Contributions of maternal and paternal adiposity and smoking to adult offspring adiposity and cardiovascular risk: the Midspan Family Study

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ABSTRACT

Objective: Obesity has some genetic basis but requires interaction with environmental factors for phenotypic expression. We examined contributions of gender-specific parental adiposity and smoking to adiposity and related cardiovascular risk in adult offspring.

Design: Cross-sectional general population survey.

Setting: Scotland.

Participants: 1456 of the 1477 first generation families in the Midspan Family Study: 2912 parents (aged 45–64 years surveyed between 1972 and 1976) who had 1025 sons and 1283 daughters, aged 30–59 years surveyed in 1996.

Main measures: Offspring body mass index (BMI), waist circumference (WC), cardiometabolic risk (lipids, blood pressure and glucose) and cardiovascular disease as outcome measures, and parental BMI and smoking as determinants. All analyses adjusted for age, socioeconomic status and family clustering and offspring birth weight.

Results: Regression coefficients for BMI associations between father–son (0.30) and mother–daughter (0.33) were greater than father–daughter (0.23) or mother–son (0.22). Regression coefficient for the non-genetic, shared-environment or assortative-mating relationship between BMIs of fathers and mothers was 0.19. Heritability estimates for BMI were greatest among women with mothers who had BMI either <25 or ≥30 kg/m². Compared with offspring without obese parents, offspring with two obese parents had adjusted OR of 10.25 (95% CI 6.56 to 13.93) for having WC ≥102 cm for men, ≥88 cm women, 2.46 (95% CI 1.33 to 4.57) for metabolic syndrome and 3.03 (95% CI 1.55 to 5.91) for angina and/or myocardial infarct (p<0.001). Neither parental adiposity nor smoking history determined adjusted offspring individual cardiometabolic risk factors, diabetes or stroke. Maternal, but not paternal, smoking had significant effects on WC in sons (OR=1.50; 95% CI 1.13 to 2.01) and daughters (OR=1.42; 95% CI 1.10 to 1.84) and metabolic syndrome OR=1.68; 95% CI 1.17 to 2.40) in sons.

Conclusions: There are modest genetic/epigenetic influences on the environmental factors behind adverse adiposity. Maternal smoking appears a specific hazard on obesity and metabolic syndrome. A possible epigenetic mechanism linking maternal smoking to obesity and metabolic syndrome in offspring is proposed. Individuals with family histories of obesity should be targeted from an early age to prevent obesity and complications.

INTRODUCTION

The prevalence of obesity worldwide has increased relentlessly over the past three decades, consuming enormous amount of healthcare resources, directly and indirectly. Excess body fatness and central fat accumulation, as reflected by high body mass index (BMI) or large waist circumference (WC), has consistently been shown to relate to metabolic disturbances which promote cardiovascular disease and premature death, a variety of other morbidities and disabilities, and poor quality of life. Obesity results from energy imbalance, when energy intake from foods is relatively greater than energy expenditure (mainly physiological metabolism and physical...
activity). Little is known about life-course factors (physical or social exposures during different stages of growth and development from gestation, early childhood to later adult life) behind energy imbalance. Heritability of BMI is high, especially in twin studies, but the relative importance of genetic make-up that predisposes individuals to increased appetite or reduced metabolism is disputed. Ultimately, genes must interact with environmental factors affecting food intake and physical activity.

Birth weight, either low or high, has been shown to associate with increased central fat accumulation and a number of health problems including hypertension, diabetes mellitus and cardiovascular disease in adult life. The possibility of epigenetic programming through fetal exposures in pregnancy is increasingly under scrutiny. Animal studies show a variety of maternal factors affecting energy balance and body fat, can be transmitted to offspring through altered DNA methylations, occurring predominantly at position C5 of cytosine in cytosine-guanine dinucleotides in DNA, and other factors and similar processes appear to operate in humans.

The major health consequences from overweight and obesity include the constellation of clinical abnormalities elevating cardiovascular risks characterised as the preventable, and reversible, metabolic syndrome. With elevated WC as its central component, metabolic syndrome approximately doubles the risk of type 2 diabetes, cardiovascular disease and mortality. Not all obese people develop the features of metabolic syndrome, so other genetic/epigenetic and environmental factors must be involved. One of these is smoking, which aggravates all features of metabolic syndrome.

A number of studies have demonstrated maternal smoking to associate with offspring adiposity in adulthood and parental BMI to associate with offspring adiposity in childhood and also in life. Parental socioeconomic factors have also been shown to influence offspring obesity status. In two previous studies from the Midspan Family Study, we have shown that parental BMI correlated highly with adult offspring BMI with a upward shift of BMI by 2-3 kg/m² in the most overweight 5% of the population. The present study further analysed the effects of parental smoking and offspring birth weight on adult offspring body fat and fat distribution. We are not aware if any previous family studies of this kind have been conducted to investigate the influences of parental, especially maternal, lifestyle factors such as smoking and socioeconomic status on adult offspring obesity. Given evidence from existing literature, it is therefore logical to examine adiposity of adult offspring, males and females separately, in conjunction with birth weight, obesity-related complications including cardiometabolic risk factors and cardiovascular disease as health consequences of paternal and maternal exposure factors (BMI and smoking habits).

The present study aimed to identify factors which operate through mothers, specifically, so potentially through epigenetic mechanisms. We investigated separately the contributions of paternal and maternal influences, from their BMI and smoking, on offspring adiposity assessed by BMI and WC, cardiometabolic risks and clinical cardiovascular disease.

**METHODS**

**Study design, setting and participants**

The Midspan Family Study is a general population survey of 1477 Caucasian families in Renfrew and Paisley (Scotland), which included 15 402 participants in the first, parental, generation (aged 45–64 years) surveyed between 1972 and 1976. In the second generation, 2338 offspring (1040 sons and 1298 daughters) aged 30–59 years were surveyed in 1996 from an eligible population of 3202 offspring aged 30–59 years who lived locally. There were 864 eligible offspring who did not take part in the study and 1358 ineligible for the study. After exclusion of 30 step and adopted children, there were 1456 families, 2912 parents with 2308 biological offspring (n=1025 sons and 1283 daughters) available for data analysis. Non-participating offspring included those aged <30 and >59 years, those who had left the area (>50 miles) and those who had not left the area but decided not to take part therefore the average number of offspring recruited per family (1.6) is below the UK national average of 2.1 children born per family in 1951. Parents of the migrants non-participants and parents of offspring participants in the present study had similar BMI, so there is no evidence of ‘migrants being leaner’.

**Anthropometry**

In the offspring study, height was measured in the Frankfort plane to the nearest millimetre using stadiometer (Holtain Ltd, Crymych, UK) and weight to the nearest 0.1 kg using digital scales (Seca, Hamburg, Germany). In the parental study, height was measured to the nearest centimetre and weight was measured to the nearest kilogram. All participants were measured without shoes and wearing indoor clothes. BMI was derived as weight (kg) divided by height squared (m²) to determine lean (<25 kg/m²), overweight (25–29.9 kg/m²) and obese (≥30 kg/m²) categories. Among offspring, WC was measured at midpoint between lowest rib margin and superior anterior suprpubric crest using non-stretchable tape measure and used to categorise by action levels as adopted for diagnosis of metabolic syndrome: action level 1: 94 cm in men, 80 cm in women; action level 2: 102 cm in men, 88 cm in women.

**Cardiometabolic risk factors, cardiovascular disease and diabetes mellitus**

Examinations were carried out in clinics by research nurses to assess the following cardiometabolic risk factors: total and high-density lipoprotein (HDL)
cholesterol, triglycerides and blood glucose were measured from non-fasting venous blood samples. Systolic (first Korotkoff sound) and diastolic (fifth Korotkoff sound) blood pressure (BP) were measured in left arm of the offspring in sitting position and rested for at least 5 min using sphygmomanometer (Dinamap 8100, Kritikon, Tampa, Florida, USA). Clinical history of angina, myocardial infarct, strokes and diabetes mellitus were recorded. People who scored 1 or 2 on the six-point Rose Angina Questionnaire were considered to have angina. Previous myocardial infarct was defined on the basis of self-reported clinical history and/or ECG evidence, identifying cardiac ischaemia or previous myocardial infarct on a score of 1 or 2 on a four-point scale (1=definite, 2=probable, 3=possible and 4=none). Metabolic syndrome was defined according to International Diabetes Federation as WC above action level 1 (men: ≥88 cm, women: ≥80 cm) or above action level 2 (men: ≥102 cm, women: ≥88 cm), angina and myocardial infarct. Additional adjustment for birth weight was also made in subgroup analysis. Both offspring sexes were analysed together only for generalised estimating equations to estimate ORs, adjusted for sex, for angina and myocardial infarct, because of the relatively low numbers of cases with the condition. Most variables had no or few missing data, except for HDL cholesterol (16%), metabolic syndrome (16.8%) and birth weight (19.8%), which were handled in analysis using a ‘listwise deletion of missing data’ approach. Analyses were performed using SPSS V.22.0 (SPSS Inc, Chicago, Illinois, USA). Statistical significance was accepted when p<0.05.

**RESULTS**

Participants’ demographic and clinical characteristics are shown in table 1 as descriptive statistics. At the times of surveys, the mean age of fathers was 55 years, mothers 53 years, and offspring 45 years. Mean BMI of fathers, mothers and daughters were almost identical (26 kg/m²) while that of male offspring was higher by 0.5 kg/m², on average, despite the male offspring being 5–10 years younger than their parents at the time of survey. The prevalence of obesity (BMI ≥30 kg/m²) was higher in offspring than in parents. Male offspring had larger WC, higher levels of triglycerides, lower HDL cholesterol, higher blood glucose and BP and higher proportion of smoking history and myocardial infarct than female offspring. Similar rates of angina among offspring were reported by both sexes.

Table 2 shows the strengths of associations between parental and offspring BMI indicated by regression coefficients (β): there were significant associations between parents and offspring of either sex. In model 1, adjusted for family clustering, parental and offspring age, smoking status and social class showed offspring BMI had greater regression coefficients for associations between father–son (β=0.35) and mother–daughter (β=0.33) than father–daughter (β=0.29) or mother–son.
relationships. In model 2, when further adjustments for mothers’ BMI in father–offspring or fathers’ BMI in mother–offspring analysis were made, there was a reduction in these regression coefficients for father–son (β=0.30), mother–daughter (β=0.33), father–daughter (β=0.25) and mother–son (β=0.22) relationships. We...

Table 1  Descriptive statistics showing characteristics of parents and offspring from the Midspan Family Study

|                          | Fathers (n=1456) | Mothers (n=1456) | Missing cases |
|--------------------------|-----------------|-----------------|---------------|
| **Mean**                 | **SD**          | **Mean**        | **SD**        | **n (%)**     |
| Age (years)              | 54.9            | 5.0             | 52.8          | 4.9           | 0                           |
| BMI (kg/m²)              | 26.0            | 3.3             | 25.9          | 4.3           | 3 (0.1)*                     |
| **n (%)**                |                 |                 |               |               |
| **BMI 25–29.9 kg/m²**    | 746 (51.3)      | 575 (39.5)      | -‡            |
| **BMI ≥30 kg/m²**        | 161 (11.1)      | 214 (14.7)      | -‡            |
| Smoking history†         | 1176 (80.8)     | 782 (53.7)      | 0             |
| **All offspring (n=2308)** |               |                 |               |
| Age (years)              | 44.8            | 6.3             | 45.2          | 6.1           | 0                           |
| BMI (kg/m²)              | 26.5            | 4.0             | 25.9          | 5.0           | 17 (0.7)‡                    |
| WC (cm)                  | 93.5            | 10.8            | 80.1          | 12.1          | 19 (0.8)‡                   |
| Triglycerides (mmol/L)   | 1.93            | 1.51            | 1.31          | 0.80          | 80 (3.5)                     |
| HDL cholesterol (mmol/L) | 1.30            | 0.33            | 1.51          | 0.36          | 370 (16.0)                   |
| Glucose                  | 5.60            | 1.94            | 5.15          | 1.18          | 69 (3.0)                     |
| Systolic BP (mm Hg)      | 131.2           | 15.3            | 123.9         | 15.6          | 25 (1.1)                     |
| Diastolic BP (mm Hg)     | 79.1            | 11.0            | 70.9          | 10.0          | 25 (1.1)                     |
| Birth weight (kg)        | 3.48            | 0.49            | 3.36          | 0.49          | 457 (19.8)                   |
| **n (%)**                |                 |                 |               |               |
| **BMI 25–29.9 kg/m²**    | 456 (44.5)      | 400 (31.6)      | -‡            |
| **BMI ≥30 kg/m²**        | 182 (17.8)      | 229 (18.1)      | -‡            |
| WC between action levels 1 and 2 (%) | 270 (26.4) | 252 (19.9) | -‡ |
| WC above action level 2 (%) | 195 (19.0) | 294 (23.2) | -‡ |
| Smoking history†         | 575 (56.1)      | 647 (50.4)      | 0             |
| Angina§                  | 39 (3.8)        | 51 (4.0)        | 0             |
| Myocardial infarct¶      | 38 (3.7)        | 8 (0.6)         | 22 (1.0)      |
| Angina and/or myocardial infarct | 73 (7.1) | 56 (4.4) | 22 (1.0) |
| Cardiac ischaemia**      | 18 (1.8)        | 25 (1.9)        | 21 (0.9)      |
| Strokes                  | 4 (0.4)         | 8 (0.6)         | 0             |
| Diabetes                 | 13 (1.3)        | 13 (1.0)        | 0             |
| Metabolic syndrome       | 201 (25.1)      | 173 (15.4)      | 387 (16.8)    |

*Missing cases are indicated in the parents BMI row.
†Smoking history=current or former smokers.
‡‡Missing cases are indicated in the offspring BMI and WC rows.
§Angina was based on Rose Angina Questionnaire.31
¶Myocardial infarct was based on clinical history and/or ECG evidence.
**Ischaemia was based on ECG evidence.
BMI, body mass index; BP, blood pressure; HDL, high-density lipoprotein; WC, waist circumference.

Table 2  Regression coefficients (β) and 95% CIs obtained from linear mixed effects model with maximum likelihood, adjusted for family clustering, parental and offspring age, smoking status and social class

|                          | Sons’ BMI | Daughters’ BMI |
|--------------------------|-----------|----------------|
|                          | β         | 95% Confidence interval | p Value | β         | 95% Confidence interval | p Value |
| **Model 1**              |           |                   |         |           |                   |         |
| Paternal BMI             | 0.35      | 0.27 to 0.42       | <0.001  | 0.29      | 0.21 to 0.38        | <0.001  |
| Maternal BMI             | 0.26      | 0.20 to 0.32       | <0.001  | 0.33      | 0.27 to 0.40        | <0.001  |
| **Model 2: additional parental BMI adjustment** |           |                   |         |           |                   |         |
| Paternal BMI (adjusted for maternal BMI) | 0.30      | 0.23 to 0.38       | <0.001  | 0.23      | 0.15 to 0.32        | <0.001  |
| Maternal BMI (adjusted for paternal BMI) | 0.22      | 0.16 to 0.28       | <0.001  | 0.33      | 0.24 to 0.37        | <0.001  |

BMI, body mass index.

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have also observed a significant association between the BMIs of fathers and mothers, who were assumed to not be genetically related (after adjustments for family clustering, parental age, smoking status and social class, $\beta=0.19$, 95% CI 0.14 to 0.25, p<0.001).

Table 3 shows heritability of BMI, estimated as $h^2$ from coefficients for offspring BMI regressed on mid-parental BMI: $h^2$ for BMI was 0.51 in sons and 0.56 in daughters. Further analysis based on parental BMI status showed a clear pattern of $h^2$ for BMI in offspring, with daughters having highest $h^2$ when both parents were lean (0.99) or when at least one parent was obese (0.76). For sons, $h^2$ was also a little above the average (0.56) with two lean parents, but lower at 0.39 for son with at least one obese parent.

Figure 1A, B shows the proportions of offspring with high WC above action level 2: $\geq 102$ cm in males, $\geq 88$ cm in females in different categories based on BMI of their fathers and mothers. With two lean parents, only 8.5% of sons and 14% of daughters had high WC. The rates rose proportionally with number of overweight/obese parents. There were greater proportions of high WC among sons and daughters with obese mothers (31–33%) than with obese fathers (22–26%). When both parents were obese, high WC was much more prevalent among both sons (60.5%) and daughters (71.4%). Figure 1C, D shows similar patterns for proportions of offspring with high BMI $\geq 30$ kg/m$^2$ in different categories based on BMI of their fathers and mothers.

Ever smoking was reported by 2015 (87.3%) in the parents’ generation, and 1222 (52.9%) among the offspring. There was a history of both parents’ smoking for 1098 offspring (47.6%). Among offspring, only 131 (10.7%) reported smoking if both parents never smoked, but 485 (39.7%) if one parent had smoked and 606 (49.6%) if both parents had a smoking history ($\chi^2=10.2$, p=0.006). After adjustment for confounding factors family clustering, sex, offspring and parental age and social class, and parental BMI, compared with offspring whose parents never smoked, offspring with at least one parent with history of smoking had OR for becoming a smoker of 1.46 (95% CI 1.10 to 1.93, p=0.008).

Table 4 shows that after adjustment for family clustering, offspring and parental age and social class, parental BMI and offspring smoking history, increased risk of high BMI $\geq 25$ kg/m$^2$ was observed in sons, but not in daughters, if fathers (OR 1.52) or similarly mothers (OR 1.46) smoked. Increased risk of high WC action level 1 was associated with maternal smoking, both in sons (OR 1.50) and in daughters (OR 1.42), but offspring WC was not related to paternal smoking. These ORs increased further for estimating the risk of offspring for having BMI above 30 kg/m$^2$ or WC above action level 2 from parental smoking history. We did not observe any associations between maternal/paternal smoking history and offspring BP, lipids, diabetes or...
stroke (results not shown). Maternal smoking history was associated with increased risk of metabolic syndrome in sons (OR 1.68) but not in daughters. Paternal smoking history did not associate with metabolic syndrome in offspring of either sex.

Birth weight was added to the analysis for subsets with birthweight data, showing that compared with male offspring whose birth weight was within the middle tertile (referent), those with birth weight in the highest tertile had a 1.98-fold increase in risk of high having WC ≥102 cm. There were no associations between birth weight with metabolic syndrome and cardiovascular disease. Birth weight somewhat diminished the relationships between parental smoking and WC above action level 2 or BMI above 30 kg/m² while accentuating the relationships between parental smoking and WC above action level 1, BMI above 25 kg/m² or metabolic syndrome. There were no associations between offspring in the lowest tertile of birth weight and increased health risk (obesity, metabolic syndrome or cardiovascular disease).

Table 5 shows that, compared with offspring with non-obese parents (BMI <30 kg/m²), the risk for offspring of having a high waist was twofold greater in those with one obese parent. With two obese parents, this risk for offspring rose to 7-fold for having WC above action level 1 and 10-fold for having WC above action level 2. Offspring with two obese parents also had increased risk of having metabolic syndrome (OR 2.46). These results included adjustments for sex, offspring and parental age, smoking status and social class. Similar risks among offspring with obese parents were observed for having high BMI.

Compared with offspring with no obese parents, offspring with two obese parents had adjusted ORs of 3–5 for having angina and/or myocardial infarct. These ORs fell to between 2 and 3 if the analysis was adjusted for BMI of offspring.

We found no differences in prevalences of individual cardiometabolic risk factors between groups of offspring categorised according to their parents’ levels of BMI: lean, overweight or obese (data not presented).

Among the offspring there were 12 who reported strokes, 26 with diabetes and 21 with cardiac ischaemia on ECG. Parental adiposity (based on BMI categories) was not associated with risk of offspring stroke, diabetes mellitus or ischaemia (data not presented).

**DISCUSSION**

**Summary of key results**

This study has demonstrated relationships between the BMI, and smoking habits, of parents on the weight status and cardiovascular risk of their offspring in early middle age. Offspring birth weight had little influences on these health outcomes in adult life. These relationships proved to be robust after adjustment for potential confounders such as age, socioeconomic status and offspring smoking. The influences from mothers’ smoking history tend to be stronger than those from fathers, which can be interpreted to suggest possible epigenetic mechanisms worthy of future exploration.

**Association between parental adiposity and offspring adiposity**

This study has valuably strengthened our understanding of the genetic-environmental influences on obesity, by assessing weight status of offspring when they are middle-aged, when adiposity status is well established.
Table 4  Generalised estimating equations to estimate ORs of the risk of offspring with parental history of smoking for having high waist circumference (WC; above action level 1 or above action level 2), high body mass index (BMI; ≥25 or ≥30 kg/m²)

|          | Sons (n, %) | Daughters (n, %) |          |
|----------|-------------|------------------|----------|
|          | OR 95% CI    | p Value          | OR 95% CI| p Value          |
| Model 1: risk of WC above action level 1 |             |                  |          |
| Paternal smoking history*              | 829 (80.9)  | 1.21 0.84 to 1.75| 0.307    | 1035 (80.7)  | 1.02 0.74 to 1.42| 0.894 |
| Maternal smoking history*              | 555 (54.1)  | 1.50 1.13 to 2.01| 0.006    | 694 (54.1)   | 1.42 1.10 to 1.84| 0.007 |
| At least one parent with smoking history | 892 (87.0)  | 1.76 1.15 to 2.70| 0.010    | 1123 (87.5) | 1.42 0.98 to 2.00| 0.061 |
| Model 2: risk of WC above action level 1 (additional birthweight variable) |             |                  |          |
| Paternal smoking history*              | 640 (80.7)  | 1.20 0.80 to 1.81| 0.378    | 848 (80.2)   | 1.13 0.79 to 1.62| 0.510 |
| Maternal smoking history*              | 428 (54.0)  | 1.56 1.13 to 2.16| 0.008    | 580 (54.8)   | 1.40 1.06 to 1.85| 0.018 |
| At least one parent with smoking history | 684 (86.3)  | 1.97 1.21 to 3.20| 0.007    | 927 (87.6)   | 1.66 1.10 to 2.51| 0.017 |
| High birth weight (highest tertile)†   | 248 (31.3)  | 1.21 0.82 to 1.78| 0.347    | 359 (33.9)   | 1.06 0.77 to 1.46| 0.720 |
| Model 1: risk of WC above action level 2 |             |                  |          |
| Paternal smoking history*              | 829 (80.9)  | 1.09 0.70 to 1.70| 0.697    | 1035 (80.7) | 1.10 0.73 to 1.66| 0.642 |
| Maternal smoking history*              | 555 (54.1)  | 1.67 1.14 to 2.45| 0.009    | 694 (54.1)   | 1.38 1.02 to 1.87| 0.036 |
| At least one parent with smoking history | 892 (87.0)  | 1.54 0.94 to 2.52| 0.086    | 1123 (87.5) | 1.52 0.91 to 2.53| 0.107 |
| High birth weight (highest tertile)†   | 248 (31.3)  | 1.98 1.23 to 3.20| 0.005    | 359 (33.9)   | 1.36 0.92 to 2.00| 0.121 |
| Model 1: risk of BMI above 25 kg/m²    |             |                  |          |
| Paternal smoking history*              | 829 (80.9)  | 1.52 1.05 to 2.20| 0.025    | 1035 (80.7) | 1.22 0.87 to 1.70| 0.244 |
| Maternal smoking history*              | 555 (54.1)  | 1.46 1.09 to 1.97| 0.013    | 694 (54.1)   | 1.13 0.88 to 1.46| 0.335 |
| At least one parent with smoking history | 892 (87.0)  | 2.01 1.33 to 3.03| 0.001    | 1123 (87.5) | 1.43 0.98 to 2.10| 0.067 |
| High birth weight (highest tertile)†   | 248 (31.3)  | 1.28 0.86 to 1.91| 0.226    | 359 (33.9)   | 1.13 0.82 to 1.56| 0.447 |
| Model 2: risk of BMI above 25 kg/m² (additional birthweight variable) |             |                  |          |
| Paternal smoking history*              | 640 (80.7)  | 1.51 1.01 to 2.27| 0.046    | 848 (80.2)   | 1.29 0.90 to 1.65| 0.172 |
| Maternal smoking history*              | 428 (54.0)  | 1.60 1.15 to 2.24| 0.006    | 580 (54.8)   | 1.30 0.93 to 1.81| 0.306 |
| At least one parent with smoking history | 684 (86.3)  | 2.21 1.39 to 3.52| 0.001    | 927 (87.6)   | 1.48 0.85 to 2.58| 0.166 |
| High birth weight (highest tertile)†   | 248 (31.3)  | 1.28 0.86 to 1.91| 0.226    | 359 (33.9)   | 1.13 0.82 to 1.56| 0.447 |
| Model 1: risk of BMI above 30 kg/m²    |             |                  |          |
| Paternal smoking history*              | 829 (80.9)  | 1.54 0.93 to 2.53| 0.091    | 1035 (80.7) | 1.14 0.73 to 1.77| 0.577 |
| Maternal smoking history*              | 555 (54.1)  | 1.79 1.23 to 2.62| 0.003    | 694 (54.1)   | 1.25 0.89 to 1.74| 0.193 |
| At least one parent with smoking history | 892 (87.0)  | 1.95 1.10 to 3.44| 0.022    | 1123 (87.5) | 1.45 0.85 to 2.49| 0.176 |
| High birth weight (highest tertile)†   | 248 (31.3)  | 1.64 0.98 to 2.72| 0.058    | 359 (33.9)   | 1.25 0.83 to 1.89| 0.295 |
| Model 1: risk of metabolic syndrome‡  |             |                  |          |
| Paternal smoking history*              | 646 (80.6)  | 1.12 0.73 to 1.73| 0.669    | 910 (81.3)   | 0.87 0.56 to 1.36| 0.544 |
| Maternal smoking history*              | 433 (54.1)  | 1.68 1.17 to 2.40| 0.005    | 602 (53.8)   | 1.18 0.81 to 1.71| 0.387 |
| At least one parent with smoking history | 692 (86.4)  | 1.72 1.01 to 2.94| 0.047    | 979 (87.4)   | 0.92 0.56 to 1.51| 0.727 |
| High birth weight (highest tertile)†   | 248 (31.3)  | 1.64 0.98 to 2.72| 0.058    | 359 (33.9)   | 1.25 0.83 to 1.89| 0.295 |
| Model 2: risk of metabolic syndrome (additional birthweight variable):‡ |             |                  |          |
| Paternal smoking history*              | 496 (80.1)  | 1.00 0.65 to 1.54| 0.995    | 746 (80.6)   | 1.00 0.63 to 1.60| 0.981 |
| Maternal smoking history*              | 330 (53.3)  | 1.83 1.28 to 2.64| 0.001    | 505 (54.5)   | 1.14 0.77 to 1.69| 0.506 |
| At least one parent with smoking history | 528 (85.3)  | 1.81 1.07 to 3.07| 0.027    | 808 (87.3)   | 1.24 0.70 to 2.19| 0.456 |
| High birth weight (highest tertile)†   | 196 (31.7)  | 1.30 0.88 to 1.93| 0.189    | 322 (34.8)   | 0.97 0.64 to 1.60| 0.884 |

Data were adjusted for offspring and parental age, social class, parental BMI, and offspring smoking history (model 1) and additional birthweight adjustment (model 2).

*Variables were entered in regression simultaneously.
†Reference group: second birthweight tertile.
‡There were 1921 cases with complete data for constructing metabolic syndrome.
This is important as most of the weight gain leading to obesity occurs in early adulthood: while only 12% of males and 20% of females have BMI >30 kg/m² when aged 16–24 years almost 40% of all UK adults become obese by late middle age. Studying offspring at younger ages cannot reliably identify those who will ultimately become obese.

The results in this Caucasian, European population show that although the BMI relationships between parents and adult offspring are statistically significant, the effect sizes relating parental to offspring BMI are relatively small (β range 0.22–0.33), and only slightly greater in size than the BMI association between their non-genetically related fathers and mothers (β 0.19). We have previously published data on parental-offspring BMI relationships showing coefficient values to be lower than those observed in the present study. These discrepancies were due to differences in selection of confounding factors and method of analysis—the study by Johnson et al included marital status and number of children as confounding factors as well as log of BMI and used linear regression models. Analysis in the present study adjusted for family clustering, social class and smoking status for both generations using linear mixed effects model. Despite these differences, the patterns of parent-offspring associations remained similar for both studies in that same sex associations (father–son, mother–daughter) were stronger than opposite sex associations.

Previous studies on familial obesity tended to focus on the mother–daughter relationship, with fewer published data on father–son relationships, especially among adult offspring. Linabery et al found that maternal BMI had a stronger influence on BMI growth of infants aged 1.5–3.5 years than paternal BMI and similarly Gaillard et al found that childhood overweight was more influenced by maternal obesity than paternal obesity. Gaillard et al suggested direct intrauterine mechanisms as an explanation for this parental difference. Perez-Pastor et al found that girls with obese mothers and boys with obese fathers were more likely to have high BMI in childhood. The much weaker relationship between BMIs of mothers and sons, or fathers and daughters, led the authors to conclude that behaviour, rather than genetics, was the dominant factor determining childhood obesity.

Fleten et al compared the maternal-offspring BMI association with the paternal-offspring BMI association when the offspring were 3 years old also concluded that maternal-offspring association may be explained by shared familial risk factors rather than by the intrauterine environment. Findings from our study in adult offspring are consistent with those from studies of children, showing that both paternal and maternal BMI had significant influences on BMI of sons or daughters. Our study found that maternal BMI had marginally greater impact on daughters than paternal BMI on sons. Our findings of BMI and prevalence of obesity of offspring being higher than that of their parents are consistent with previous studies. The observation that maternal BMI was more strongly associated with offspring birth weight than was paternal BMI, but no
differences in parent-offspring associations for BMI when offspring were between 3 and 39 years led Kívimäki et al. to conclude that higher adult BMI for offspring than for parents is likely explained by environmental influences. Our study found that the parental influence on adiposity was far greater when two parents were obese, but the father–daughter and mother–son influences on BMI were much weaker than with like-sex offspring.

Heritability, genetics and epigenetics of obesity
Heritability estimates were generally higher for daughters than for sons, with highest values in women when their mother’s BMI values were either below 25 or above 30 kg/m². For both sexes, heritability was higher among offspring with thinner parents than among those with fatter parents. Our results support the general conclusion from other studies, that genetic contributions to obesity are rather less than environmental ones, and that the high heritability of obesity, especially documented among twins is more to do with shared environmental factors. In line with studies among children, the present study has shown that intergenerational relationships of BMI were strongest between mothers and daughters and between fathers and sons, which may point to epigenetic effects. Conventional genetics does not appear to provide full explanations for obesity. The most powerful common single-gene influences on human obesity, for example, FTO, have only small effects on BMI, increasing risk of obesity by 1.2-fold. Since obesity is a polygenic disorder, individuals who carry many variants (more than 10) are more susceptible to gaining weight than those who carry only one or two variants. Although obesity clearly has some genetic aetiology, its manifestation always requires an interaction with adverse environmental factors to upset energy balance during weight gain, and to permit weight maintenance at a high level. Elks et al. reviewed and performed metaregression of 88 twin studies and 27 family studies and found that BMI heritability estimates were highly variable and were generally higher from twin studies (0.47–0.90) than family studies (0.24–0.81). Our family study found $h^2$ for BMI to be within this range (0.51 in men to 0.56 in women). We saw a clear contrast in $h^2$ for BMI between women and men, when there were two lean parents ($h^2=0.99$ in women and 0.56 in men) and also when there was at least one obese parent ($h^2=0.76$ in women and 0.39 in men). It should be emphasised that $h^2$ estimate for BMI represents not only parents’ and offspring’s share of half of their genome but also many common environmental factors that are transmitted to the next generation. The association between BMI of fathers and mothers who are not genetically related suggests that a substantial part of the association between parental BMI and offspring BMI were due to non-genetic familial influences, reflecting shared habits and environments which are likely to be passed on to offspring as well. Further evidence to support the implications that adverse lifestyles shared by family members that contribute to obesity arises from the observation of greater risk of offspring obesity from parental smoking history and low social class. Undoubtedly, family traditions of poor diet quality and physical inactivity would play major roles in familial obesity. We did not have data to analyse these aspects, but the very different influences from maternal and paternal smoking weigh against purely social mechanisms. It has been suggested by others that ‘missing heritability’, not fully accounted for by conventional genetics, may be explained by epigenetic mechanisms. The original Renfrew & Paisley Study database did not include information about smoking of their parents (ie, grandparents of the Midspan offspring). It was therefore not possible to adjust the estimates of $h^2$ for parental smoking in both generations.

Parental smoking and offspring adiposity
A number of studies have indicated parental smoking as risk of obesity in offspring. Studies of women aged 17–47 and 13–28 years found that daughters of mothers who smoked during pregnancy had increased the risk of obesity.

Our observation of increased risk of high WC in adult offspring whose mothers smoked is novel and consistent with recent studies of more than 5000 adolescents aged 10–18 years which showed that prenatal maternal exposure to smoking, including passive smoke was associated with increased risk of obesity in adolescents, independent of birth weight. Similar evidence arise from other studies showing that either parent or mothers who smoked during pregnancy had increased risk of obesity in adults. Maternal smoking is also associated with shorter offspring height. It is unlikely that height has significant influences on the relationship between maternal smoking and offspring adiposity since height only explained 2% of the variance in WC in male and 0% in female offspring, and 0% of the variance in BMI in male and 1% in female offspring. In our study, after adjustments for parental and offspring age and social class and parental BMI, maternal smoking had significant effects on sons’ and daughters’ WC, but paternal smoking had no effect. When at least one of the parents smoked, the risk of high BMI was increased in sons indicating interaction or additive effect of both parents’ smoking history. This effect was not observed in daughters. The lack of a similar effect on WC from fathers who smoked could again point towards an epigenetic effect on increased adult central fat distribution, mediated by exposure to
exogenous free radicals in pregnancy resulted in changes such as DNA methylation. This hypothesis would be consistent in general terms with the programming hypothesis of early uterine environment, whereby stress at critical stages of pregnancy resulted in the later adverse health consequences of metabolic syndrome. We found no significant associations between parental smoking history and individual features of metabolic syndrome in offspring, although parental obesity was associated with high WC and coronary heart disease in their offspring. However, we have found that maternal smoking history was significantly associated with increased risk of metabolic syndrome in sons.

**Influences of offspring birth weight on offspring adiposity and health outcomes**

It has been observed that the genetic contributions are more evident for obesity manifesting in early childhood than adult life. Gestational weight gain is related to increased offspring BMI in childhood, adolescence and young adults. Fraser *et al.* analysed over 5000 mother–offspring pairs from a UK prospective pregnancy cohort and found that greater maternal pregnancy weight and gestational weight gain were associated with greater offspring adiposity and adverse cardiovascular risk factors at 9 years of age. These findings have suggested epigenetic programming of early-onset offspring obesity.

A previous study of women has shown an inverse relationship between birth weight and intra-abdominal fat measured by MRI. Evidence from our study of offspring birth weight associating only with increased risk of high WC in male adult life does not add support to the general programming hypothesis of Barker. However, the relationship between birth weight and adult health is complex since abnormal birth weight may mark a range of environmental insults to the fetus. Birth weight does not reflect the complete picture of body morphology such as body lengths or circumferences.

**Parental adiposity and cardiovascular disease**

We found a relationship between parental obesity and offspring angina and myocardial infarct (MI), which was largely explained by relationships between parental obesity with offspring smoking habits and adiposity (either by BMI or WC). However, this relationship remained significant after adjustments for a number of confounding factors that are known to affect BMI including offspring and parental age, smoking history, social class and offspring BMI. This remaining relationship suggests possible underlying genetic/epigenetic factors linking parental obesity and offspring’s risk of heart disease. Various criteria for MI, either on the basis of clinical history, ECG evidence or both, were analysed in the present study, all of which showed similar outcomes in relation to parental adiposity. We have also observed significantly greater risk of metabolic syndrome in offspring with two obese parents.

**Strengths and limitations**

Our study included a large number of well-characterised participants in a stable population, with high response rates (>70%). Its design allowed, unusually and importantly, outcome analysis among offspring as adults, with relevant data for analysis of familial obesity and related diseases and risk factors. To interpret the results, we were able to adjust our data for major confounders, but the possibility of residual confounding always exists in hypothesis-generating research of this kind, so confirmation in future research would be welcomed. It is important to recognise that this study was restricted to a Caucasian population under prevailing conditions stretching back 50–70 years, so caution must be exercised in generalising the results. Bias might have been introduced from outcome measures which were recalled or self-reported and from the lack of information of physical activity. Data were not available to assess maternal smoking specifically during pregnancy. Some mothers may have discontinued smoking during pregnancy which could underestimate our findings, but there was little advice or pressure on women to discontinue smoking in pregnancy at the time of data collection, so most probably continued. Passive smoking (largely from fathers) was also likely to persist during pregnancy. It is possible that for mothers who abstained from smoking during pregnancy, their placental and fetal development could still endure from chronic effects of smoking. We do not have direct evidence for this but this area of study deserves further investigations. On the other hand, it is possible that early childhood contact may be closer with the mothers than fathers could influence offspring behaviours including dietary, smoking and physical activity habits that result in adult obesity.

We assumed that all offspring were biologically related to both parents in the present study and that the parents were not related to one another. Parental differences in the association with their offspring’s BMI could be due to non-paternity leading to weaker paternal-offspring BMI relationship than that of maternal-offspring BMI relationship. There are no specific data from which to estimate the rate of non-biological fathers, so we assume non-paternity in this study would be similar to those in the UK at the time (between 2 and 13%).

At the time of questionnaire completion, the ages of offspring (mean 45, SD 6 years) meant that prevalence of cardiovascular disease was still relatively low, particularly in women, so conclusions can only be drawn for early-onset CHD, and true associations with CHD may still exist. In the present study, h² was estimated from regression coefficients derived from linear mixed model by regressing mid-parental BMI on offspring BMI as suggested by Visscher *et al.* This technique has certain limitations and seems to overestimate h² as observed in some of the values approaching 1 due to high correlations between parental and offspring BMI and when the sample size is small in subgroup analysis.
CONCLUSIONS
Evidence from our study supports the existence of genetic/epigenetic-environmental interaction in the genesis of obesity and coronary heart disease, but points to a dominant environmental influence with rather limited genetic influences, most marked between mothers and daughters, fathers and sons. A possible epigenetic mechanism linking maternal smoking to obesity and metabolic syndrome in offspring is proposed. Given that most obese adults have gained weight during adulthood, individuals with family history of obesity should be targeted from an early age to prevent obesity and complications.

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Contributors
TSH and MEJL articulated the conceptual framework and wrote the first draft. TSH developed the analytical approach and analysed the data. TSH, CLH, CH, JL, MNU, GCMW and MEJL contributed to the final study design, interpretation of data and added intellectual content during manuscript preparation. All authors read and approved the final manuscript. TSH and MEJL are the guarantors for the study.

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