EXPERIMENTAL STUDY

Does maternal dietary mineral restriction per se predispose the offspring to insulin resistance?

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Abstract

Background: Maternal undernutrition is hypothesized to predispose the offspring to disease in adult life. The relevance of maternal macronutrient deficiency has been well studied but not that of micronutrients.

Objective: To assess the effect of maternal dietary mineral restriction per se on oral glucose tolerance (OGT), insulin resistance (IR) and fat metabolism in offspring.

Design: Female weanling Wistar/NIN rats received a control or a 50% mineral-restricted (MR) diet for 12 weeks, by which time MR rats had lower plasma Fe, Zn, Mg and Ca concentrations. Following mating with control males, a third of the MR dams were shifted to the control diet from parturition. Half of the pups born to the remaining MR dams were weaned onto the control diet while the other half continued on the MR diet.

Results: Pregnant MR dams had a higher abortion rate, body weights of their pups at birth and weaning were lower and rehabilitation had no beneficial effect. No offspring had impaired OGT, and IR status was comparable among different groups on postnatal days 40, 70, 100 or 180. Compared with controls, total body electrical conductivity measurements indicated significantly higher body fat %, lower lean body mass and fat-free mass in MR offspring besides elevated plasma triacylglycerols. Mineral rehabilitation from parturition or weaning had little effect on these changes, which did not appear to be due to increased oxidative stress.

Conclusions: Maternal MR per se resulted in an increase in body fat and in plasma triacylglycerol concentrations in the offspring. These changes had, however, no discernable effect on insulin sensitivity over the first 180 days of life.

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Introduction

Exposure of the fetus to maternal malnutrition is a well-known causal factor for intrauterine growth retardation, both in humans and other animals (1, 2). Abundant epidemiological evidence in humans has shown an association between coronary heart disease, hypertension, non-insulin-dependent diabetes and impaired glucose tolerance in adults on the one hand, and low birth weight and altered fetal growth on the other (3–5). Elevated blood pressure, glucose intolerance and dyslipidaemia occur together frequently in humans and are collectively termed as the ‘metabolic syndrome’ (6, 7). The fetal origin of adult disease hypothesis proposes that these disorders derive from fetal adaptations in utero to maternal undernutrition, which permanently alters growth characteristics and postnatal metabolism and physiology (8, 9). Indeed, a number of animal models have been established to try to understand the mechanistic basis of this relationship (10, 11), but most of them considered only the maternal deficiency of macronutrients. For example, maternal dietary restrictions of calories (12) and protein (13), and umbilical artery ligation (14) have all been shown to cause fetal growth retardation, alter fat and glucose metabolism and elevate blood pressure in the offspring. The data from these studies suggest that poor maternal nutrition during pregnancy may lead to metabolic and morphological changes in the fetus, with lifelong consequences.

It is well recognized that micronutrients, especially the minerals, play an important role in the structural and metabolic activities of the organism. For example, zinc, magnesium, manganese, iron and copper are known to modulate the synthesis, storage, release and conformational integrity of insulin (15). Chromium increases insulin binding due to an increase in insulin receptor number, activation of insulin receptor kinases and inhibition of insulin receptor phosphatases (16). Vanadium shows insulin mimetic effects in vivo and in vitro when used in pharmacologic doses (17). Recently, dietary iron restriction in rat dams has been shown to
increase hypertension and alter lipid metabolism in the offspring (18).

Moreover, mineral deficiencies and anemia are common in the developing world, particularly during pregnancy and lactation (19). It is also known that maternal mineral deficiencies are associated with low birth weight and increased rates of perinatal mortality and morbidity (20). Further, in developing countries, the prevalence of low birth weight varies from 13% to 30% (21). Despite this and the known effects of minerals per se on insulin synthesis/release/action, the role of maternal mineral deficiencies in the etiology of insulin resistance in the offspring has not yet been explored.

In view of previous studies, we hypothesized that maternal dietary mineral restriction per se predisposes the offspring to insulin resistance in later life. The present study was conducted in albino rats to validate/negate this hypothesis.

Materials and methods

Animals: feeding, maintenance and breeding

The animal experimental protocol and all the procedures involved were approved by the National Institute of Nutrition ethical committee on animal experiments.

Twenty, female, weaning Wistar/NIN (WNIN) rats were obtained from the National Centre for Laboratory Animal Sciences, National Institute of Nutrition, Hyderabad, India. They were housed individually in polypropylene cages with wire mesh bottom and maintained at 22±2°C under standard lighting conditions (12 h light: 12 h darkness). Of these, 14 were fed for 12 weeks on a control diet (based on the American Institute of Nutrition AIN-93G diet) (22) with 50% restriction of mineral mixture and they were provided with deionized water. The remaining six animals were pairfed with six rats fed the mineral-restricted (MR) diet. At the end of 12 weeks of feeding, blood was collected from the supra-orbital sinus for determining the concentrations of iron, zinc, copper, magnesium and calcium in plasma, in addition to the following biochemical parameters: haemoglobin, glucose, insulin, cholesterol and triacylglycerols.

After assessing their mineral status, the rats were mated with control males and maintained on their respective diets throughout gestation. At parturition, one-third of the MR dams (n = 4) were shifted to control diets (MSP) and the remaining two-thirds of the dams continued on restricted diets throughout lactation. A uniform litter size was maintained in all groups by adjusting the number of offspring per litter to eight on postnatal day 3. At 21 days of age (weaning), half of the pups born to MR mothers were weaned onto the control diet (MSW) and the remaining half continued on restricted diets (MR). From weaning onwards, in each group eight male pups were maintained until postnatal day 180. The feeding protocol used in this experiment is depicted schematically in Fig. 1.

Diet intake and body weights were recorded daily and weekly respectively in both mothers and offspring. At the end of 100 and 180 days of feeding of the offspring, body length was measured as the length between the tip of nose and the centre of the anus. Body mass index (BMI) was calculated from the formula: BMI = body weight in kg/body length in m².

Oral glucose-tolerance test

An oral glucose-tolerance test (OGTT) was performed in the offspring on postnatal days 40, 70, 100 and 180. After an overnight fast, glucose (300 gm/l) was administered orogastrically at a dose of 2.5 g/kg body weight and blood samples were obtained from the orbital sinus at 0, 60 and 120 min for determination of plasma glucose and insulin concentrations. The glucose and insulin responses during the OGTT were calculated by estimation of the total area under curves (AUC) for glucose and insulin respectively, using the trapezoidal method (23).

Physiological indices of insulin resistance/glucose tolerance

Indices based on fasting glucose and insulin concentrations (homeostasis model of assessment of insulin resistance (HOMA-IR)) as well as those which take into account the response to a challenge with oral glucose (glucose AUC/insulin AUC) were computed as follows. (i) HOMA-IR: insulin resistance was assessed from fasting glucose and fasting insulin concentrations as follows: HOMA IR = (fasting insulin (μU/ml) × fasting glucose (mM))/22.5 and (ii) ratio of glucose AUC to insulin AUC: AUC for glucose and insulin during the OGTT
were calculated by the trapezoidal method (23) and the ratio of glucose AUC/insulin AUC was computed.

**Measurements in plasma**

Blood was collected from offspring at 40, 70, 100 and 180 days of age following an overnight fast; plasma was separated and stored at –20°C until analysis. Iron, zinc, copper, magnesium and calcium concentrations were measured in maternal plasma by atomic absorption spectroscopy (Varian spectrAA 220) (24).

Plasma glucose and high density lipoprotein (HDL) cholesterol levels were measured enzymatically using commercially available kits (Dr Reddy Laboratories, Hyderabad, India) according to the manufacturer’s instructions. Total cholesterol and triacylglycerols were measured in fasting plasma using commercially available kits from Biosystems, Barcelona, Spain. Plasma insulin was measured using a radioimmunoassay kit from BRIT, Mumbai, India.

**Body composition of the offspring**

Total body composition of the offspring was determined at 100 and 180 days of age using a total body electrical conductivity (TOBEC) small animal body composition analysis system (EM-SCAN, Model SA-3000 multi detector, Springfield, IL, USA) (25, 26). The difference between the impedance measured when the animal is inside the electromagnetic field and when the chamber is empty is an index of the total electrical conductivity (E) of the body which, in turn, is proportional to the animal’s lean body mass (LBM).

Prior to TOBEC measurement, the rats were anesthetized lightly with ether and placed in a carrier in such a way that the animal’s body was stretched to its maximum comfortable length. Then the carrier was placed in the TOBEC chamber and 10–12 recordings/scans were taken from each rat. The highest and lowest readings were excluded and the remaining ten were averaged. The intra-assay coefficient of variation was less than 3.0%. Prior to the in vivo measurement, the instrument was calibrated with a standard coil (ID 3076) supplied with the instrument and the empty chamber read 0–2 units.

The following body composition parameters were obtained mathematically according to the methods of Morbach et al. (27). (i) LBM = 0.5 E + (0.3 × total body weight); (ii) total body fat = total body weight – LBM; total body fat % = (total body fat × 100)/total body weight; (iii) fat-free mass (FFM) = 16.28 + 0.4 E.

**Oxidative stress and antioxidant status**

The animals were killed on postnatal day 180 and livers were dissected out immediately, washed thoroughly with ice-cold saline, frozen in liquid nitrogen and stored at –80°C until analysis.

A part of the liver was weighed, minced and homogenized (10% w/v) in 50mM phosphate buffer (pH 7.0). The homogenate was centrifuged at 1000 g for 20 min at 4°C. A part of the supernatant was used for estimation of lipid peroxidation and protein carbonyls. The remaining supernatant was further centrifuged at 12 000 g for 20 min at 4°C to obtain the post-mitochondrial supernatant, which was used for the estimation of reduced glutathione (GSH) and activities of the antioxidant enzymes: catalase, superoxide dismutase (SOD) and glutathione peroxidase (GPx).

Lipid peroxidation was measured by the thiobarbituric acid color reaction for malondialdehyde (MDA) (28). Protein carbonyl content was measured spectrophotometrically using 2,4-dinitrophenyl-hydrazine (29). GSH was determined by its reaction with o-phenthaldehyde (30). Total SOD activity was assayed by monitoring the rate of inhibition of pyrogallol reduction (31). One unit of SOD represents the amount of enzyme required for 50% inhibition of pyrogallol reduction/min. Catalase activity was assayed by monitoring the disappearance of H2O2 at 240 nm (32). One unit of catalase represents the decrease of 1 μmol H2O2/min. The activity of total GPx was assayed using cumene hydroperoxide and GSH as substrates (33). One unit of GPx was defined as μmol NADPH oxidized/min. The protein content was determined using the modified Lowry method (34).

**Statistical analysis**

For maternal and neonatal data, differences between the control and restricted groups were analyzed using Student’s t-test. Data collected from the offspring after weaning were analyzed using one-way ANOVA followed by the multiple range test/least significance difference (LSD) method. Wherever heterogeneity of variance was observed, differences between groups were tested by non-parametric Mann–Whitney U tests. Comparisons considered were: control (MC) vs MR, MR vs MSP and MR vs MSW. All values are presented as means±S.E. The differences were considered significant if P was at least <0.05.

**Results**

**Parameters in mothers**

Maternal growth, mineral status and glucose homeostasis At the end of 3 months of feeding there was no significant difference in the body weight gain between the control and the MR rats. However, hemoglobin concentrations were significantly lower (P < 0.001) in MR animals than in controls. In addition, food intake was not significantly different
Reproductive performance of dams

Conception was 100% in both control and the MR groups. Weight gain during pregnancy was not significantly different between control and MR dams. Although the abortion rate was 14% (2/14) in MR animals (compared with none in controls), there were no significant differences between the groups in litter size (5–11), percent of still births (3–4%) and percent of deaths during lactation (4–5%). However, the mean birth weight of pups was significantly lower in the MR group than in the controls (5.52±0.11 vs 6.08±0.14 g; P < 0.01).

Parameters in offspring

Growth characteristics of the offspring

The offspring of the MR dams which had a lower birth weight continued to weigh less until postnatal day 180, whether they were kept on an MR diet or shifted to the control diet from parturition or weaning (Fig. 2). Although there were no differences in BMI among the different groups measured at postnatal day 100, the offspring of MR, MSP and MSW dams had a lower BMI compared with controls at postnatal day 180 (Fig. 2). Although there were no differences in their diet intake, the haemoglobin concentrations of MR pups continued to be lower than those of controls until postnatal day 180 and rehabilitation either from parturition or weaning completely corrected these changes (data not shown).

Glucose tolerance and insulin resistance

There were no significant differences in the fasting glucose concentrations between MC and MR, or MSP and MSW vs MR on days 40 and 70 of postnatal life. However, on postnatal days 100 and 180, the offspring of the MSP and MSW groups had significantly lower fasting glucose concentrations (P < 0.05) compared with controls (Fig. 3A). There was, however, no significant differences in fasting insulin concentrations among the four groups at any of the time-points tested (Fig. 3B). The AUC for glucose and insulin during an OGTT were, in general, not significantly different among the four groups at any time-point tested (Fig. 4A and B), except for some transient changes seen in the MSP and MSW groups on postnatal days 40 and 180 in glucose AUC values. It was indeed interesting that none of the animals in any of the groups had impaired OGT at any of the time-points tested.

Nevertheless, insulin resistance status was assessed among the offspring of the control and the other groups by computing indices, such as HOMA, based on fasting glucose and insulin or the ratio of the glucose AUC/insulin AUC which take into account values during OGTT. In general, MR pups were not significantly different from controls in both the indices tested at any of the four time-points tested (Figs 3C and 4C), nor did rehabilitation from parturition (MSP) or weaning (MSW) have any impact on these indices.

Body composition of the pups

Interestingly, maternal MR had a significant effect on the body composition of
offspring measured on postnatal days 100 and 180. Indeed, body fat % was significantly higher ($P < 0.01$) in MR offspring compared with controls (Fig. 5A). Other markers of adipogenesis such as LBM and FFM (Fig. 5B and C) were significantly lower ($P < 0.001$) in the MR offspring compared with controls. However, rehabilitation of MR dams from parturition (MSP) or weaning MR pups onto the control diet (MSW) seemed to have no significant effect on any of these parameters at both the time-points tested.

**Plasma lipids** In keeping with high body fat %, plasma triacylglycerol concentrations were significantly higher in the MR offspring than controls (Fig. 6) ($P < 0.05$) at all the four time-points tested. Interestingly, mineral rehabilitation from parturition (MSP) or weaning (MSW) appeared to correct the changes almost completely. Plasma total cholesterol and HDL cholesterol were, in general, not significantly different among the groups at any of the time-points tested (data not shown).

**Oxidative stress and antioxidant status** Chronic maternal MR resulted in a significant ($P < 0.05$) decrease in reduced GSH concentrations in the MR offspring and an increase in the concentrations of protein carbonyls (Table 2), albeit not statistically significant. Interestingly, rehabilitation from parturition or weaning could not prevent either the decrease in GSH or the increase in the concentrations of protein carbonyls. Lipid peroxides (MDA) and activities of the antioxidant enzymes catalase, SOD and GPx were similar among all the groups (Table 2).

**Discussion**

The present study was designed to assess the hypothesis that 50% restriction of maternal dietary minerals *per se* predisposes the offspring to insulin resistance in later life. We have assessed the effect of maternal MR during the phases of growth, pregnancy and lactation, on glucose homeostasis, insulin resistance and fat metabolism/adipogenesis in the offspring at 40, 70, 100 and 180 days of age. The results revealed that although 50% maternal MR *per se* did not affect glucose homeostasis and insulin resistance in the offspring until postnatal day 180, it significantly increased their body fat content and concentrations of plasma triacylglycerols compared with controls. Further, these changes could be due to increased oxidative stress and/or decreased antioxidant defence in these offspring. It was interesting that neither rehabilitation of the MR

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**Figure 3** Fasting (A) glucose, (B) insulin and (C) HOMA-IR of the rat offspring of different groups on different postnatal days. Values are means±S.E. ($n=6$). aMC vs MR, bMSP vs MR and cMSW vs MR, significant at $P < 0.05$ by ANOVA/multiple range test/LSD.

**Figure 4** AUC of (A) glucose, (B) insulin and (C) the ratio of glucose AUC to insulin AUC during the OGTT in the offspring on different postnatal days. Values are means±S.E. ($n=6$). aMC vs MR, bMSP vs MR and cMSW vs MR, significant at $P < 0.05$ by ANOVA/multiple range test/LSD.
mothers from parturition nor weaning the MR pups onto the control diet had any beneficial effect in mitigating these changes.

There are a number of similarities between our results and the long-term effects reported in the offspring of mothers fed low protein diets (35–37). Both the models induced early growth retardation, and altered body composition and lipid profile. Shifting them to control diets from parturition or weaning appeared to have no significant effect on body weights.

Maternal MR in general had no significant effect on fasting glucose and insulin values until postnatal day 70. However, on postnatal days 100 and 180, MR, MSP and MSW offspring had significantly lower values for fasting glucose but not insulin, compared with controls. Although the HOMA-IR values computed based on fasting glucose and insulin were not statistically different among the groups, MR, MSP and MSW offspring had lower values compared with controls.

In keeping with the observations made above, there were no differences among the offspring of different groups in their OGT until postnatal day 70 and this was true even on postnatal day 100. However, on postnatal day 180, the MSP offspring had the highest glucose AUC while the MC, MR and MSW pups were comparable. MR, MSP and MSW offspring, having a lower insulin AUC at this time-point compared with controls (of which only MSP and MSW were significant), probably indicates that maternal MR decreased the capacity of the offspring to respond to a challenge of oral glucose, and rehabilitation from parturition or weaning may not mitigate the defect. These observations are comparable with the previous reports on maternal protein and calorie malnutrition (37, 38). As a consequence of this, the ratio of glucose AUC to that of insulin was the highest in MSP offspring while that of MR and MSW was higher than that of MC, but not significant. These results based on OGT not only corroborate those from fasting concentrations of glucose and insulin, but also appeared to suggest that rehabilitation of MR mothers from parturition may worsen the insulin response of the offspring to a glucose challenge.

Despite the changes observed above, it was interesting to note that none of the offspring in any of the groups had impaired glucose tolerance until postnatal

![Figure 5](A) Body fat %, (B) LBM and (C) FFM of the offspring on postnatal days 100 and 180 as determined by TOBEC. Values are means ± S.E. (n = 6). aMC vs MR and aMSW vs MR, significant at P < 0.05 by ANOVA/multiple range test/LSD.

![Figure 6](Plasma triacylglycerols in the offspring of different groups on postnatal days 40, 70, 100 and 180. Values are means ± S.E. (n = 6). aMC vs MR, aMSP vs MR and aMSW vs MR, significant at P < 0.05 by ANOVA/multiple range test/LSD.

### Table 2

<table>
<thead>
<tr>
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<th>MDA (nmol/mg protein)</th>
<th>Protein carbonyls (nmol/mg protein)</th>
<th>Reduced GSH (μmol/mg protein)</th>
<th>Catalase (units/mg protein)</th>
<th>SOD (units/mg protein)</th>
<th>GPx (units/mg protein)</th>
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</thead>
<tbody>
<tr>
<td>MC</td>
<td>0.097 ± 0.006</td>
<td>4.95 ± 0.72</td>
<td>2.61 ± 0.07</td>
<td>4.89 ± 0.58</td>
<td>15.26 ± 0.70</td>
<td>0.119 ± 0.025</td>
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<tr>
<td>MR</td>
<td>0.088 ± 0.007</td>
<td>6.19 ± 1.04</td>
<td>1.73 ± 0.05</td>
<td>4.45 ± 0.61</td>
<td>16.28 ± 1.11</td>
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<tr>
<td>MSP</td>
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<td>2.49 ± 0.41</td>
<td>5.54 ± 0.28</td>
<td>13.64 ± 1.34</td>
<td>0.074 ± 0.024</td>
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<tr>
<td>MSW</td>
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<td>6.07 ± 1.16</td>
<td>2.31 ± 0.31</td>
<td>4.60 ± 0.51</td>
<td>14.22 ± 1.32</td>
<td>0.093 ± 0.037</td>
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</table>

*MC vs MR, significant at P < 0.05 by ANOVA/multiple range test.*
day 180. This may indicate the need for a greater duration of feeding for impaired glucose tolerance or insulin resistance to be manifest in the offspring. In this respect, our results are in partial agreement with those of Langley et al. (37) who showed that maternal protein restriction did not affect the glucose tolerance in the offspring at 3 months of age, but impaired it only at 15 months of age, at which time they were indeed diabetic. Our results are also in line with those of Lewis et al. (18) who demonstrated, in the offspring of iron-restricted rat dams, increased hypertension and altered serum lipids at 3 months of age but not impaired glucose tolerance.

In the present study, although maternal MR had no effect on glucose metabolism in the offspring, significant increases were observed in plasma triacylglycerols and body fat %. Indeed, body fat % as computed by the TOBEC measurements is well validated and is an indicator of visible fat. The finding in our study that both LBM (which includes tissue-associated fat) and FFM showed a significant decrease in MR compared to MC appears to suggest that tissue-associated fat may be decreased in MR rats, whereas visible fat percent increased. These changes in body fat % are in agreement with those found by Lucas et al. (35) who observed altered lipid metabolism in the offspring of rat dams subjected to protein malnutrition during pregnancy.

The high body fat % along with low body weight in the MR offspring observed here appear to be similar to those reported in the ‘thin fat babies’ seen in developing countries such as India, an abnormal condition attributed to maternal malnutrition (39). Though it is not immediately clear how maternal mineral restriction altered lipid metabolism, it appears that it may not be due to the increased oxidative stress in these animals. To the best of our knowledge, this is the first report showing that chronic maternal MR per se during the phases of growth, pregnancy and lactation affected lipid metabolism and antioxidant status in the offspring.

There is abundant material in the literature that indicates that altered adipogenesis/lipid metabolism is the earliest change seen much before tissue insulin resistance manifests (40–43). Indeed, insulin resistance is hypothesized to originate in impaired adipogenesis/lipid metabolism (42–44). However, in our study, MR offspring had neither impaired glucose tolerance nor insulin resistance until postnatal day 180, although they had high body fat content and altered lipid metabolism. Further, it was somewhat perplexing that the rehabilitation of MR dams from parturition or weaning MR pups onto the control diet almost completely reversed the plasma triacylglycerol concentrations, but had little effect on the body fat %, LBM and FFM, indicating only partial reversibility of the changes. Thus, maternal MR per se had no discernible effect on glucose tolerance or insulin resistance in the offspring, although it altered the body fat composition and lipid metabolism.

Nevertheless, in view of the earlier reports in the literature that changes in adipogenesis/fat metabolism are the earliest events in the manifestation of insulin resistance, our observations are not necessarily in conflict with the hypothesis that maternal MR per se may predispose the offspring to insulin resistance in later life.

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References


