Maternal obesity and diabetes induces latent metabolic defects and widespread epigenetic changes in isogenic mice

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Abbreviation: Avy, agouti viable yellow

Intrauterine nutrition can program metabolism, creating stable changes in physiology that may have significant health consequences. The mechanism underlying these changes is widely assumed to involve epigenetic changes to the expression of metabolic genes, but evidence supporting this idea is limited. Here we have performed the first study of the epigenomic consequences of exposure to maternal obesity and diabetes. We used a mouse model of natural-onset obesity that allows comparison of genetically identical mice whose mothers were either obese and diabetic or lean with a normal metabolism. We find that the offspring of obese mothers have a latent metabolic phenotype that is unmasked by exposure to a Western-style diet, resulting in glucose intolerance, insulin resistance and hepatic steatosis. The offspring show changes in hepatic gene expression and widespread but subtle alterations in cytosine methylation. Contrary to expectation, these molecular changes do not point to metabolic pathways but instead reside in broadly developmental ontologies. We propose that, rather than being adaptive, these changes may simply produce an inappropriate response to suboptimal environments; maladaptive phenotypes may be avoidable if postnatal nutrition is carefully controlled.

Introduction

Obesity and its accompanying morbidities, such as type 2 diabetes, are now pandemic. While there is no doubt that obesity has a genetic component, its rapidly increasing prevalence indicates a major environmental contribution; the etiology and familial patterns of obesity and diabetes are thus of great interest. Extensive evidence indicates that maternal nutrition and metabolism can stably affect the phenotypes of offspring, a phenomenon termed fetal programming. Rodent models of maternal overnutrition—diet-induced obesity and diabetes (through high fat and/or sugar diets), genetic manipulation (e.g., ob or db mice) and chemical/surgical induction of diabetes—show a variety of metabolic consequences in offspring, including increased body weight, adiposity, hypertension, insulin resistance, hyperglycemia, hyperphagia and leptin resistance (reviewed in refs. 1 and 3). Of these models, diet-induced maternal obesity and diabetes is most likely to reflect the metabolic syndrome seen in human populations but cannot distinguish the effect of excess fat consumption from the effect of maternal obesity and diabetes per se. Increasing the fat content of a diet also necessitates a relative reduction of other components in the diet. Genetic and pairwise high fat-feeding models have suggested that maternal obesity per se may cause programming effects that are distinct from those of dietary fat intake.

Since fetal programming appears to involve stable changes in phenotype, it has been widely speculated that epigenetic modifications induced by the intrauterine environment are responsible. Epigenetic modifications specify stable states of gene expression; they are laid down largely during early development and remain relatively fixed over a lifetime (and sometimes into succeeding generations). In this view, fetal programming stems from induced epigenetic changes in genes that regulate metabolic pathways and has an adaptive function. But evidence for such targeted epigenetic change is scarce, and studies coupling the physiological and epigenomic consequences of exposure to fetal...
overnutrition, particularly in models of natural-onset obesity and diabetes, are lacking. It is at present unclear to what extent maternal obesity impacts on offspring phenotype via changes to epigenetic states.

We have previously characterized changes in cytosine methylation patterns in isogenic C57BL/6J mice in response to dietary intervention. Cytosine methylation is a key epigenetic modification in vertebrates, acting as a focal point for mechanisms that suppress gene expression. We found spontaneous and widespread methylation variability in genes with functions related to gene expression and development; similar variation has also been observed in humans and methylation at four such methylation-variable loci appear to correlate with body mass index. We found that exposure to dietary methyl donors dramatically increases the extent of this epigenetic variability. We therefore wondered whether similar epigenomic effects would be manifest with exposure to maternal obesity.

Here we have studied the phenotypic and epigenomic effects of maternal obesity in a model of natural-onset obesity and type 2 diabetes, the A\(^p\) mouse. Our system allows comparison of genetically identical mice whose mothers either did, or did not, acquire diabetes and obesity eating a standard chow diet. We find that the offspring of obese mothers have a cryptic phenotype that is unmasked by exposure to a Western-style high fat diet: they remain nearly normal on a standard diet, but on a high fat diet they develop insulin resistance, glucose intolerance and hepatic steatosis. This predisposition is associated with gene expression and epigenomic changes affecting a broad variety of functions.

### Results

#### A model of natural-onset maternal obesity

To observe the phenotypic and epigenomic consequences of natural-onset maternal obesity and diabetes, we exploited the obese yellow phenotype exhibited by A\(^p\) mice. The A\(^p\) allele is unusual in that it is subject to stochastic epigenetic silencing in an isogenic background, so that genetically identical mice can display markedly different phenotypes. Mice carrying an active allele have a characteristic syndrome of fully yellow pelage, obesity and type 2 diabetes; mice carrying a silent allele have agouti fur and a metabolically normal phenotype, termed pseudoagouti. The A\(^p\) allele is congenic with nonagouti (a), and A\(^p\) mice are typically maintained as A\(^p\)/a heterozygotes; in a cross of an A\(^p\)/a dam with an a/a sire, 50% of offspring will be heterozygous for the A\(^p\) allele, and the other 50% will be homozygous a/a, on an isogenic C57BL/6J background.

In this experiment, we compared the a/a offspring of obese yellow diabetic A\(^p\)/a dams with genetically identical a/a offspring of lean pseudoagouti A\(^p\)/a dams (Fig. 1A). In this way, phenotypic and epigenomic parameters could be assessed without confounding genetic variation. Furthermore, the induction of obesity and diabetes in yellow A\(^p\) mice does not involve any external manipulation: the mice are hyperphagic and by 3 mo of age they are obese, demonstrably hyperglycemic, hyperinsulinemic and insulin resistant; thus yellow coat color is inseparable from type 2 diabetes in adult A\(^p\) mice. We chose the A\(^p\) model because the progressive development of metabolic syndrome in yellow A\(^p\) mice more accurately recapitulates the gradual onset of metabolic disease in humans with adult-onset obesity and diabetes. This is in contrast to genetic models of obesity and diabetes, which produce early-onset metabolic disease, or dietary manipulation where animals are forced onto a hypercaloric diet.

#### A Western-style diet precipitates metabolic disease in obese-born offspring

In contrast to the overt defects in glucose metabolism seen in offspring of rodents fed a hypercaloric diet, mice born of obese yellow A\(^p\)/a mothers showed no evidence of disturbed glucose homeostasis when compared with offspring of lean A\(^p\)/a mothers (Fig. 1; Fig. S1). However, offspring of obese mothers (ObC) were slightly but significantly heavier than offspring of lean mothers (LnC) before weaning, and males remained so afterwards (Fig. 1B). We asked if this indicated a cryptic or latent predisposition to metabolic disease that might be unmasked by a dietary challenge. We exposed both groups to a Western-style high fat diet and found that while both groups (LnHF and ObHF) gained more weight on this diet, male ObHF mice became significantly heavier than LnHF by 12 weeks (Fig. 1B). After only three weeks on the Western diet (at 6 weeks of age) male ObHF mice demonstrated glucose intolerance (Fig. 1C) and insulin resistance (Fig. 1D); their female siblings displayed slight insulin resistance but no impaired response to glucose (Fig. 1C and D). Male (and to a lesser extent female) ObHF mice also exhibited a spike in blood glucose soon after intraperitoneal injection with insulin, indicating an impairment in their insulin response (Fig. 1D).

To further assess the metabolic phenotype induced by natural-onset maternal diabetes and obesity, we sacrificed animals at 12 weeks of age for tissue analysis. We analyzed only male offspring as they demonstrated the most marked impairment in metabolic homeostasis. Liver lipidomic profiling revealed that total triacylglycerides (TAGs) and diacylglycerides (DAGs) were significantly higher in offspring of obese mothers, and levels were further elevated by the Western diet (Fig. 2A and B; Fig. S2). H&E stained sections of liver were scored using pathological criteria for nonalcoholic steatohepatitis in humans. The Western diet induced steatosis, which was markedly more severe in ObHF mice (Fig. 2C and D).

We also assessed serum leptin, an adipokine that circulates at levels proportional to body fat. Leptin was higher in ObC mice than controls (Fig. 2E) and was further elevated by the Western diet. It was also raised by the Western diet in offspring of lean mothers. Leptin correlated with body mass in all of the groups (Fig. 2F), suggesting that the observed increases in body weight are due to an increase in adiposity rather than an increase in lean mass.

Taken together, these results indicate that the offspring of obese mothers have a latent defect in lipid and glucose metabolism that is unmasked by an energy-dense Western-style diet.

#### Maternal obesity and diabetes alters hepatic gene expression in male offspring

The phenotypic defects in ObC and ObHF mice cannot have a genetic basis, since the mice are genetically identical to LnC and LnHF mice. In order to obtain a more
detailed picture of the defects induced by maternal obesity and diabetes, we assessed hepatic gene expression in 16 mice (4 from each group) using Affymetrix GeneChip Mouse Gene 1.0 ST arrays, which provide whole transcript coverage of 28,853 genes in the mouse genome. Using the statistical packages Messina and LimmaGP (see Materials and Methods), we identified 45 transcripts (42 unique genes) whose expression was significantly altered by maternal obesity (ObC vs. LnC; Fig. 3A). Candidate gene expression changes identified by LimmaGP were, as expected, concordant across all animals within a group (because it uses a moderated t-statistic), whereas changes identified by Messina had larger effect sizes but more heterogeneity within groups (because it allows for intra-group variation); Messina candidates were also more likely to be associated with a CpG island (Fig. 3A). qPCR was used to interrogate five candidate genes (chosen at random) associated with CpG islands, four of which exhibited altered expression in ObC relative to LnC as predicted by the array (Dbp, Gm129, Gna14, Zbtb16; Fig. S3). Validation included many animals that were not used on the array (17/22), and the same intragroup variation was observed in this larger cohort, particularly in the ObC group; this is consistent with a stochastic epigenetic response to maternal obesity.

A similar pattern of expression changes was found in control mice exposed to the Western diet (LnHF vs. LnC); using the same array analysis parameters we find that 30 of the 45 transcripts (67%) identified as altered in ObC are also significantly altered in LnHF. This suggests that intrauterine and preweaning exposure to maternal obesity induces expression changes in a set of genes that are also responsive to overnutrition in later life. A much larger number of genes exhibited altered expression in the ObHF group compared with the other groups (Fig. S4); this is likely to reflect the obvious liver pathology in the ObHF mice (see Fig. 2C and D), which confounds any attempt to link gene expression changes to the latent predisposition in this group.

Gene ontology (GO) analysis of genes whose expression was altered in ObC as compared with LnC obtained a large set of ontologies, within which metabolic functions were present but not prominent (Fig. 3B). The two genes with the most strongly downregulated expression were ATPASE6 and CYTB, both encoded by the mitochondrial genome. We asked if this decrease reflected a lower mitochondrial copy number in hepatocytes of
We compared the genome wide patterns of CpG island methylation in the livers of ObC and LnC mice (high fat-fed mice were not examined, as their overt metabolic disease would make it impossible to distinguish between methylation changes that were potentially causal of the latent phenotype, and those merely consequential to their disease). As previously described, we made libraries enriched for the unmethylated fraction of DNA by sequential digestion with \( Hpa \) II and \( Mcr \) BC, followed by ligation-mediated PCR. Libraries were hybridized to Agilent Mouse CpG Island 105K arrays covering approximately 16,000 CpG islands. We analyzed 8 male LnC mice and 8 male ObC mice; pooled libraries from 10 LnC controls acted as the reference sample for each array.

We first used principal components analysis (PCA) to visualize the overall distribution of array data. PCA reduces many variables within a complex data set to a few artificial variables, which together account for most of the variance in the data. The first four principal components of our data accounted for over ObC mice, but found no difference in mitochondrial genome copy number between LnC and ObC (Fig. 3C). Among the most strongly upregulated genes are major urinary proteins (MUPs), which have been implicated in regulation of energy expenditure.

Maternal obesity causes widespread epigenetic changes in male offspring. The characteristics of the mice exposed to maternal obesity and diabetes demonstrate that they are programmed in fetal life to respond to an energy-dense diet by developing adiposity, insulin resistance, glucose intolerance and hepatic steatosis. This programming may involve stable alterations in gene expression states, mediated by epigenetic modifications. The molecular basis of epigenetic gene regulation is complex and incompletely characterized; however the role of cytosine methylation has been extensively investigated and is best understood in the suppression of transcription initiation. Cytosine methylation in mammals is almost completely confined to CpG dinucleotides, which are concentrated in “CpG islands” that are transcriptional regulatory elements.

![Figure 2](image-url). Offspring exposed to maternal obesity and diabetes have defects in lipid metabolism which are exacerbated by a Western diet. Total liver (A) triacylglycerol and (B) diacylglycerol levels (LnC n = 4, ObC n = 6, LnHF n = 12, ObHF n = 8), and (C) steatosis scores (LnC n = 4, ObC n = 9, LnHF n = 10, ObHF n = 10) in livers of 12 week old male animals. Representative liver histology sections are shown in (D) (scale bar, 100 μm). (E) Serum leptin levels and (F) relationship between serum leptin levels and body weight at 12 weeks (LnC n = 6, ObC n = 7, LnHF n = 12, ObHF n = 12). Box and whisker plots show the median, 25th and 75th percentile values with whiskers indicating the maximum and minimum; means are indicated by an X. Statistically significant differences are indicated by asterisks: *p < 0.05, **p < 0.01, ***p < 0.001. Right-tailed Fisher’s exact test; LnHF c.f. ObHF steatosis scores: p = 0.012.
half (54.4%) of the variability and are shown in two-dimensional score plots (Fig. 4A). The intermingling of ObC and LnC mice in principal components (PC) 1 and 2 indicates that most variation is common within these two groups. This spontaneous inter-individual variation is reminiscent of the variation we have previously described in C57BL/6J mice.8 However, distinct clustering and separation of ObC and LnC offspring is evident in PC 3 and 4. Taken together, the intermingling in PC 1 and 2 and separate clustering in PC 3 and 4 indicate that intergroup variation resides within a larger context of the spontaneous variation we and others8,10 have previously described. A GO analysis of the 5% most methylation-variable loci in LnC and ObC, which are responsible for PC 1 and 2, produces a set of ontologies related to developmental functions (Fig. 4B) that are nearly identical to the set identified in our previous study of intrinsic epigenetic variation in the C57BL/6J strain.8

We investigated the array probes responsible for the separate clustering of ObC and LnC in PC 3 and 4. Statistical
testing by LimmaGP revealed an enrichment of small \( p \)-values (Fig. 4C), indicating that there are significant methylation differences between the two groups. Limma detected methylation differences at 2350 probes corresponding to 1598 genes (at \( p < 0.01 \)). Given the substantial intragroup variation, we also used Messina to identify stochastic changes in methylation that affected most but not all individuals in the ObC group. Most margin scores were very small (Fig. 4D); only 36 loci showed a margin of > 1.5-fold change. The many more significant methylation changes identified by LimmaGP (Fig. 4C) are therefore likely to be small in magnitude. Validation by bisulfite allelic sequencing showed that 6 of 9 loci examined exhibited small methylation differences (0.9–15.3%, Fig. S5) in ObC animals in the direction predicted by the array. The verification rate (FDR ~0.33) and the small magnitude of the changes are consistent with that of other studies of complex phenotypes using genome-wide strategies.8,27-30 There was no significant overlap between the genes whose expression was changed in the ObC group and the genes whose methylation states differed. However, GO analysis of the genes identified by LimmaGP as differentially methylated (at \( p < 0.01 \)) between the ObC and LnC groups again revealed significant enrichment in ontologies associated with development (Fig. 4E); these are largely the same ontologies (8/10) as those in the intrinsically methylation-variable loci (Fig. 4B). Despite the prominent metabolic phenotype in Ob mice, metabolic functions are not represented in these ontologies. This may suggest that the effects of maternal obesity are much broader than the metabolic aberrations uncovered by a Western diet.

Discussion

We have found that natural-onset maternal obesity and diabetes induces a latent defect in glucose and lipid metabolism in offspring of \( A^{vy} \) mice. This defect is unmasked by a Western-style diet, which causes mice to become insulin resistant and develop hepatic steatosis. Changes in gene expression induced by maternal obesity point to a broad set of functional ontologies, not limited to energy metabolism. Genome-wide assay of CpG island methylation demonstrates that maternal obesity induces widespread methylation changes, which affect functional ontologies relating to development and gene expression. Thus the latent defect in energy metabolism is not prominently reflected in the expression and epigenomic changes induced by maternal obesity.

The latent metabolic phenotype we observed in this study contrasts with those of other rodent models using high fat and sugar diets to mimic maternal overnutrition: in these cases glucose handling defects are apparent in offspring even when they are weaned onto a normal chow diet.31-33 The syndrome of obese yellow \( A^{vy} / a \) mice more closely models the gradual onset and progression of obesity and type 2 diabetes in humans. In this model, \( a / a \) offspring exhibit only subtle defects in lipid metabolism and do not progress to a disease state if they are maintained on a standard chow diet. Exposure to an energy-dense “Western” diet produces overt effects, which result in mice exhibiting the equivalent of nonalcoholic steatohepatitis (NASH) in humans. If this effect occurs in humans, it would suggest that the children of obese and diabetic mothers are at risk of metabolic syndrome but can be protected by avoiding an energy-dense diet.

Fetal programming by maternal nutrition has been proposed as a predictive-adaptive response to environmental conditions, whereby developing offspring adjust their physiology to suit the expected postnatal environment. While the underlying mechanisms are not clear, it is now generally supposed that epigenetic modification of metabolically relevant genes is involved.52 But concrete examples are limited to a handful of genes, where the observed changes are small33-36 and very few exist in the context of maternal overnutrition.57,58 Here we deliberately took an unbiased, genome-wide approach to ask whether epigenetic mechanisms are involved in the response to maternal obesity. While we found that maternal obesity induced methylation changes across the genome and expression changes at a variety of genes, metabolic genes were notable by their relative absence. Instead, the genes whose methylation is affected by maternal obesity cluster in developmental ontologies. This may indicate a greater susceptibility of developmental processes to environmental perturbation, consistent with our previous finding that methylation patterns at genes in developmental ontologies are intrinsically more variable and susceptible to dietary supplements than other parts of the genome.8 The systemic consequences of a perturbation in developmental processes could well underlie the metabolic phenotype and perhaps other cryptic phenotypes programmed by maternal obesity and diabetes. Recent large-scale epidemiological studies on the prevalence of non-metabolic disorders, such as asthma and autism in children of obese mothers, support the idea that other defects may be programmed by maternal obesity.59,60

The methylation changes we observed were small in magnitude, consistent with previous reports of epigenetic programming by us and others.5,11,36,41,42 This raises the obvious question as to the relevance of the methylation changes to the phenotype. Certainly, we did not observe a large overlap between the genes with methylation changes and those with changed expression. This is reminiscent of the findings of Rando and colleagues, where widespread methylation changes in response to paternal high-fat feeding did not specify individual gene expression changes, but were nevertheless linked to a metabolic phenotype in offspring.41 Notably, few metabolic genes were altered in this study, and those that showed a magnitude of methylation change similar to that we observe here.

The latency of the metabolic phenotype in our model permits the examination of methylation and gene expression changes that occur prior to the onset of disease, allowing us to avoid detecting changes that are consequential to disease. Consistent with their metabolic phenotype, the high-fat fed group exhibited gene expression changes related to metabolism; this is a common finding in fetal programming studies, and such changes are often assumed to be inborn. But in our control fed offspring, where disease is not yet present, we find that metabolic genes are practice ably absent from comparisons between lean- and obese-born mice. The possible exceptions are the mitochondrially-encoded genes involved in oxidoreductase activity: mitochondrial activity
is known to play an important role in the pathogenesis of metabolic disease and particularly, in NASH. We speculate that the changes we observe in expression of genes related to non-metabolic functional ontologies may either play an indirect role in regulating metabolism, or indicate other cryptic defects in these mice, programmed by maternal obesity but possibly unrelated to energy metabolism.

The predictive-adaptive view of fetal programming has been heavily influenced by the idea that the infants of undernourished mothers are adapted to an environment with scarce food resources. An alternative possibility is that inappropriate nutrition (undernutrition, overnutrition or a modified supply of key nutrients) programs generalized defects that impair responses to environmental stresses or conditions. In this view, the relative paucity of metabolic genes in our lists of affected loci is not surprising. Since diet is always likely to be among the most prominent environmental conditions, defects related to the handling of nutrients may be the most readily unmasked; in our study this seems to particularly be the case with male offspring. Sexually dimorphic responses have been observed in a number of fetal programming studies (reviewed in ref. 44); while the effect of sex hormones is often invoked as the mediator of sexual dimorphism in the setting of metabolic programming, it may also be that sex-specific epigenetic profiles render either the male or female genomes more vulnerable to the effects of specific environmental insults, either in utero or postnatally.

The fetal response to maternal obesity is clearly maladaptive and the epigenomic response most likely represents some form of damage induced by a suboptimal intrauterine environment. However, the maladapted phenotype is not a fait accompli: When maintained on a healthy diet, the mice never develop metabolic disease. Thus, although risk is programmed, overt disease is avoidable; if this translates to human populations, it highlights the need for multigenerational affirmative intervention to break the cycle of obesity.

Materials and Methods

Mouse breeding and diets. All animals were handled in accordance with good practice as defined by the National Health and Medical Research Council (Australia) Statement on Animal Experimentation and requirements of state government legislation. The study was approved by the Garvan/St. Vincent’s Animal Ethics Committee (06/12 and 09/12).

The A/a mice used in this study were descended from an inbred C57BL/6 colony at Oak Ridge National Laboratories and have been maintained at the Victor Chang Cardiac Research Institute since 2001. All parental mice were fed ad libitum on NIH-31 control diet (5% w/w fat, 13.5 MJ/kg). a/a offspring were weaned onto either NIH-31 or SF00-219 high-fat diet designed to mimic a Western fast-food diet [equivalent to Harlan Teklad TD88137; 22% w/w fat (40% digestible energy), 0.15% w/w cholesterol, 19.4 MJ/kg]; A/a littermates were culled at weaning. Feeds were manufactured by Specialty Feeds.

Glucose and insulin tolerance tests. For metabolic testing, mice were fasted for 6 h and given an intraperitoneal injection of either 1.5 g glucose/kg mouse weight (for glucose tolerance test) or 0.75 units of insulin/kg mouse weight (for insulin tolerance test). Blood glucose levels were then measured at 15 min (glucose) or 10 min (insulin) intervals using an Accu-Chek II Performa® glucose monitor (Roche Diagnostics GmbH).

Lipid profiling. Lipidomic profiling was performed on liver samples as described previously. Liver histology. Haematoxylin and eosin stained sections were scored by an anatomical pathologist (M.E.B.) for nonalcoholic steatohepatitis according to standardized pathological criteria.

Mitochondrial copy number. Mitochondrial copy number was assessed by qPCR quantification of mitochondrial genes Atp6 and Nd6 normalized to the nuclear gene Ndafv1. Each reaction was done in triplicate with approximately 10 ng of DNA per reaction. See Table S1 for primer sequences.

Leptin assay. Serum was collected at 12 weeks, and leptin concentrations were measured using the Mouse Leptin ELISA (Millipore) following manufacturer’s instructions.

Statistical analysis for physiological data. All comparisons were made between age-matched, gender-matched, genetically identical mice. Statistical significance was determined by two-way ANOVA with two-tailed Student’s t-tests for glucose and insulin tolerance tests, by two-tailed Student’s t-tests for the other metabolic tests and by a right-tailed Fisher’s exact test for comparison of steatosis scores.

Gene expression. Total RNA was extracted from liver tissue and analyzed with GeneChip® Mouse Gene 1.0ST arrays (Affymetrix). Labeling and hybridization were performed by the Ramaciotti Centre for Gene Function Analysis (University of New South Wales). Microarray data was preprocessed and normalized using robust multichip average (RMA), via the NormalizeAffymetrixST (version 2.0, available at pwbc.garvan.unsw.edu.au/gp) GenePattern module (version 3.2.346). Differential gene expression analysis was performed using LimmaGP and Messina (see below).

LimmaGP analysis. LimmaGP (version 19.3, available at pwbc.garvan.unsw.edu.au/gp; Cowley et al., manuscript in preparation) is a GenePattern module for identifying differential gene expression using limma. Briefly, limma combines linear models to flexibly represent the experimental design of a microarray study in a mathematical framework, with an empirical Bayes, moderated t-statistic, which has more power than the Student’s t-test or ANOVA for analyzing microarray data. We chose a cutoff of FDR < 0.05, with p < 1 × 10−5 and a minimum fold change of 1.3, to capture genes with a consistent change between groups.

Messina analysis. We used Messina as a complimentary candidate identification method. Messina takes a machine learning approach and allows for intra-group heterogeneity by permitting a user-specified, low level of misclassification. This allows detection of candidate genes with inconsistent changes in expression, so is particularly suited to studies where changes may be stochastic. Messina represents the magnitude of differential expression by a margin score (representing the difference in expression between the closest included members of the two groups).
Figure 4. Livers of obese-born males exhibit widespread changes in cytosine methylation. (A) Two-dimensional plots showing principal component (PC) scores of LnC and ObC animals (n = 8 per group). Three plots are shown: scores for PC 1 vs. PC 2 (left), PC 2 vs. PC 3 (middle) and PC 3 vs. PC 4 (right). The amount of variability accounted for by each PC is indicated on the axes. The wireframe ellipses surrounding PC scores for each group are indicative of the overall standard deviation within each group. (B) Bar graph showing −log p-values (with Benjamini–Hochberg correction for multiple testing) for gene ontologies significantly overrepresented in methylation variable genes in LnC and ObC animals. A p-value cutoff of 0.05 is indicated by the red dashed line. (C) Histogram showing the distribution of LimmaG p-values across all probes on the methylation array. A p-value cutoff of 0.01 is indicated by the red dashed line. (D) Histogram showing the distribution of Messina margin scores from the methylation array. Inset shows enlarged view of the histogram tails. (E) Bar graph showing −log p-values (with Benjamini–Hochberg correction for multiple testing) for gene ontologies identified as being significantly overrepresented in genes differentially methylated in ObC vs. LnC (as detected by LimmaG, p < 0.01). A p-value cutoff of 0.05 is indicated by the red dashed line.
set a maximum misclassification of one animal out of four in both control and test groups (i.e., specificity and sensitivity equals 0.75), and applied a margin of ≥ 0.585 (equivalent to ≥ 1.5-fold change).

Microarray validation. One microgram total RNA was reverse transcribed and assayed by qPCR. Each reaction was done in triplicate with approximately 10 ng of cDNA per reaction. Samples were normalized to the geometric mean of four housekeeping genes (Actb, Hprt, Gapdh and Capn1); see Table S1 for primer sequences.

Methylation analysis. DNA was extracted, enriched for unmethylated sequence by HpaII and MspIBC digest, and hybridized to two-color Agilent Mouse 105K CpG island arrays as previously described. Each sample was hybridized against a reference pool made by combining DNA libraries from 10 Lnc offspring. Data was analyzed with LimmaGP and Messina algorithms as described above. Principal component analysis was performed in Partek Genomics Suite (v6.5).

Bisulfite sequencing validation. Allelic methylation patterns at nine candidate loci were assessed by bisulfite allelic sequencing. Two micrograms DNA was treated with sodium bisulfite using E. coli Two micrograms DNA was treated with sodium bisulfite using E. coli HpaII and unmethylated sequence by both control and test groups (i.e., specificity and sensitivity set a maximum misclassification of one animal out of four in both control and test groups (i.e., specificity and sensitivity equals 0.75), and applied a margin of ≥ 0.585 (equivalent to ≥ 1.5-fold change).

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Gene ontology. Gene ontology analysis was performed on gene expression and methylation candidates using the standard workflow in Ingenuity Pathways Analysis (Ingenuity Systems).

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Supplemental Materials
Supplemental materials may be found here: www.landesbioscience.com/journals/epigenetics/article/24656
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