Newborns of obese parents have altered DNA methylation patterns at imprinted genes

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Abstract

BACKGROUND—Several epidemiologic studies have demonstrated associations between periconceptional environmental exposures and health status of the offspring in later life. Although these environmentally related effects have been attributed to epigenetic changes, such as DNA methylation shifts at imprinted genes, little is known about the potential effects of maternal and paternal preconceptional overnutrition or obesity.

OBJECTIVE—We examined parental preconceptional obesity in relation to DNA methylation profiles at multiple human imprinted genes important in normal growth and development, such as: maternally expressed gene 3 (MEG3), mesoderm-specific transcript (MEST), paternally expressed...
gene 3 (PEG3), pleiomorphic adenoma gene-like 1 (PLAGL1), epsilon sarcoglycan and paternally expressed gene 10 (SGCE/PEG10) and neuronatin (NNAT).

METHODS—We measured methylation percentages at the differentially methylated regions (DMRs) by bisulfite pyrosequencing in DNA extracted from umbilical cord blood leukocytes of 92 newborns. Preconceptional obesity, defined as BMI ≥30 kg m$^{-2}$, was ascertained through standardized questionnaires.

RESULTS—After adjusting for potential confounders and cluster effects, paternal obesity was significantly associated with lower methylation levels at the MEST ($\beta = -2.57$; s.e. =0.95; $P =0.008$), PEG3 ($\beta = -1.71$; s.e. =0.61; $P =0.005$) and NNAT ($\beta = -3.59$; s.e. =1.76; $P =0.04$) DMRs. Changes related to maternal obesity detected at other loci were as follows: $\beta$-coefficient was +2.58 (s.e. =1.00; $P =0.01$) at the PLAGL1 DMR and −3.42 (s.e. =1.69; $P =0.04$) at the MEG3 DMR.

CONCLUSION—We found altered methylation outcomes at multiple imprint regulatory regions in children born to obese parents, compared with children born to non-obese parents. In spite of the small sample size, our data suggest a preconceptional influence of parental life-style or overnutrition on the (re)programming of imprint marks during gametogenesis and early development. More specifically, the significant and independent association between paternal obesity and the offspring’s methylation status suggests the susceptibility of the developing sperm for environmental insults. The acquired imprint instability may be carried onto the next generation and increase the risk for chronic diseases in adulthood.

Keywords
epigenetics; DNA methylation; imprinted genes; offspring; epidemiology; Newborn Epigenetics STudy

INTRODUCTION

Epidemiologic studies, some with long-term follow-up, have demonstrated the importance of periconceptional nutrition of the mother as a crucial factor for optimal development and the long-term health status of the next generation. Early developmental exposures to maternal undernutrition or conditions of famine have been related to coronary heart disorders, increased body mass index (BMI), hypertension, elevated lipid profiles in female offspring, higher risk for breast cancer and mental health problems in later life. These observations have led to concepts of ‘developmental programming through intrauterine exposures or maternal nutrition’ and are consistent with Barker’s ‘thrifty phenotype hypothesis’. The transgenerational effects of poor fetal nutrition have in part been attributed to epigenetic changes at imprinted genes, among which there are persistent shifts in DNA methylation. However, the effects of overnutrition are less understood. It is widely accepted that the current ‘westernized’ diet, characterized by high caloric intake and imbalanced nutritional supply, contributes to adverse health effects in the individual, although the effects on the next generation are poorly understood. Epidemiologic and animal studies indicate an association between preconceptional maternal obesity or overnutrition and cardiovascular diseases, metabolic disorders, congenital abnormalities and behavioral problems, such as autism and attention-deficit hyperactivity disorder, in the...
offspring. El Hajj and Haïf demonstrated aberrant methylation at the \textit{mesoderm-specific transcript (MEST)} imprinted gene in newborns from mothers suffering gestational diabetes; and research in animal models recently showed that methylation is affected at the \textit{paternally expressed gene 3 (PEG3)} in oocytes from obese mice. Although the notion of ‘the developmental programming through nutritional exposures’ is confined to maternal exposures only, the paternal impact has not yet been studied in humans. A handful of studies in animal models suggest that preconceptional nutritional conditions of the father may alter metabolic mechanisms in the offspring. Earlier analysis of the Newborn Epigenetics STudy (NEST) subcohort revealed for the first time in humans that paternal BMI may also contribute to the offspring’s epigenetic profile. Lower methylation was observed at the \textit{IGF2} differentially methylated region (DMR) in children from obese fathers, compared with children from non-obese fathers. DMRs are regions of the genome at which multiple adjacent CpG sites show parent-of-origin-specific methylation. Genomic imprinting is defined by expression from either the paternal or the maternal allele. The imprint methylation marks at DMRs are established during gametogenesis, hence we hypothesize that the paternal allele is also susceptible to environmentally induced modifications or damage during sperm development. The aim of the present study was to evaluate the possible effects of parental obesity on DNA methylation at seven imprinted DMRs in NEST newborns, and hence evaluate an extended version of Barker’s hypothesis, namely that nutrition of both parents before conception is important in the developmental programming of the future child. In this analysis we included human imprinted genes involved in early growth regulation, such as \textit{MEST}, \textit{PEG3}, \textit{pleiomorphic adenoma gene-like 1 (PLAGL1)}, \textit{epsilon sarcoglycan} and \textit{paternally expressed gene 10 (SGCE/PEG10)}, \textit{neuronatin (NNAT)} and \textit{maternally expressed gene 3 (MEG3)}. The latter includes two DMRs: one located at the intergenic region between \textit{Delta-like 1 homolog (DLK1)} and \textit{MEG3} (referred as the \textit{MEG3-IG DMR}), and one located at the \textit{MEG3} promoter (referred as the \textit{MEG3 DMR}). Deregulated expression of either of these genes has been associated with enlarged adipocyte size, obesity or several tumor types.

**SUBJECTS AND METHODS**

**Study participants**

Our study includes the first 98 children who enrolled in the prospective cohort NEST, born between July 2005 and November 2006 at Duke University Hospital, Durham, NC, USA. Recruitment strategies have been described in detail elsewhere. In brief, English-speaking pregnant women, who were at least 18 years old and intending to use one of the two obstetric facilities in Durham County, NC, for their obstetric care were recruited by a trained interviewer at the prenatal clinics. Mothers self-administered a questionnaire and were helped by a trained nurse, if needed. Questionnaire data included socio-demographic information (for example, race/ethnicity and education), lifestyle characteristics (cigarette smoking) and detailed anthropometrics (the mother’s and the father’s height, highest and lowest weight ever, and current and usual weight). Medical records were abstracted to verify medical conditions, birth weight and the newborn’s gender. BMI was calculated from the data obtained from maternal or paternal heights and the mother’s weight before pregnancy or the father’s current weight; obesity was defined as BMI $\geq 30$ kg m$^{-2}$. Our analytical study
cohort includes only newborns from whom we obtained either maternal or paternal BMI. Of the 98 infants, 6 participants had missing height or weight information; hence, our study population includes a total of 92 newborns. Seventy-eight mothers (86%) provided detailed information about the biological fathers. One mother gave birth to twins, bringing the total number of newborns with matching paternal data to 79. The study was approved by the Duke University Institutional Review Board.

**Specimen collection**

The umbilical vein was punctured, and cord blood samples were collected within minutes of delivery in a vacuum blood collection tube, coated with K$_3$EDTA. The tubes were inverted gently to mix the anticoagulant with the blood. After centrifugation, the leukocyte-containing buffy coat was isolated and stored at −80 °C. Genomic DNA was extracted using Gentra Puregene Reagents (Qiagen, Valencia, CA, USA).

**DNA methylation analysis**

Bisulfite pyrosequencing assay development and validation have been previously described$^{29,30}$ Methylation at CpG sites was quantitatively measured at seven DMRs. These DMRs included two involved in regulating the DLK1/MEG3 imprinted domain on chromosome 14q32.2 (the MEG3-IG DMR and the MEG3 DMR), one within the MEST promoter at 7q32.2, one at the NNAT locus at 20q11.23, one within the PEG3 promoter region at 19q13.43, one at the PLAGL1 locus at 6q24.2 and one at the SGCE/PEG10 promoter region at 7q21.3. The number of CpG sites studied at each DMR is depicted in Table 2. Genomic DNA (800 ng) was treated with sodium bisulfite, using the EZ DNA Methylation Kit as per the manufacturer’s instructions (Zymo Research, Irvine, CA, USA) to convert unmethylated cytosines to uracils, leaving methylated cytosines unchanged. Bisulfite-converted DNA (40 ng) was amplified by PCR using the PyroMark PCR Kit (Qiagen, Valencia, CA, USA). Specific positions of the primers used have been published earlier.$^{29,30}$ Pyrosequencing was performed in duplicate using a Pyromark Q96 MD pyrosequencing instrument (Qiagen). Of the 92 DNA samples, methylation data were complete for 83 subjects at the MEG3-IG DMR, 86 subjects at the MEG3 DMR, 84 subjects at the MEST DMR, 75 subjects at the NNAT DMR, 84 subjects at the PEG3 DMR, 82 subjects at the PLAGL1 DMR and 84 subjects at the SGCE/PEG10 DMR.

**Statistical methods**

Variables were defined as follows: marital status (living with partner or married versus single), obtained a college degree or not (high education versus low education), maternal age (<30 and ≥30), race/ethnicity (African American, Caucasian and other or not specified), gestation time/age (<37 weeks and ≥37 weeks), maternal smoking status (never/quit smoking when pregnant/continued smoking during pregnancy), gender of the baby (male or female), birth weight of the baby (<2.5 kg versus ≥2.5 kg) and pre-pregnancy maternal or paternal obesity (<30 versus ≥30 kg m$^{-2}$) or BMI (Table 1).

We computed the least square means (or estimated marginal means) of the methylation percentages at each CpG site, accounting for potential effects from multiple laboratory tests on different plates, wells and dates. In order to take into account potential cluster effects, we
used mixed models, where well numbers and dates were included as random effects. The methylation outcomes for children born to non-obese parents are presented in Table 2 and were used as a baseline in Figure 1, where differences in methylation percentage by obesity of either of the parents were computed. For further assessment of the effects of maternal and/or paternal obesity on the methylation levels at each DMR we used multivariate procedures, adjusting for potential confounding and accounting for cluster effects from same person and experimental unit, as well as the covariance at the individual CpGs (Table 3). DNA methylation was the dependent variable, and parental obesity (or BMI) and co-variables were included as described above. Potential confounders were selected on the basis of known or observed associations with DNA methylation at these or other loci and with maternal or paternal obesity. A comparison of the different multivariate analyses, with exclusion or inclusion of most of our variables, did not change our results. We further explored the effects of parental obesity by race and repeated the multivariate analyses in Caucasians and African Americans. However, our statistical power calculations indicated that a stratified analysis represents unstable estimates, caused by the small numbers of obese parents by race. All analyses were based on the available laboratory data for each CpG site at the DMRs. Statistical analyses were conducted in SAS v9.3 (SAS Institute Inc., Cary, NC, USA).

RESULTS

Characteristics of study participants

About 32% of the newborns were born to mothers who were obese before pregnancy, whereas only 20% were born from a father reported as obese; in 12% both parents were obese. Maternal and paternal preconceptional obesity was strongly associated: the odds ratio was 4.9 (95% CI =1.5–15.9). The distributions of the characteristics of the study participants by parental obesity status are shown in Table 1. A little over 50% of the mothers were younger than 30 years old, independent of the obesity subgroup studied. Nearly 43% of the obese mothers were single, whereas only 12% were single in the non-obese subgroup (P =0.001). Maternal obesity was also strongly associated with race: most obese mothers were African American (71%), and most non-obese mothers were Caucasian (73%) (P<0.001). Although not statistically significant, 68% of the obese mothers had no college degree at the time of pregnancy, whereas 49% had not earned a college degree in the non-obese subgroup (P =0.10). Similarly, a non-significant inverse association was found between maternal obesity and smoking: 65% of the obese mothers never smoked, whereas only 44% reported ‘never smoking’ in the non-obese subgroup (P =0.11). About 90% of the pregnancies were full-term (gestation time was ≥37 weeks). Approximately 17% of the newborns had a birth weight lower than 2.5 kg, and gender was equally distributed. Only one child was conceived using assisted reproductive technology, the parents were not obese (data not shown). Paternal obesity was not associated with any of the included characteristics, with the exception of maternal obesity, as described above.

Methylation levels at imprinted genes by obesity of the parents Methylations profiles of newborns with non-obese parents are presented in Table 2, by individual CpG sites at each DMR studied. The calculated changes related to maternal or paternal obesity are shown in
Figure 1. At the $PLAGL1$ DMR, we measured a mean methylation percentage of 59.8% for children born to non-obese parents; and if the mother was obese, an increase of 3.9% in DNA methylation was detected (s.e. =1.6, $P =0.01$). At the $PEG3$ DMR, children born to non-obese parents showed a mean methylation of 37.5%, and if the father was obese, a decrease of 1.5% was detected (s.e. =0.6; $P =0.02$). A mean methylation change of −1.9% was also measured at the $MEST$ DMR of children born to obese fathers, although this was not significant (s.e. =1.0; $P =0.07$).

To determine the independent effects of preconceptional maternal or paternal obesity on offspring DNA methylation at the seven DMRs of the imprinted genes studied, we controlled for potential confounding owing to maternal age, smoking, education, newborn’s gender and race/ethnicity, in addition to experimental batch effects and the different CpGs studied. We found significant associations between paternal obesity and lower DNA methylation at $MEST$ ($P =0.003$), $NNAT$ ($P =0.04$) and $PEG3$ ($P =0.007$) (model 1, Table 3); whereas maternal obesity was associated with an increase in methylation at $PLAGL1$ ($P =0.0005$) and a borderline decrease in methylation at $MEG3$ ($P =0.06$) (model 2, Table 3). Notably, adding both maternal and paternal obesity in the model did not markedly change our results with regard to the exposure through the father, suggesting that the effects of paternal obesity are generally independent from the maternal exposures. If the newborn’s father was obese, the $\beta$-coefficient for $MEST$ was −2.57 ($P =0.008$), the $\beta$-coefficient for $NNAT$ was −3.59 ($P =0.04$) and the $\beta$-coefficient for $PEG3$ was −1.71 ($P =0.005$) (model 3, Table 3). At $PLAGL1$, the individual effect of maternal obesity showed an increase in methylation, with a $\beta$-coefficient of +2.58 ($P =0.01$) (model 3, Table 3). Furthermore, maternal obesity was associated with a decrease in methylation at $MEG3$: the $\beta$-coefficient was −3.42 ($P =0.04$) (model 3, Table 3). At the same locus, an opposite effect was seen in children from obese fathers, with a $\beta$-coefficient of +2.75 ($P =0.11$), but this was not significant.

We further extended our analyses by replacing the dichotomous obesity variable by the continuous BMI variable and found similar results. For instance, after controlling for the same variables as before, our model 3 showed that paternal BMI was related with a decrease in DNA methylation at the $MEST$ DMR, $\beta$-coefficient was −0.16 ($P =0.01$); and at the $PEG3$ DMR, $\beta$-coefficient was −0.10 ($P =0.03$). Although an increase in DNA methylation at the $SGCE/ PEG10$ DMR was positively associated with paternal BMI, $\beta$-coefficient was +0.12 ($P =0.04$). In general, maternal BMI did not affect DNA methylation, with one exception at the $PLAGL1$ DMR, where $\beta$-coefficient was +0.13 ($P =0.32$) and +0.01 ($P =0.91$), respectively. However, we lost significance at the $NNAT$ DMR for paternal BMI and at the $MEG3$ DMR for maternal BMI, $\beta$-coefficients were −0.13 ($P =0.32$) and +0.01 ($P =0.91$), respectively.

**DISCUSSION**

We hypothesized that not only *in utero* exposures but also preconceptional exposures through the father may induce epigenetic shifts at DMRs of imprinted genes in the offspring. Our key finding is that newborns from obese fathers are hypomethylated at the $MEST$, $PEG3$ and $NNAT$ DMRs, independent of maternal obesity and other potential confounders. Obesity
in mothers was associated with an increase in methylation at the PLAGL1 DMR and a decrease at the MEG DMR. When using pre-pregnancy BMI instead of obesity in our multivariate models, we also found a decrease in DNA methylation at the MEST and PEG3 DMRs by paternal BMI and a positive association between DNA methylation at the PLAGL1 DMR and maternal BMI. Our analyses further revealed a positive association between DNA methylation at the SGCE/PEG10 DMR and paternal BMI, suggesting that the effect can go either way (hypo- or hypermethylation), depending on the DMR studied.

Our results are consistent with our earlier findings on the IGF2 DMR, where differences in DNA methylation were detected in children from obese fathers or mothers. Although not reported here, we repeated the experimental analyses on the IGF2/H19 DMRs in a subset of the study population and were able to reconfirm these trends in methylation changes. Although alterations are subtle, methylation differences of similar magnitude have been reported earlier, inclusive of studies on in utero or periconceptional exposures to nutritional deprivation, folate supplements, gestational diabetes, tobacco use, and maternal use of medications including antidepressants and antibiotics. In general, changes in DNA methylation at imprint regulatory regions or loss of imprinting may persist through life and have been correlated with cardiovascular diseases, behavioral disorders, ovarian cancer, cervical cancer, Wilms’ tumor and colorectal cancer. These health defects were mainly related with IGF2 deregulation, but loss of imprinting or abnormal regulation of MEST and PEG3 has also been associated with cancer such as rhabdomyosarcoma and glioma. Loss of expression of the PLAGL1 gene, which was hypermethylated in children from obese mothers, has been detected in basal cell carcinoma, breast cancer, head and neck cancer, and ovarian tumors. Interestingly, PLAGL1, MEST and PEG3, which normally exhibit imprinting with silencing (or methylation) of the maternal allele and expression of the paternal allele, belong to a network of imprinted genes important in embryonic growth that are mainly expressed when rapid body growth is important, during embryogenesis and early postnatal life. It has been suggested that alteration of one member of the network may modulate the expression of several other imprinted genes in this network, leading to several imprinting-related pathologies. MEST and PEG3 are additionally involved in social behavioral phenotypes such as nest-building and lactation in mice. Studies in mouse models suggest that demethylation of the MEST promoter may lead to overexpression of the gene, causing enlargement of adipocytes and enhanced expression of genes related to metabolic conditions, such as diabetes or deficiencies in energy-uptake regulation. Also found to be hypomethylated in offspring from obese fathers, has also been associated with adipocyte and metabolic regulation, as well as with childhood obesity. Follow-up of the NEST population over time will further elucidate potential associations between paternal obesity and phenotypes in these children, such as obesity, diabetes or other disorders. Of note, after adjusting for potential confounders, our regression analyses indicated an inverse but insignificant effect of parental obesity on the MEG3 DMR (Table 3). Loss of imprinting at this locus has also been correlated with tumorigenesis.

In general, the literature supports the idea that maternal lifestyle is important for optimal development of the fetus. Few epidemiologic studies have investigated the potential
influence of paternal lifestyle or body composition on the next generation. Figueroa-Colon et al.\textsuperscript{58} showed that the paternal body fat may be a predictor for percentage of body fat in the offspring. Long-term cohort studies show that overeating before puberty may increase the risk of death from cardiovascular disease or diabetes in the grandchildren; this transgenerational association was only detected through the male line.\textsuperscript{59} Paternal obesity and metabolic effects in the offspring were also studied in the Framingham Heart Study, where aberrant levels of circulating alanine transaminase, a biomarker for liver function and risk for obesity, were measured in offspring from obese fathers.\textsuperscript{60} Observations in animal models are in line with the theory of a paternal impact on the next generation’s gene expression regulation or its metabolism. High-fat diet in male rats results in offspring with altered methylation at a putative regulatory region of the \textit{Interleukin 13 receptor alpha 2} gene and impaired insulin secretion.\textsuperscript{19} Similarly, Wu and Suzuki\textsuperscript{61} showed that parental high-fat diet before pregnancy was associated with aberrant fat accumulation in the rat offspring; although they did not distinguish between maternal and paternal diet, they suggested that high-fat diet may alter the parent’s insulin and glucose metabolism, causing an incomplete erasure of epigenetic marks during gametogenesis in associated genes. Other imbalanced diets, such as a low protein diet in male mice, resulted in changes in hepatic expression of genes involved in lipid and cholesterol biosynthesis.\textsuperscript{20} Similarly, paternal food deprivation in mice affected metabolism-related factors in the offspring, represented by low levels of serum glucose.\textsuperscript{18} The obesity burden and concomitant health problems are global issues. Male obesity interferes with fertility and sperm quality, but these can be bypassed with artificial reproductive techniques regardless of potential obesity-induced epigenetic changes. Our study cohort included only one child conceived through artificial reproductive techniques; in this particular case, parents were not obese. Our cohort further included one pair of twins. In order to exclude any potential epigenetic effects of either of these subjects, we repeated our analyses by eliminating twins and/or the child conceived through artificial reproductive techniques. Our findings were unaltered when analyses were restricted to non-twins and/or children not conceived through any artificial reproductive techniques.

The main limitation of our study is its small sample size; however, we explored multiple DMRs and found results that were consistent with our earlier report,\textsuperscript{21} suggesting that the paternal impact is strong enough to be detected in this small study cohort, whereas an association with maternal obesity is probably more complex and needs a larger sample size and alternative cohort(s) for appropriate evaluation. Other potential concerns are proof of paternity and the fact that paternal anthropometric data were reported by the mothers. However, we do not expect that the methylation outcomes are differential with respect to potential misclassification of these exposures. Consideration should also be given to the ethnicity of our population. Missing paternal data were most prominent among African American women and mothers with lower education. Hence, future studies should address the effects by racial/ethnic groups and level of education. A potential limitation of our study includes also the use of cord blood, with its differential cell counts, as a marker for the newborn’s epigenetic status. However, we used isolated leukocytes, and the epigenetic profile of imprinted genes is expected to be similar across all cell types. We earlier verified the DMR methylation profiles in several cell fractions of umbilical cord blood and found no significant differences across blood fractions at imprinted gene DMRs examined here.\textsuperscript{29}
Despite the limitations, our results suggest that preconceptional parental lifestyle, or body fat composition, may cause trans-generational epigenetic effects. Although the underlying molecular processes are yet unclear, we believe that the conditions associated with obesity, such as elevated hormone levels, insulin resistance or diabetes, not measured in this study, may affect the epigenetic machinery at the level of the developing germ cells, finally causing DNA methylation shifts in the offspring. Further research is necessary to understand the biological mechanisms or targets of obesity-related exposures during gametogenesis, especially in the male germ line. We hypothesize that aside from earlier described maternal effects, also paternally induced modifications in epigenetic patterns may increase the susceptibility for diseases in the offspring at later age.

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Figure 1.
Changes in DNA methylation percentage at the DMRs of imprinted genes by parental obesity. Difference in methylation percentages between children born from obese parents compared with non-obese parents are shown by CpG site for each DMR studied, adjusted for cluster effects. The methylation percentages at baseline, representing the outcome for non-obese parents, are shown in Table 2. Bars represent s.e.
Table 1

Parental and newborn characteristics of the NEST subcohort 2005–2006

|                        | Mothers |                        | Fathers |                        |
|------------------------|---------|-------------------------|---------|-------------------------|
|                        | Obese   | Not obese               | Obese   | Not obese               |
|                        | n       | %                       | n       | %                       |
| Age                    |         |                         |         |                         |
| <30 years              | 16      | 57.1                    | 9       | 56.2                    |
| ≥30 years              | 12      | 42.9                    | 7       | 43.8                    |
| Marital status         |         |                         |         |                         |
| Living with partner    | 16      | 57.1                    | 13      | 81.2                    |
| Single                 | 12      | 42.9                    | 3       | 18.8                    |
| Race                   |         |                         |         |                         |
| Caucasian              | 8       | 28.6                    | 9       | 56.2                    |
| African American       | 20      | 71.4                    | 7       | 43.8                    |
| Other or not specified | 0       | 0                       | 0       | 3                       |
| Education              |         |                         |         |                         |
| Low (no college degree)| 19      | 67.9                    | 8       | 50.0                    |
| High (at least college degree) | 9     | 32.1                    | 8       | 50.0                    |
| Gestation time         |         |                         |         |                         |
| Preterm (<37 weeks)    | 2       | 7.1                     | 2       | 12.5                    |
| Normal (≥37 weeks)     | 26      | 92.9                    | 14      | 87.5                    |
| Smoking                |         |                         |         |                         |
| Mother never smoked    | 17      | 65.4                    | 10      | 62.5                    |
| Quit smoking when pregnant | 6   | 23.1                    | 3       | 18.8                    |
| Smoked during pregnancy| 3       | 11.5                    | 3       | 18.8                    |
| Baby gender            |         |                         |         |                         |
| Male                   | 13      | 46.4                    | 9       | 60.0                    |
| Female                 | 15      | 53.6                    | 6       | 40.0                    |
| Birth weight           |         |                         |         |                         |
| <2.5 kg                | 5       | 17.9                    | 3       | 20.0                    |
| ≥2.5 kg                | 23      | 82.1                    | 12      | 80.0                    |
Abbreviation: NEST, Newborn Epigenetics Study. The NEST subcohort includes families in which babies were born at Duke University Hospital between July 2005 and November 2006. Characteristics are shown for newborns from whom we had either maternal or paternal BMI.
Table 2

Methylation profiles in children from non-obese parents at the DMRs of imprinted genes

| Baseline methylation percentage in children from non-obese parents | MEG3-IG | MEG3 | MEST | NNAT | PEG3 | PLAG1 | SGCE/PEG10 |
|---|---|---|---|---|---|---|---|
| % | s.e. | % | s.e. | % | s.e. | % | s.e. | % | s.e. |
| CpG 1 | 47.4 | 0.9 | 51.1 | 1.4 | 45.6 | 0.5 | 61.5 | 0.8 | 36.0 | 0.4 | 62.8 | 1.3 | 50.2 | 0.5 |
| CpG 2 | 42.5 | 0.9 | 67.9 | 1.6 | 46.4 | 0.5 | 56.0 | 0.8 | 38.5 | 1.2 | 61.2 | 1.3 | 49.0 | 0.5 |
| CpG 3 | 60.2 | 0.9 | 73.4 | 1.1 | 45.8 | 0.5 | 53.0 | 0.7 | 38.3 | 0.4 | 61.3 | 1.3 | 50.9 | 0.5 |
| CpG 4 | 53.0 | 1.1 | 70.5 | 1.2 | 48.5 | 0.6 | 35.3 | 0.5 | 59.1 | 1.2 | 47.2 | 0.5 |
| CpG 5 | 75.5 | 0.8 | 55.0 | 0.9 | 41.0 | 0.4 | 58.6 | 1.0 | 49.4 | 0.7 |
| CpG 6 | 78.3 | 1.0 | 55.8 | 0.5 | 56.2 | 1.1 | 53.0 | 0.5 |
| CpG 7 | 83.0 | 0.6 | 35.9 | 0.4 |
| CpG 8 | 74.6 | 0.9 | 33.1 | 0.6 |
| CpG 9 | 39.8 | 0.4 |
| CpG 10 | 41.2 | 0.6 |
| Mean | 50.8 | 0.7 | 71.8 | 1.0 | 46.6 | 0.5 | 56.9 | 0.8 | 37.5 | 0.3 | 59.8 | 1.1 | 50.0 | 0.5 |

Abbreviations: MEG3, maternally expressed gene 3; MEST, mesoderm-specific transcript; NNAT, neuronatin; PEG3, paternally expressed gene 3; PLAG1, pleiomorphic adenoma gene-like 1; SGCE/PEG10, epsilon sarcoglycan and paternally expressed gene 10. The estimated means of methylation percentages for children born from non-obese parents are presented for each DMR, by the individual CpG site studied. The calculated mean for each DMR is shown at the bottom line.
Table 3

Multivariate analyses: DNA methylation at the DMRs in the offspring in relation to obesity of the parents

|          | MEG3-IG | MEG3 | MEST | NNAT | PEG3 | PLAGL1 | SGCE/PEG10 |
|----------|---------|------|------|------|------|--------|------------|
|          | β       | s.e. | P    | β    | s.e. | P      | β         | s.e. | P    | β    | s.e. | P      | β    | s.e. | P  |
| Model 1  |         |      |      |      |      |        |           |      |      |      |      |        |      |      |    |
| Father obese | +0.38  | 1.43 | 0.79 | +1.04| 1.47 | 0.48   | -2.50     | 0.84 | 0.003| -3.04| 1.45 | 0.04   | -1.70| 0.62 | 0.007| +1.39| 1.47 | 0.34 | +1.26| 0.99 | 0.10 |
| Model 2  |         |      |      |      |      |        |           |      |      |      |      |        |      |      |    |
| Mother obese | +0.20  | 1.26 | 0.87 | -2.94| 1.57 | 0.06   | -0.98     | 0.66 | 0.14 | -0.88| 1.26 | 0.48   | -0.02| 0.56 | 0.97 | -3.03| 0.86 | 0.0005| +1.08| 0.68 | 0.11 |
| Model 3  |         |      |      |      |      |        |           |      |      |      |      |        |      |      |    |
| Father obese | +0.34  | 1.71 | 0.84 | +2.75| 1.74 | 0.11   | -2.57     | 0.95 | 0.008| -3.59| 1.76 | 0.04   | -1.71| 0.61 | 0.005| +2.19| 1.33 | 0.10 | +1.26| 0.80 | 0.12 |
| Mother obese | +0.11  | 1.57 | 0.95 | -3.42| 1.69 | 0.04   | +0.37     | 0.81 | 0.65 | +0.85| 1.60 | 0.59   | +0.46| 0.60 | 0.47 | +2.58| 1.00 | 0.01 | +1.02| 0.75 | 0.17 |

Abbreviations: MEG3, maternally expressed gene 3; MEST, mesoderm-specific transcript; NNAT, neuronatin; PEG3, paternally expressed gene 3; PLAGL1, pleiomorphic adenoma gene-like 1; SGCE/PEG10, epsilon sarcoglycan and paternally expressed gene 10. Obesity was defined as BMI ≥ 30 kg m⁻². All the models were adjusted by age, race, education, smoking status of the mother, gender of the offspring, clusters of experimental tests and CpG sites measured. Model 1 includes paternal obesity, model 2 includes maternal obesity and model 3 includes both paternal and maternal obesity.