Maternal Diet-Induced Obesity Alters Mitochondrial Activity and Redox Status in Mouse Oocytes and Zygotes

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Abstract

The negative impact of obesity on reproductive success is well documented but the stages at which development of the conceptus is compromised and the mechanisms responsible for the developmental failure still remain unclear. Recent findings suggest that mitochondria may be a contributing factor. However to date no studies have directly addressed the consequences of maternal obesity on mitochondria in early embryogenesis. Using an established murine model of maternal diet-induced obesity and a live cell dynamic fluorescence imaging techniques coupled with molecular biology we have investigated the underlying mechanisms of obesity-induced reduced fertility. Our study is the first to show that maternal obesity prior to conception is associated with altered mitochondria in mouse oocytes and zygotes. Specifically, maternal diet-induced obesity in mice led to an increase in mitochondrial potential, mitochondrial DNA content and biogenesis. Generation of reactive oxygen species (ROS) was raised while glutathione was depleted and the redox state became more oxidised, suggestive of oxidative stress. These altered mitochondrial properties were associated with significant developmental impairment as shown by the increased number of obese mothers who failed to support blastocyst formation compared to lean dams. We propose that compromised oocyte and early embryo mitochondrial metabolism, resulting from excessive nutrient exposure prior to and during conception, may underlie poor reproductive outcomes frequently reported in obese women.

Introduction

Obesity and related metabolic disorders are a major health issue worldwide. With increasing prevalence in all populations and age groups, the proportion of women of reproductive age who are obese is rising [1]. Evidence is growing that excessive body fat has a detrimental effect on female fertility and pregnancy [2]. Obese women take longer to conceive and have a higher risk of miscarriage compared to lean women [3]. Obesity also impairs the immediate outcome of assisted reproductive technologies suggesting that maternal body mass index (BMI) may influence the potential for fertilisation and viability of early embryos [4]. Since the earliest stages of embryo development are primarily controlled by the oocyte, it is likely that a sub-optimal environment within the ovary and/or oviduct accounts for these poor reproductive outcomes.

As recently reviewed [5], understanding of the effects of maternal obesity on the structure and metabolism in oocytes and pre-implantation embryos is very limited. It has been proposed that a high plane of nutrition might lead to excessive enrichment of embryonic metabolism. Mitochondria are likely candidates for compromised mitochondrial sources of NAD(P)H along with mitochondrial oxidative phosphorylation, a mechanism coupling the oxidation of nutrients and reducing equivalents (NAD(P)H, FADH2) with the phosphorylation of adenosine triphosphate (ATP) by oxidative phosphorylation, a mechanism coupling the oxidation of nutrients and reducing equivalents [nad/p/h, fadh2] with the phosphorylation of adenosine diphosphate. Both cytosolic and mitochondrial sources of NAD(P)H along with mitochondrial FADH2 stimulate the mitochondrial electron transport chain to pump H+ out of the mitochondrial matrix thereby hyperpolarising the inner mitochondrial membrane and generating the proton-motive force used to generate ATP. Electron donors NAD(P)H and FADH2 besides being used for energy production set the oxidative stress that is a key factor in the development of obesity.

metabolism may compromise oocyte and embryo development [7,8]. Mitochondria are likely candidates for compromised metabolism in the embryo; these organelles are exclusively maternal in origin, and thus a deleterious influence of maternal BMI on mitochondria in the oocyte would strongly influence embryonic metabolism. Mitochondria also perform numerous regulatory functions during oocyte maturation [9], fertilization, initiation and progression of preimplantation embryos [10]. As energy producer, the central and most important function of mitochondria is the synthesis of adenosine triphosphate (ATP) by oxidative phosphorylation, a mechanism coupling the oxidation of nutrients and reducing equivalents (nad/p/h, fadh2) with the phosphorylation of adenosine diphosphate. Both cytosolic and mitochondrial sources of NAD(P)H along with mitochondrial FADH2 stimulate the mitochondrial electron transport chain to pump H+ out of the mitochondrial matrix thereby hyperpolarising the inner mitochondrial membrane and generating the proton-motive force used to generate ATP. Electron donors NAD(P)H and FADH2 besides being used for energy production set the intracellular redox state. NADH oxidation in the mitochondria will produce ROS whereas NADPH oxidation (in the cytosol and mitochondria) serves to rejuvenate the antioxidant defence by reducing peroxiredoxins, thioredoxin and oxidised glutathione.
Mitochondrial functions have, therefore, a dual impact on the intracellular redox state via regeneration antioxidant systems and via ROS production [11].

Mitochondria not only supply cells with their ATP, but are also the source of cellular guanosine-5’-triphosphate (GTP) as well as site of amino acid synthesis and reservoir of cell calcium. Thus, changes in mitochondrial activity can alter cell function in a dramatic way. The importance of mitochondria in oocyte quality and embryo development is highlighted by reports showing that defects in mitochondrial biogenesis together with insufficient mitochondrial mass are associated with oocyte maturation failure and abnormal embryo development [12,13]. Both the quality and quantity of mitochondria are therefore an essential prerequisite for successful fertilization and embryo development [14].

Studies in vitro have also highlighted the susceptibility of mitochondria within the oocyte and developing embryo to environmental stressors and have shown that even low-level acquired mitochondrial injuries may persist into embryonic life [15,16]. Potential influences of maternal nutritional status in obesity are indicated by reports showing that periconceptual exposure to high energy substrates such as fatty acids [17] and proteins [18] results in perturbed oocyte and embryo mitochondrial metabolism. Hitherto, mitochondrial abnormalities of the oocyte and early embryo have not been identified as a direct consequence of maternal obesity. Using an established murine model of maternal diet induced obesity [19] and a live cell dynamic fluorescence imaging techniques coupled with molecular biology we have investigated the effects of maternal obesity on mitochondria 1 metabolism and biogenesis in oocytes and pre-implantation embryos. Our study is the first to show that maternal obesity during the periconceptional period resulted in an increased mitochondrial potential, biogenesis and damaging level of ROS in oocytes and zygotes. These changes were associated with reduced fertility and impaired embryo viability. Our findings have identified altered mitochondrial status as one of the probable mechanisms of obesity-associated reproductive and developmental failure.

Results

Maternal metabolic profile

At conception, female mice fed an obesogenic diet had a 43% increase in body weight, 2.5 fold increase in fat pad and a significantly higher concentration of serum fatty acids (p<0.05) compared with chow fed controls (Table 1). Although maternal serum leptin concentrations were similar in obese and lean mice, the concentration of leptin in the oviductal fluid was significantly elevated in obese females in comparison with lean females (p<0.05).

Mitochondrial status in oocytes and zygotes of obese female mice

Live cell dynamic fluorescence imaging was employed to study effects of maternal obesity on mitochondrial function in oocytes and embryos. This technique together with a range of targeted fluorescent probes permitted comprehensive evaluation of mitochondrial function with simultaneous measurement of multiple mitochondrial variables in a single oocyte and embryo. The common vital mitochondrial membrane-specific dyes; MitoTrackers and JC1 have been extensively used to study mitochondrial dynamics and function in oocytes and embryos. However, prolonged excitation of cells loaded with MitoTrackers may impair mitochondrial function [20] whereas the JC1 dye appears sensitive to factors other than inner mitochondrial membrane potential (ΔΨm) [21] and may inhibit mitochondrial complex 1 [22]. We therefore chose to measure ΔΨm in single oocytes and embryos using a low toxicity potentiometric fluorescent dyes-tetramethyl rhodamine methyl ester (TMRM; [23]).

Maternal diet-induced obesity led to a dramatic increase in ΔΨm in oocytes and zygotes. The intensity of mitochondrial localised TMRM fluorescence in oocytes and zygotes of obese females increased by 147% (p<0.01) and 74% (p<0.01), respectively, compared with oocytes and zygotes of lean females (Figure 1A and 1B). Eggs from obese mice also had a different

Table 1. Maternal weights and metabolic parameters.

| Parameter                  | Control females | Obese females | Significance |
|---------------------------|-----------------|---------------|-------------|
| Body weight (g)           | 22.9±1.2        | 32.8±1.3      | P<0.001 (10)|
| Fat pads weight (g)       | 0.91±0.01       | 2.32±0.01     | P<0.01 (10) |
| Serum glucose (mmol/L)    | 6.68±0.81       | 8.99±0.95     | P<0.05 (10) |
| Serum FFA (mmol/L)        | 0.67±0.06       | 0.89±0.08     | P<0.05 (10) |
| Serum triglycerides (mmol/L) | 0.93±0.08       | 0.94±0.09     | P<0.05 (10) |
| Serum lepton (pg/ml)      | 1172±270        | 1492±170      | P<0.05 (8)  |
| Oviductal lepton (pg/ml)  | 1067±293        | 1916±221      | P<0.05 (7)  |
| Oviductal glucose (mmol/L)| 6.11±0.72       | 10.90±2.01    | P>0.05 (7)  |

Data expressed as mean ± SEM. All serum measurements were fasting. Values in parentheses indicate n/group.

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Figure 1A. Maternal diet-induced obesity leads to increased mitochondrial activity in the oocytes and zygotes. The measurements of inner mitochondrial membrane potential (ΔΨm) were made using confocal imaging of TMRM fluorescence. The signal intensity was quantified per pixel in a confocal slice after thresholding to remove background signal. (A) – oocytes and (B) zygotes derived from lean n=15 cells/group and obese females (n=15 cells/group). Relative intensity of TMRM fluorescence is expressed as a percentage of the signal from oocytes of lean mice. (C) Representative confocal images of mitochondria distribution in oocytes from lean and obese mice. * p<0.05. Data are mean ± SEM.

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pattern of mitochondrial distribution as visualised by the distribution of TMRM staining (Figure 1C and 1D). Mitochondria were distributed evenly throughout the ooplasm of the eggs from lean females. In contrast, mitochondria in eggs from obese females had a discontinuous distribution with high density clusters localised to the cortical ooplasm and surrounding the nucleus. Since the organisation of mitochondria within cells is important for the signalling events associated with fertilisation [10], this abnormal localisation of mitochondria may be detrimental for the preimplantation embryo. Indeed, a similar aggregated mitochondrial structure was identified in preimplantation embryos undergoing arrest [24].

An increase in ΔΨm may have a number of origins including a higher supply of substrates and increased mitochondrial respiratory activity or inhibition of ATP synthase activity [25]. To differentiate between different mechanisms we analysed the redox state in oocytes and zygotes, measured as the autofluorescence signal derived from NAD(P)H or FAD2⁺ [23]. Confocal laser-scanning microscopy (CLSM) revealed that the NAD(P)H in oocytes and zygotes from obese mice was more oxidised than in eggs from lean mice (oocytes - 36±3% vs 53±4%, p<0.05; zygotes - 27±2% vs 86±5%, p<0.05 on a scale that runs from 0% for full oxidation to 100% for full reduction) (Figure 2B). Imaging FAD2⁺ autofluorescence also revealed that flavoproteins were more oxidised in oocytes and zygotes from obese females compared to eggs from lean females (oocytes - 74±6 vs 41±3%, p<0.05; zygotes- 56±4% vs 40±4%, p<0.05: note that for this signal, 100% is fully oxidised and 0%, fully reduced) (Figure 2B).

These changes indicate increased oxidation of the pyridine nucleotide and flavoprotein pools [23] and suggest a shift in the intracellular redox status towards oxidation in the eggs from obese mice.

Studies in mitochondria respiratory chain function in vitro [26] and in vivo [11,23] have shown that the redox state of the pyridine nucleotide and flavoprotein pools reflects the balance between the rate of reduction by substrate utilization and the rate of oxidation by mitochondrial respiration. The shift of the redox balance towards a net reduced state occurs as a consequence of up-regulated substrate processing and inhibition of respiration, whereas an increase in mitochondrial respiratory rate favours the shift of the redox potential towards the oxidised state. Therefore it is conceivable that oxidised state of the pyridine nucleotide and flavoprotein pools in oocytes and zygotes from obese females may be attributable to increased mitochondrial respiratory chain activity.

In order to evaluate the level of oxidative stress and antioxidant defence in oocytes and zygotes we measured rates of intracellular ROS generation using dihydroethidium (HE), a non-fluorescent derivative of ethidium which is oxidised to a fluorescent product by superoxide. In oocytes and zygotes of obese mice the rate of ROS production was significantly increased by 2.1 (p<0.05) and 1.6 (p<0.05) fold respectively compared to the eggs of lean mice (Figures 3A and 3B). We measured levels of the antioxidant glutathione (GSH) using the monochlorobimane (MCB) which forms a fluorescent adduct following an enzyme catalysed reaction with GSH. In parallel with the increase in the oxidative load, GSH was depleted in oocytes and zygotes from obese females when compared with eggs from lean females (oocytes – 80±27 vs 100±26, p<0.05; zygotes- 68±4% vs 93±6%, p<0.05) (Figures 3C and 3D).

Alterations in the rates of intracellular ROS generation are associated with changes in mitochondrial abundance and mtDNA copy number. Oxidative stress damages bases as well as causing single or double-strand breaks in mtDNA which are mutagenic and can inhibit mtDNA replication [27]. However, excessive ROS generation has been associated with an increase in mtDNA copy number in aging tissues as a result of a feedback response which compensates for defective mitochondria bearing impaired respiratory chain or mutated mtDNA [28].

To test the hypothesis that maternal obesity-associated oxidative stress in oocytes and zygotes may affect mitochondrial biogenesis we measured mtDNA copy number and expression of key genes involved in the regulation of the replication and transcription of the mitochondrial genome. mtDNA copy number was significantly increased in oocytes from obese compared to lean mice (Figure 4A), and the expression of nuclear genes encoding mtDNA transcription factors - mtTFAM and NRF1 - was also elevated in oocytes from obese females suggesting upregulation of mitochondrial biogenesis (Figure 4B and 4C). Interestingly, mtDNA copy number, TFAM and NRF1 expression were not altered in zygotes from obese mice.

Effects of maternal obesity on pre-implantation embryos

No significant differences were detected in the number of zygotes between lean and obese females. However, the ability of zygotes to develop to the blastocyst stage was reduced in obese mice (Table 2). Thus, the number of obese females who failed to produce blastocysts 4 days after mating was higher in comparison
with controls (5 vs 2). In three of 5 obese mice no blastocyst were found in either uterus and oviduct whereas in two obese females a small number of arrested fragmented embryos at various stages of development was recovered from the oviduct. In the remaining 6 obese (54%) females blastocyst development per mouse was similar to that in lean females. Differential nuclear labeling did not reveal any differences in the number of cells within trophectoderm and inner cell mass lineages of blastocysts from obese and lean mice (Table 2).

Discussion

The negative impact of obesity on reproductive success is well documented [5] but the stages at which development of the conceptus is compromised and the mechanisms responsible for the developmental failure remain unclear. Using our established model we have identified altered mitochondrial activity as one of the probable mechanisms of obesity-associated reproductive and developmental failure.

We report that blastocyst development was reduced in maternal diet-induced obesity and was associated with altered mitochondrial distribution and striking hyperpolarisation of the mitochondrial membrane, oxidised redox state and oxidative stress in both oocytes and zygotes. Regulation of all of these parameters is required for normal development [13]. We also report increased mitochondrial biogenesis in oocytes as evidenced by high mtDNA copy number and up-regulation of NRF1 and TFAM transcripts.

It is currently unclear what mechanisms underlie such large differences in Δψ between control oocytes and embryos and those derived from obese females. Nor is it entirely clear how these differences are established and maintained. Studies in mitochondria in somatic cells [29] and embryos [30,31] have shown that the magnitude of Δψm is related to the level of mitochondrial respiration so the differences in mitochondrial respiratory activity may account for the hyperpolarisation observed. The increased oxidation of reducing equivalents - NAD(P)H and FADH2 in oocytes and zygotes from obese females have also provided indirect evidence for activation of mitochondrial respiratory

![Figure 3. Maternal diet induced obesity increases rates of ROS generation and depletes glutathione in oocytes and zygotes.](https://example.com/figure3.png)
activity. Carbohydrates and fatty acids are the principal substrates for mitochondrial oxidation and an increased availability of these energy substrates is a well known mechanism of up-regulation of mitochondrial respiration [32]. The presence of excessive fuel within the oocyte reproductive environment may therefore increase substrate influx through the metabolic mitochondrial pathway, leading to activation of mitochondrial respiratory activity which is reflected in a hyperpolarised state of the mitochondrial membrane. Whether oocytes and embryos that show intense mitochondrial hyperpolarisation also have abnormally higher ATP content remain to be determined.

While the increase in $\Delta\psi_m$ in mouse oocytes and zygotes of obese mice may be due to increased energy substrate load to mitochondria [33], others have suggested that mitochondrial hyperpolarisation might be associated with early molecular events that precede development impairment and the induction of cell death [31]. Recently, Schienke et al. [34] has also reported reduced differentiation potential in mouse embryonic stem cells with a high $\Delta\psi_m$ and overall rate of mitochondrial metabolism.

Maternal diet-induced obesity was associated with increased oxidation of NADP/H in both oocytes and zygotes. NADP/H has direct antioxidant properties and also ensures regeneration of GSH, a developmentally critical antioxidant molecule [35]. Therefore, NADP/H down-regulation and reduced antioxidant capacity may contribute to oxidative stress in the oocytes and zygotes from obese females, as observed through direct measurements of ROS production and GSH concentration in the eggs. Unlike somatic cells, preimplantation embryos cannot synthesize GSH de novo [36] and may therefore be very sensitive to ROS, even at low concentrations [7].

ROS can be toxic when in excess but may also play a regulatory role in the control of mitochondria activity, particularly in mitochondrial biogenesis [37]. In cell lines carrying different common mouse mtDNA haplotypes a direct correlation between ROS generation and mtDNA content has been shown [38]. This is in agreement with our observation that oocytes from obese females producing more ROS had a higher mtDNA copy number than oocytes from lean mice. Expression of the nuclear genes involved in mitochondrial biogenesis, specifically PGC-1, NRF-1 and mtTFA are up-regulated in some human cell types [39] and in rat hepatocytes [40] in response to oxidative stress. Hence, the increased mtDNA copy number as well as NRF1 and TFAM mRNA expression in the oocyte of the obese female may be attributable to an oxidative stress-mediated increase in the transcription of genes involved in mitochondrial biogenesis. Surprisingly, increased mitochondrial biogenesis was present in the oocytes but not the zygotes of the obese females. Mitochondria replication and the synthesis of maternal nuclear encoded transcripts associated with mtDNA replication are ongoing processes in the growing oocyte [10] and therefore can be susceptible to ROS or nutritional and hormonal factors present in obese reproductive environment [12]. The normalisation of mtDNA and mRNAs associated with mtDNA replication after

Table 2. The influence of maternal diet-induced obesity on early embryo development.

| Parameter                          | Control females | Obese females | Significance |
|-----------------------------------|-----------------|---------------|-------------|
| Number of females with zygotes    | 100% (7)        | 100% (7)      | P>0.05      |
| Zygote recovered/mouse            | 7.9±0.1 (7)     | 8.9±1.7 (7)   | P>0.05      |
| Number of females with blastocysts/mated | 82% (9/11)  | 54% (6/11)    |             |
| Number of females without blastocysts/mated | 18% (2/11) | 46% (5/11)    | P<0.02 (11) |
| Blastocyst recovered/mouse        | 6.7±1.2 (7)     | 7.7±1.6 (6)   | P>0.05      |
| Blastocyst total cell number      | 45.8±2.1 (7)    | 44.8±3.2 (6)  | P>0.05      |
| ICM cell number                   | 14.3±1.2(7)     | 14.1±0.6 (6)  | P>0.05      |
| TE cell number                    | 31.7±1.6 (7)    | 30.6±0.8 (6)  | P>0.05      |
| ICM/TE cell number (%)            | 24.1±5.2 (7)    | 22.2±1.3 (6)  | P>0.05      |

Data expressed as mean ± SEM. The number of zygotes was determined in the morning after natural mating. Blastocysts were recovered from the uterus on day 4 after mating. The number of cells per blastocyst and their distribution between the inner cell mass and the trophectoderm were analysed in 3–5 blastocysts per mouse. Values in parentheses indicate a number of mice/group.

1In both control and in 2 out of 5 obese mothers a small number of fragmented embryos at various stages of development was recovered from the oviducts.

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fertilization may be explained by a combination of our recent findings [13] and established molecular characteristics of the early embryo [41]. We have previously reported a period of mtDNA turnover where both mtDNA synthesis and destruction occur for a short period after fertilization [15]. This period of mtDNA turnover may offer a mechanism by which abnormal levels of mtDNA could be normalised in zygote stage embryos from obese dams. Additionally, it is well established that all maternally inherited transcripts are degraded during the 1–2 cell stage prior to the onset of the main burst of zygotic transcription during the late two cell stage [41]. Thus the normalisation of nuclear encoded transcripts associated with mitochondrial biogenesis in zygotes from obese reflects this documented global maternal mRNA destruction.

In this study a significant impairment in the ability to support embryo development to the blastocyst stage has been seen in 46% of the obese mothers. Similar numbers of zygotes were readily recovered from both groups, however. Thus, the absence of blastocysts in 46% of obese females is not attributable to anovalation but more likely due to increased embryonic death since fragmented embryos were found in the oviducts of 2 out of 5 obese females that failed to produce blastocysts. In support of this idea, a recent study by Minge et al. [42] has reported poor oocyte quality, reduced blastocyst survival rates and abnormal embryonic cellular differentiation in obese female mice. In women obesity is also associated with poor pre-implantation embryo quality and a reduced rate of embryo survival [5].

We also found that the obeseogenic diet increased serum fatty acid concentration and caused a marked increase in the oviductal leptin concentration. Leptin is essential for normal pre-implantation development of mouse embryos [43]. However, exposure to a higher leptin concentration in vivo [44] and in vitro [45] impairs embryo development, reducing the rate of blastocyst formation. In obese women, hyperleptinaemia is associated with poor fertility and increased risk of early pregnancy loss [46]. Leptin is also crucial to mitochondrial function. In rodents, leptin increases the expression of enzymes contributing to fatty acid oxidation [47] and up-regulates mitochondrial biogenesis through expression of PPARγ coactivator-1α [48]. Hyperlipidaemia, also frequently associated with obesity, increases fatty acids flux into oocytes and embryos which may increase ROS generation or trigger embryo apoptosis through mitochondria-dependent pathways [49].

In conclusion, exposure of oocytes and embryos to an obese reproductive environment was associated with qualitative and quantitative changes in mitochondria, oxidised redox state, increased oxidative load and impaired antioxidant capacities. Oocytes and embryo with compromised mitochondrial activity may not be able to exert tight regulation of focal substrate supply and demand and, as a result, generate ROS at rates that become developmentally toxic after fertilization [7]. Embryo apoptosis or arrest may then ensue. Further investigation of mitochondrial functions in oocytes and embryos are required, particularly with regard to obesity-related alterations in mitochondrial gene and protein expression that inappropriately regulate mitochondrial energy metabolism.

We propose that altered oocyte and early embryo mitochondrial metabolism, resulting from excessive nutrients exposure prior to and during conception may be responsible for poor reproductive outcomes frequently reported in obese women.

Materials and Methods

Experimental animals and diets

This study was conducted in accordance with the UK Home Office Animal (Scientific Procedures) Act 1986. Six week-old virgin female C57BL/6J mice (Charles River Laboratories, UK) were fed either a standard chow diet (7% simple sugars, 3% fat, 50% polysaccharide, 15% protein [w/w]) RM1, Special Dietary Services, n = 10) or highly palatable obesogenic diet (10% simple sugars, 20% animal lard, 28% polysaccharide, 23% protein [w/w]), Special Dietary Services, n = 10) supplemented with sweetened condensed milk and micronutrient mineral mix (AIN93G, Special Dietary Services) [19]. After 6 weeks of diet, all females were induced to superovulate by consecutive ip injections of 10 IU pregnant mare’s serum (Dunlops) and 10 IU human chorionic gonadotrophin (hCG, Dunlop). Prior to embryo collection mice were celled by cervical dislocation. Blood samples were taken via cardiac puncture. Abdominal and inguinal fat pads and body weights were recorded. Twelve hours post hCG fully grown oocytes were collected by puncturing pre-ovulatory follicles with sterile needles and treated with hyaluronidase (0.5 mg/ml) to remove surrounding cumulus cells. Zygotes and blastocysts were collected after successful mating with males at 24 and 84 h after post-hCG as described [15].

Oviduct fluid was collected as described in [50] with minor modifications. Briefly, pregnant female mice (day 4) were celled and oviducts were ligatured at uterotubal junction with 6-0 black suture silk. Oviducts with attached ovaries were excised, rinsed in saline and placed under mineral oil on a watch glass. Ovaries were separated and the thread was removed at the cut end of the uterus under the microscope. A curved blunt-end metal capillary connected to a 1-ml syringe was inserted into infundibulum and oviducts were carefully flushed with 50 μl of flushing medium (PBS supplemented with 0.3% polysvinylpyrrolidione). Flushing from both oviducts was collected and placed into a sterilized 0.5-ml microcentrifuge tube and centrifuged at 10,000 g for 10 min to remove cellular debris, and the supernatant was aspirated into another sterilized 0.5-ml microcentrifuge tube and stored at −80°C until assayed for concentrations of leptin and glucose.

Assessment of mitochondrial morphology and metabolism

Mitochondria of living oocytes and zygotes were imaged using a Zeiss 510 uv-Vis CLSM META and a range of targeted fluorescent probes (Molecular Probes) as described previously [23,51]. Some measurements were made using a cooled CCD camera (Orca ER).

The distribution of active mitochondria and ΔΨm were analysed in eggs incubated with tetrathyl rhodamine methyl ester (TMRM; 25 nM) in M2 medium at 37°C for 30 min. The TMRM is a fluorescent lipophilic cation and accumulates into mitochondria in response to the negative mitochondrial membrane potential. TMRM was excited using the 543 nm laser line and fluorescence measured using a 560 long-pass filter.

Measurements of NADP/H and FAD2* autofluorescence intensity in oocytes and zygotes were used to estimate the mitochondrial redox potential. The reduced forms of pyridine nucleotides (NADP/H) are excited by ultraviolet light at 351 nm excitation line of the CLSM and measured between 435–485 nm. The oxidised form (NADP+) is non-fluorescent. In contrast to NADP/H, it is the reduced form of flavoproteins (FADH2) that is non-fluorescent. Fluorescence of the oxidised form of flavoproteins (FAD2*) was excited at 438 nm and emitted fluorescence was collected throughout the 505–550 nm bandpass filter.

The resting redox state was defined as a function of the maximally oxidised and maximally reduced signals which were obtained by adding FCCP (1 μM) to drive the signals to maximal oxidation followed by addition of 1 mM NaN3 which drives the signals to a maximally reduced state. The fluorescence signals are...
then normalised between 100 (maximal reduction to NAD(P)H and maximal oxidation to FADH2) and 0 (maximal oxidation to NAD(P)H and maximal reduction to FADH2), giving a value which is a measure of the resting relative redox state [23].

For measurement of cytosolic ROS production, HEt (2 μM) was added to M2 medium and remained present throughout the experiment (15 min). This is a non-fluorescent derivative of the red fluorescent ethidium, and so an increase in red fluorescence (excited at 543 nm and measured at >560 nm) gives a measure of the rate of oxidation of the dye and therefore of the rate of ROS generation.

In order to measure GSH, cells were incubated with 50 μM MCB in M2 medium at 37°C for 40 min, or until a steady state had been reached before images were acquired. Non-fluorescent MCB undergoes a reaction with glutathione catalysed by glutathione-s-transferase to yield a fluorescent adduct which therefore gives a measure of GSH content [52]. MCB fluorescence was excited at 351 nm and measured at 430–480 nm.

Fluorescent images were obtained from at least from 3 oocytes/zygotes from 5 females per diet. Image analysis, differentiation and exponential curve fitting were performed using Origin 8 software (OriginLab Corporation).

Quantification of mtDNA copy number
Total DNA was extracted from groups of two oocytes and zygotes (n = 8 females/group) as described [53]. A mtDNA content was measured by qPCR on Thermal Cycler Corbett Rotorgene TM 6000 (Corbett research) using QuantiFast SYBR Green PCR Kit (Qiagen) and primers corresponding to 16S ribosomal gene. Sample copy number was determined with mtDNA standard curves using Rotorgene 6000 software.

Gene expression analysis
Poly(A)+ RNA was isolated from snap-frozen oocytes (n = 20 oocytes/8 females/group) and zygotes (n = 10zygotes/8 females/group) using magnetic oligo(dT) beads (Dynabeads mRNA DIRECT Kit). cDNA synthesis was performed by random hexamer priming and the Transcriptor First Strand cDNA Synthesis Kit (Roche). qPCR took place on a Chromo4 thermocycler (MJ) using the Precision SybrGreen Master Mix (Primerdesign). Each assay was performed in duplicates using intron-spanning primers (Operon Biotechnologies GmbH). Stability of housekeeping genes was validated using geNorm application and H2afz mRNA was selected for normalization. Relative mRNA abundance was determined using the comparative deltatCt method. A list of genes investigated and Primer sequences are shown in Table 3.

Differential nuclear staining
The number of cells per blastocyst and their distribution between the inner cell mass and the trophectoderm were counted by differential fluorochrome nuclear labelling [54].

Metabolic Studies
Fasted glucose, fatty acids and triglycerides concentrations were assessed using autoanalyser (LX20, Beckman Coulter) and the assay kits (Glucose; UV-hexokinase; nr. GLU 1442640; triglycerides; enzymatic GPO method; nr. TG 445850; total cholesterol; enzymatic method; nr. CHOL 467825) as described in [19]. Concentrations of leptin were measured by a sandwich ELISA using paired leptin antibodies (Duoset, R&D Systems Ltd). Serum and oviduct fluid samples from the mice were diluted 1:20 for the assay and leptin concentration of samples was calculated from a standard curve constructed with mouse leptin standards.

Statistics
All results are expressed as mean ± SEM. Data were analysed by Student’s t-test after testing for normal distribution using Graphpad Prism v. 2.01 (Graphpad Software, USA). A value of p<0.05 was considered significant.

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Author Contributions
Conceived and designed the experiments: NI MRD JM. Performed the experiments: NI AYA. Analyzed the data: NI AYA JJE. Contributed reagents/materials/analysis tools: AYA LP JJE TPF MRD JM. Wrote the paper: NI.

References
1. Ramsay JE, Greer I, Sattar N (2006) ABC of obesity. Obesity and reproduction. Bmj 333: 1159–1162.
2. Baker P, Balen A, Poston L, Sattar N (2007) Obesity and Reproductive health: RCOG Press. 286 p.
3. Metwally M, Li TC, Ledger WL (2007) The impact of obesity on female reproductive function. Obes Rev 8: 515–523.
4. Maheshwari A, Stollberg L, Bhattacharya S (2007) Effect of overweight and obesity on assisted reproductive technology: a systematic review. Hum Reprod Update 13: 433–444.
5. Robker RL (2008) Evidence that obesity alters the quality of oocytes and embryos. Pathophysiology 15: 115–121.
6. Robker RL, Akinson LD, Bennett BD, Thrupp PN, Chura LR, et al. (2009) Obese women exhibit differences in ovarian metabolites, hormones, and gene expression compared with moderate-weight women. J Clin Endocrinol Metab 94: 1533–1540.
7. Leese HJ, Baumann CG, Brison DR, McEvoy TG, Sturmer RG (2008) Metabolism of the viable mammalian embryo: quietness revisited. Mèh Reprod 14: 667–672.
8. Fleming TP, Kwong WY, Potter R, Ursell E, Fesenko I, et al. (2004) The embryo and its future. Biol Reprod 71: 1046–1054.
9. Cummins JM (2004) The role of mitochondria in the establishment of oocyte functional competence. Eur J Obstet Gynecol Reprod Biol 115 Suppl 1: S23–29.
10. Dumollard R, Duchen M, Carroll J (2007) The role of mitochondrial function in the oocyte and embryo. Curr Top Dev Biol 77: 21–49.

11. Dumollard R, Carroll J, Duchen MR, Campbell K, Swann K (2009) Mitochondrial function and redox state in mammalian embryos. Semin Cell Dev Biol 20: 346–353.

12. Jansen RP, Burton GJ (2004) Mitochondrial dysfunction in reproduction. Mitochondrion 4: 577–600.

13. Van Blommen J (2008) Mitochondria as regulatory forces in oocytes, preimplantation embryos and stem cells. Reprod Biomed Online 16: 555–569.

14. Tachibana M, Sparmian, M, Sirnataunomchi H, Ma H, Clepper L, et al. (2009) Mitochondrial gene replacement in primates offspring and embryonic stem cells. Nature 461: 367–372.

15. McConnel M, Petrie L (2004) Mitochondrial DNA turnover occurs during preimplantation development and can be modulated by environmental factors. Reprod Biomed Online 9: 418–424.

16. Thomas GA, Trounson AO, Jones GM (2006) Developmental effects of sublethal mitochondrial injury in mouse oocytes. Biol Reprod 74: 969–977.

17. Wakefield SL, Lane M, Schulz SJ, Hebart ML, Thompson JG, et al. (2008) Maternal supply of omega-3 polyunsaturated fatty acids alter mechanisms involved in early embryonic development in the mouse. Am J Physiol Endocrinol Metab 294: E425–E434.

18. Mitchell M, Schulz SL, Armstrong DT, Lane M (2009) Metabolic and mitochondrial dysfunction in early mouse embryos following maternal dietary protein intervention. Biol Reprod 80: 622–630.

19. Samuelson AM, Matthews PA, Argenton M, Christie MR, McConnell JM, et al. (2008) Diet-induced obesity in female mice leads to offspring hyperphagia, adiposity, hypertension, and insulin resistance: a novel murine model of developmental programming. Hypertension 51: 303–392.

20. Minamikawa T, Williams DA, Bowser DN, Nagley P (1999) Mitochondrial permeability transition and swelling can occur reversibly without inducing cell death in intact human cells. Exp Cell Res 246: 26–37.

21. Keelan J, Allen NJ, Antcliffe D, Pal S, Duchen MR (2001) Quantitative imaging of mitochondrial function during preimplantation development and early embryonic development. Proc Natl Acad Sci U S A 98: 11848–11853.

22. Brannian JD, Schmidt SM, Kreger DO, Hansen KA (2001) Baseline non-fasting serum leptin concentration to body mass index ratio is predictive of IVF outcomes. Hum Reprod 16: 1819–1826.

23. Mayevsky A, Rogatsky GG (2007) Mitochondrial function in vivo evaluated by mitochondrial membrane potential, uncoupling and permeability transition. Biochim Biophys Acta 1757: 156–162.

24. Holmgren A, Johansson C, Berndt C, Lonn ME, Hudemann C, et al. (2005) Mitochondrial metabolism modulates differentiation and teratoma formation capacity in mouse embryonic stem cells. J Biol Chem 280: 20556–20512.

25. Holmgren A, Johansson C, Berndt C, Lonn ME, Hudemann C, et al. (2005) Thiol redox control via thioredoxin and glutaredoxin systems. Biochem Soc Trans 33: 1373–1377.

26. Gardner CS, Reed DJ (1995) Synthesis of glutathione in the preimplantation mouse embryo. Arch Biochem Biophys 318: 30–36.

27. Lee HC, Wei YH (2005) Mitochondrial biogenesis and mitochondrial DNA maintenance of mammalian cells under oxidative stress. Int J Cell Biol 37: 822–834.

28. Schieke SM, Ma M, Gao L, McCoy JP, Jr., Liu C, et al. (2008) Mitochondrial metabolism modulates differentiation and teratoma formation capacity in mouse embryonic stem cells. J Biol Chem 283: 28506–28512.

29. Van Blommen J (2008) Mitochondria as regulatory forces in oocytes, preimplantation embryos and stem cells. Reprod Biomed Online 16: 555–569.

30. Holmgren A, Johansson C, Berndt C, Lonn ME, Hudemann C, et al. (2005) Thiol redox control via thioredoxin and glutaredoxin systems. Biochem Soc Trans 33: 1373–1377.

31. McConnel M, Petrie L (2004) Mitochondrial DNA turnover occurs during preimplantation development and can be modulated by environmental factors. Reprod Biomed Online 9: 418–424.

32. Thomas GA, Trounson AO, Jones GM (2006) Developmental effects of sublethal mitochondrial injury in mouse oocytes. Biol Reprod 74: 969–977.

33. Wakefield SL, Lane M, Schulz SJ, Hebart ML, Thompson JG, et al. (2008) Maternal supply of omega-3 polyunsaturated fatty acids alter mechanisms involved in early embryonic development in the mouse. Am J Physiol Endocrinol Metab 294: E425–E434.

34. Mitchell M, Schulz SL, Armstrong DT, Lane M (2009) Metabolic and mitochondrial dysfunction in early mouse embryos following maternal dietary protein intervention. Biol Reprod 80: 622–630.

35. Samuelson AM, Matthews PA, Argenton M, Christie MR, McConnell JM, et al. (2008) Diet-induced obesity in female mice leads to offspring hyperphagia, adiposity, hypertension, and insulin resistance: a novel murine model of developmental programming. Hypertension 51: 303–392.

36. Minamikawa T, Williams DA, Bowser DN, Nagley P (1999) Mitochondrial permeability transition and swelling can occur reversibly without inducing cell death in intact human cells. Exp Cell Res 246: 26–37.

37. Keelan J, Allen NJ, Antcliffe D, Pal S, Duchen MR (2001) Quantitative imaging of mitochondrial function during preimplantation development and early embryonic development. Proc Natl Acad Sci U S A 98: 11848–11853.

38. Brannian JD, Schmidt SM, Kreger DO, Hansen KA (2001) Baseline non-fasting serum leptin concentration to body mass index ratio is predictive of IVF outcomes. Hum Reprod 16: 1819–1826.

39. Minokoshi Y, Kim YB, Peroni OD, Fryer LG, Muller C, et al. (2002) Leptin stimulates fatty-acid oxidation by activating AMP-activated protein kinase. Nature 415: 339–343.

40. Lee Y, Xu X, Gonzales F, Mangedorf DJ, Wang MY, et al. (2002) PPAR alpha is necessary for the lipolytic action of hyperleptinemia on white adipose and liver tissue. Proc Natl Acad Sci U S A 99: 11348–11353.

41. Di Paola M, Lorusso M (2006) Interaction of free fatty acids with mitochondria: coupling, uncoupling and permeability transition. Biochim Biophys Acta 1737: 1330–1337.

42. Harris CE, Gopichandran N, Picton HM, Leece HJ, Orii NM (2003) Nutrient concentrations in murine follicular fluid and the female reproductive tract. Theriogenology 62: 992–1006.

43. Abramov AV, Scrofani A, Duchen MR (2007) Three distinct mechanisms generate oxygen free radicals in neurons and contribute to cell death during anoxia and reoxygenation. J Neurosci 27: 1129–1130.

44. Kekal J, Allen NJ, Antcliffe D, Pal S, Duchen MR (2001) Quantitative imaging of glutathione in biphasocampal and glia in culture using monoclonal- ohbainan. J Neurosci Res 66: 873–894.

45. Shiata H, Kanceda H, Sato A, Inoue K, Ogura A, et al. (2000) Selective and continuous elimination of mitochondria microinjected into mouse eggs from spermatids, but not from liver cells, occurs throughout embryogenesis. Genetics 150: 1277–1294.

46. Handside AH, Hunter S (1984) A rapid procedure for visualising the inner cell mass and trophectoderm nuclei of mouse blastocysts in situ using polynucleotide- staining. J Reprod Fertil 73: 156–165.