Chautard T, Conte-Devolx B, Oliver C. Maturation of the pituitary-adrenal function in rat fetuses. Neuroendocrinology. 1988; 48:417-22.

25. Georgiades P, Ferguson-Smith AC, Burton GJ. Comparative developmental anatomy of the murine and human definitive placentae. Placenta. 2002; 23:3-19.

26. Jauniaux E, Watson A, Burton G. Evaluation of respiratory gases and acid-base gradients in human fetal fluids and uteroplacental tissue between 7 and 16 weeks' gestation. Am J Obstet Gynecol. 2001; 184:998-1003.

27. Liao C, Ashley N, Diot A, Morten K, Phadwal K, Williams A, Fearnley I, Rosset L, Lowndes J, Fratter C, Ferguson DJ, Vay L, Quaghebeur G, et al. Dysregulated mitophagy and mitochondrial organisation in optic atrophy due to OPA1 mutations. Neurology. 2017; 88:131-42.

28. Ozanne SE, Hales CN. Early programming of glucose-insulin metabolism. Trends Endocrinol Metab. 2002; 13:368-73.

29. Landrigan PJ, Forman J, Galvez M, Newman B, Engel SM, Chemtob C. Impact of September 11 World Trade Center disaster on children and pregnant women. Mt Sinai J Med. 2008; 75:129-34.

30. Bennetto-Hood C, Bryson YJ, Stek A, King JR, Mirochnick M, Acosta EP. Zidovudine, lamivudine, and nelfinavir concentrations in amniotic fluid and maternal serum. HIV Clin Trials. 2009; 10:41-7.

31. Aiken CE, Cindrova-Davies T, Johnson MH. Variations in mouse mitochondrial DNA copy number from fertilization to birth are associated with oxidative stress. Reprod Biomed Online. 2008; 17:806-13.

32. Kudo Y, Boyd CA, Kimura H, Cook PR, Redman CW, Sargent IL. Quantifying the syncytialisation of human placental trophoblast BeWo cells grown in vitro. Biochim Biophys Acta. 2003; 1640:25-31.

33. Mortiboys H, Thomas KJ, Koopman WJ, Klaffke S, Abou-Sleiman P, Olpin S, Wood NW, Willems PH, Smeitink JA, Cookson MR, Bandmann O. Mitochondrial function and morphology are impaired in parkin-mutant fibroblasts. Ann Neurol. 2008; 64:555-65.