Nutrition in pregnant or lactating rats programs lipid metabolism in the offspring

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Epidemiological studies in humans show that size in early life is related to blood cholesterol concentrations in adult life, raising the hypothesis that early nutrition programs later lipid metabolism, affecting risk for later vascular disease. Here, we tested the hypothesis that nutrition during pregnancy or lactation in the rat programs lipid metabolism in the offspring, studied in adult life (mean 6 months). Rats \( (n = 35) \) from normally-fed dams (controls) were compared with (1) rats \( (n = 22) \) from dams protein-restricted in pregnancy and lactation; (2) rats \( (n = 9) \) born to normally-fed mothers crossed to protein-restricted lactating dams and (3) those \( (n = 9) \) born of protein-restricted dams and crossed to normally-fed lactating animals. In these latter three groups the offspring showed long-term reduction in plasma cholesterol, HDL-cholesterol and triacylglycerol concentrations compared with controls. The effects were predominantly in males. These findings suggest that in the rat the sensitive period for nutritional programming of cholesterol and triacylglycerol metabolism is both pre- and postnatal (pre-weaning) and that rats may be ‘indirectly’ programmed by altering the maternal nutritional milieu during gestation or lactation. Whilst it has been hypothesized that early human undernutrition programs risk for vascular disease, one aspect of undernutrition, low maternal protein intake, in this rat model programmed lower plasma cholesterol and triacylglycerol concentrations.

Maternal diet: Lipid metabolism: Nutritional programming

Whether early nutrition has a long-term ‘programming’ effect (Lucas, 1994) on risk factors for adult degenerative vascular disease is currently an area of major public health concern (Fall et al. 1992). Epidemiological studies in humans show that size in early life is related to adult plasma levels of cholesterol, LDL-cholesterol and apoprotein B, recognized risk factors for cardiovascular disease. Thus weight at 1 year is inversely related to apoprotein B (Barker et al. 1993a); and subjects who had a small abdominal circumference have the highest levels of total and LDL-cholesterol in adult life (Barker et al. 1993c). In addition, men in the 60’s who were breast fed and not weaned by 1 year and those who were bottle fed had higher plasma total and LDL-cholesterol concentrations; the highest values being seen in those heavy at birth and light at 1 year (Barker et al. 1993a). These retrospective epidemiological findings generate the hypothesis that early nutrition has a role in programming long-term lipid metabolism in humans (Hales & Barker, 1992).

Currently, this hypothesis is being tested prospectively in humans (A. Lucas, R. Morley and M. Fewtrell, unpublished results). However, studies in animal models are important in defining effective nutritional interventions, sensitive periods and fundamental mechanisms for such programming events. In the early 1970s, Hahn (1984) reduced litter size in rats to induce experimental overfeeding during the sucking period: rats from small litters (four
pups) had, in adult life, a permanent elevation in total plasma cholesterol concentration compared with values found in adult rats from normal litters (fourteen pups). Mott et al. (1991) assigned baboons to breast or formula feeding and at the end of infancy fed both groups on a ‘Western’-style, high-saturated-fatty-acid diet. Compared with those fed on formula, the previously breast-fed group, in adult life, had higher serum levels of LDL- and VLDL-cholesterol, lower values of HDL-cholesterol, greater absorption of cholesterol from the gut, lower cholesterol excretion and, at post mortem, significantly more atherosclerosis (Lewis et al. 1988). These two examples in animal models provide compelling experimental evidence that adult cholesterol metabolism may be programmed by early nutrition.

In the animal studies cited the early dietary manipulation was applied to the infant. There has been speculation in humans however, (Hales & Barker, 1992) that nutrition of the mother could program long-term health outcomes in her progeny. In the current study we tested the hypothesis in an established rat model (Snoeck, 1990; Dahri et al. 1993; Desai et al. 1995) that maternal nutrition (low protein intake), either in pregnancy or lactation, could have a permanent programming effect on circulating lipids in the offspring.

METHODS

The methods used have been described in detail by Desai et al. (1996). Briefly, virgin female Wistar rats, weighing 240–260 g, were mated with normal males. Day 0 of gestation was taken as the day on which vaginal plugs were expelled. The rats were either fed on a diet containing 200 g protein/kg (‘regular’ diet) or an isoenergetic diet containing 80 g protein/kg (‘low-protein’ diet) throughout pregnancy and lactation. The diet was purchased from Hoppe Farms BV, Hoge Rijndijk 14, 3440 HD Woerden, The Netherlands. Details of the diets are available from the manufacturers and summarized elsewhere (Desai et al. 1996). These diets are identical to those used by Snoeck et al. (1990). Casein was used as the protein source. The 80 g protein/kg diet was made isoenergetic using cerelose (glucose) and its fat content was similar to the 200 g protein/kg diet.

Spontaneous delivery took place on day 22 of gestation. Feeding an 80 g protein/kg diet to pregnant rats did not have any adverse effects on the ability of the dams to carry litters, nor was perinatal mortality changed (Desai et al. 1996). Pregnant rats fed on an 80 g protein/kg diet showed a decreased gain in total body weight with no significant differences in total feed intake and litter size when compared with the pregnant rats fed on a 200 g protein/kg diet (Desai et al. 1996).

After birth, at 3 d of age, large litters were reduced to eight pups, ensuring a standard litter size per dam. Thus each mother nursed a total of eight pups, four males and four females. At 21 d of age two males and two females were studied for other variables and the remaining offspring were followed into adult life. A large random subsample of these latter animals was included in the current study on the early dietary effects on blood lipids. Four groups were followed into adult life. The two main groups were as follows: group 1 (controls) comprised thirty-five pups from ten dams that received the regular, 200 g protein/kg chow during both pregnancy and lactation; group 2 (low protein group) comprised twenty-two pups from seven dams that received the low-protein (80 g/kg) chow during pregnancy and lactation. Two smaller crossover groups were designed as follows: group 3 (postnatal low protein group) comprised nine pups born of three pregnant dams fed on a regular 200 g protein/kg diet that were the suckled by the dams fed on the low-protein diet throughout gestation and lactation, and group 4 (prenatal low protein group) comprised nine pups born to three dams on a low protein (80 g/kg) diet and then suckled by the dams fed on the regular 200 g protein/kg diet throughout. From 21 d the offspring
were weaned onto a normal laboratory chow (LAD 1: Rat and Mouse Breeder Diet, Special Diet Services, Witham, Essex). They continued on this diet into adulthood.

The present paper focuses solely on the impact of early nutrition on long-term lipid metabolism. However, the nutritional interventions affected other outcomes including early and later growth as reported by Desai et al. (1996). Briefly, since handling pups immediately after birth was found, in a pilot study, to cause loss of pups (resulting from mothers eating their pups or refusal to nurse them), the first body weights were recorded at 3 d of age to avoid undue stress to the mothers and pups. These weights were significantly different between groups 1 and 2 (9.3 (SD 1.4) and 6.2 (SD 0.8) g respectively $P < 0.001$). Animals from groups 2 and 3 continued to have lower body weights than group 1 (controls) even as adults despite weaning them onto a normal laboratory chow (LAD1 containing 200 g protein/kg). Their feed intake was also lower than that of group 1.

All animal work was undertaken with appropriate approval and in accordance with Home Office regulations. For analytical purposes, groups 2–4 were each compared separately with the controls, and also combined to produce a 'pre- or postnatal low protein group' ($n$ 40) for comparison with controls ($n$ 35). Statistical analyses included ANOVA, Student's $t$ tests and regression analysis.

**Blood samples and assays**

Blood samples from the offspring were taken from the tail, by amputation of the tip, at a mean of 6 months of age (184 (SD 50) d). Samples were separated, the plasma stored at $-85^\circ$ and subsequently analysed for total cholesterol, HDL-cholesterol and triacylglycerols. The assays were done on a Roche 'Cobas Bio' centrifugal analyser using Roche kits (Roche Diagnostics, Welwyn Garden City, Herts.). Instrument precision tests were made before each batch of analyses. The instrument coefficients of variation were 0.3–0.5%.

Triacylglycerols were analysed by an enzymic colorimetric test with glycerol phosphate oxidase (EC 1.1.3.21) and 4-aminophenazone. Plasma triacylglycerols were converted to glycerol by lipoprotein lipase (EC 3.1.1.34). The glycerol was converted to glycerol-3-phosphate by glycerol kinase (EC 2.7.1.30) and then to dihydroxyacetone phosphate and $\text{H}_2\text{O}_2$ which, in the presence of peroxidase (EC 1.11.1.7), effects the oxidative coupling of 4-chlorophenol and 4-aminophenazone to form a red-coloured quinoneimine derivative. The colour intensity, determined at a wavelength of 520 nm, is proportional to triacylglycerol concentration.

Total cholesterol was analysed by an enzymic colorimetric test. Cholesterol ester was converted to cholesterol by cholesterol esterase (EC 3.1.1.13) and then to 4-cholesten-3-one and $\text{H}_2\text{O}_2$ by cholesterol oxidase (EC 1.1.3.6). Liberated $\text{H}_2\text{O}_2$ in the presence of peroxidase effects the oxidative coupling of phenol and 4-aminantipyrine to form a red-coloured quinoneimine derivative. The colour intensity, determined at a wavelength of 500 nm, is directly related to cholesterol concentration.

LDL and VLDL were precipitated using a phosphotungstate–MgCl$_2$ reagent. After centrifugation at 4000 rev./min for 10 min, the HDL remains in the supernatant fraction. HDL-cholesterol was then assayed enzymically as described for cholesterol.

Quality control sera ('Precinorm' supplied by Boehringer Mannheim UK, Diagnostics and Biochemistry Ltd, Lewes, E. Sussex) were included with each batch of samples. Inter-assay coefficients of variation were all below 2%.
RESULTS

Main comparison

The two main groups (group 1: controls; group 2: maternal low protein intake during pregnancy and lactation) were sampled at corresponding ages in adult life. Group means for plasma lipids are compared in Fig. 1; there were significantly lower values in rats from mothers given low-protein v. control diets for plasma cholesterol; control v. low protein (mmol/l): 2.20 (SE 0.19) v. 1.69 (SE 0.10), 95% CI 0.09–0.93 (P = 0.019); for HDL-cholesterol: 1.88 (SE 0.19) v. 1.43 (SE 0.32), 95% CI 0.05–0.85 (P = 0.029); and for plasma triacylglycerol: 2.95 (SE 0.34) v. 1.70 (SE 0.17), 95% CI 0.48–2.01 (P = 0.002).

Analyses including crossover groups

Fig. 1 also shows group means for the two further crossover groups: group 3, postnatal low protein group (rats who were fed by mothers who received the low-protein diet in lactation) and group 4, the prenatal low protein group (rats born to dams given low protein in pregnancy). ANOVA for each lipid type showed significant differences between the four comparison groups, reflecting the higher values in the controls. Indeed, there were no significant differences between groups 2, 3 and 4, suggesting that giving dams a low-protein diet postnatally (group 3), prenatally (group 4) or both (group 2) had the same impact on plasma cholesterol, HDL-cholesterol and triacylglycerol concentrations in the offspring. Despite the small sample size, compared with controls (using Student’s t test) the postnatal low protein group (group 3) had significantly lower plasma cholesterol (P = 0.04), HDL-cholesterol (P = 0.05) and triacylglycerol (P = 0.004) concentrations. Results for group 4 (prenatal low protein) had a larger standard deviation, and a significant reduction compared with controls was only found for triacylglycerol (P = 0.03).

Combined groups

Since lipid values for groups 2, 3 and 4 did not differ, they were combined and compared with those in group 1 (controls) (Fig. 1). The reductions in plasma cholesterol, HDL-cholesterol and triacylglycerol concentrations in this combined group were again significant compared with controls (P = 0.02, P = 0.03 and P = 0.0006 respectively). All these differences between groups and combined groups observed were also found after adjusting for minor chance differences between groups in age at blood sampling.

Influence of sex

Table 1 shows that males had significantly higher plasma cholesterol and triacylglycerol concentrations than females (P = 0.04 and P = 0.001 respectively) and a trend to higher values of HDL-cholesterol (P = 0.09).

Analyses were performed to explore whether the maternal dietary effect on the offspring was influenced by their sex. For these analyses group 1 (control group) was compared with three reduced maternal protein intake groups (2, 3 and 4) combined, as described previously. Table 1 shows that the maternal dietary effect on plasma cholesterol and HDL-cholesterol in the offspring was only seen in males. Whilst the dietary effect was significant for plasma triacylglycerol in both males and females, there was a greater effect in males. Indeed there was a significantly greater effect of diet in males than females (interaction between sex and diet) for HDL-cholesterol (P = 0.03) and some evidence for this interaction for total cholesterol (P = 0.09) and triacylglycerol (P = 0.09). The interaction between sex and diet justifies analysing the data according to sex.
The present study shows that, in the rat, maternal nutrition in pregnancy and lactation has a long-term programming effect on lipid metabolism in the offspring. Dams given a low-protein diet (80 g/kg chow) in both pregnancy and lactation had offspring that in adult life (mean age 6 months) had significant reductions in plasma concentrations of cholesterol, HDL-
Table 1. Plasma values of cholesterol, HDL-cholesterol and triacylglycerols in male and female rats of mean age 6 months in all four experimental groups combined, together with the difference, and 95% confidence interval (CI) for difference, in plasma cholesterol, HDL-cholesterol and triacylglycerol levels between feed group 1 (controls) and the three reduced maternal protein intake groups (2, 3 and 4) combined according to sex†

(Mean values with their standard errors)

<table>
<thead>
<tr>
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<th>All four experimental groups combined</th>
<th>Difference between feed groups: 1 v. 2 + 3 + 4§</th>
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<tr>
<td></td>
<td>Males (n 33)</td>
<td>Females (n 42)</td>
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<td></td>
<td>Mean  se</td>
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<tr>
<td></td>
<td></td>
<td>95% CI for difference between males and females</td>
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<tr>
<td>Cholesterol (μmol/l)</td>
<td>2.21 0.20</td>
<td>1.76* 0.07</td>
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<tr>
<td>HDL-cholesterol (μmol/l)</td>
<td>1.86 0.20</td>
<td>1.50 0.06</td>
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<tr>
<td>Triacylglycerols (μmol/l)</td>
<td>3.05 0.34</td>
<td>1.67*** 0.16</td>
</tr>
</tbody>
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Mean value was significantly different from that for males: * P < 0.05, *** P < 0.001.
Difference was statistically significant: † P < 0.05, †† P < 0.01.
†† For details of procedures, see pp. 606–607.
§ A positive value denotes a higher level in controls.
cholesterol and triacylglycerols compared with those of controls. Whether this pro-
gramming effect was due to pre- or postnatal dietary manipulation was explored in two
smaller crossover groups in which the progeny of dams fed on regular 200 g protein/kg
chow were suckled by dams fed on an 80 g protein/kg chow (postnatal low protein group)
and vice versa (prenatal low protein group). In these groups, plasma values of the three
lipids in the progeny were low and similar to those of the progeny of dams who received
the low-protein diet during both pregnancy and lactation, providing evidence that, in
the rat, the critical period for long-term programming of lipid metabolism extends from fetal
life into the sucking period.

We chose a low-protein dietary intervention because use of this model had proven
successful in demonstrating programming, in the offspring, of insulin secretion and diabetic
tendency (Snoeck et al. 1990), blood pressure (Langley-Evans et al. 1994), and obesity
(Anguita et al. 1993). It might be argued that this model has limited application to humans
in whom isolated protein deficiency would be uncommon. However, the present study
supports the general concept that, in rats at least, manipulation of the nutritional milieu of
the fetus and suckled animal could have lifelong effects. It remains important to dissect
which specific classes of nutrients are critical in the programming process, and further
nutrients are now being investigated. We emphasize that the present study explores the
impact of specific nutritional intervention in the dam. We recognize that protein deficiency
in the dam could affect availability of many other nutrients (i.e. cause more general
undernutrition) in the fetus or sucking animal; we are currently exploring this.

The observed interaction between sex and diet is of wider biological interest in relation
to the hitherto unexplained increased vulnerability to early nutrition of males (Smart, 1977;
Lucas et al. 1990). For instance, using the same model, Anguita et al. (1993) showed a
diet–sex interaction in relation to the development of obesity and Desai et al. (1996) showed
such an interaction in relation to growth and development of specific organs, notably
muscle and pancreas. Possibly nutritional sensitivity in the male relates to the faster growth
of tissues and hence more critical nutritional needs; or perhaps additional genetic material
on the second X-chromosome has protective value in the female.

Our observation that cholesterol and triacylglycerol metabolism may be programmed in
both fetal and sucking animal suggests that in each case the initial programming trigger(s)
may be effective when received either by intravascular (transplacental) or enteral route. The
lifetime programming of lipid metabolism after a brief early dietary manipulation leaves
intriguing questions about the underlying mechanisms. Which organs or organ systems are
programmed? Where does the 'memory' of the early nutritional event reside, given
repeated cell replacement during life? Could the initial event be a programmed change in
gene expression, clonal selection within developing tissues or differential proliferation of
cell lines with different metabolic functions (Lucas, 1994; Desai et al. 1995). Our further
work focuses on these issues.

Our original intention was to test the hypothesis, generated by human epidemiological
work (Fall et al. 1992), that early malnutrition might program later risk of vascular disease,
including elevated plasma cholesterol concentrations. We have apparently shown the
opposite; plasma cholesterol was reduced in the offspring of protein-undernourished dams.
Our findings, however, are consistent with those of Hahn (1984), who showed that
overnutrition in neonatal rats, achieved by reducing litter size during suckling, was
associated with long-term elevation of total cholesterol in plasma. Thus, in both studies, the
less nourished group had lower plasma cholesterol later in life. Using a similar but not
identical rat model to ours, Langley-Evans et al. (1994) found that early maternal
malnutrition programmed higher blood pressure in the offspring; but in our experiments
on the same animals studied here we found the opposite (results not shown), in parallel with
our findings here on cholesterol.
Whilst we have not, therefore, provided direct support for this early malnutrition hypothesis in humans, our findings do at least implicate early nutrition in long-term programming of factors which would be cardiovascular risk markers in man. Our long-term clinical prospective studies in which we have applied randomized early nutritional interventions will eventually provide direct evidence on nutritional programming of cardiovascular disease in humans (Lucas et al. 1990; Lucas & Morley, 1994). We suggest, however, that use of this rat model could be of value in providing insights into the mechanisms of nutritional programming of cardiovascular risk factors and may point to future pertinent clinical studies.

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REFERENCES


