Differential DNA methylation profile in infants born small-for-gestational-age: association with markers of adiposity and insulin resistance from birth to age 24 months

Marta Diaz,1,2 Edurne Garde,1,2 Abel Lopez-Bermejo,3 Francis de Zegher,4 Lourdes Ibañez 1,2

ABSTRACT

Introduction Prenatal growth restraint followed by rapid postnatal weight gain increases lifelong diabetes risk. Epigenetic dysregulation in critical windows could exert long-term effects on metabolism and confer such risk.

Research design and methods We conducted a genome-wide DNA methylation profiling in peripheral blood from infants born appropriate-for-gestational-age (AGA, n=30) or small-for-gestational-age (SGA, n=21, with postnatal catch-up) at age 12 months, to identify new genes that may predispose to metabolic dysfunction. Candidate genes were validated by bisulfite pyrosequencing in the entire cohort. Endocrine-metabolic variables and body composition (dual-energy X-ray absorptiometry) were assessed at birth and at 12 and 24 months.

Results GPR120 (cg14582356, cg01272400, cg23654127, cg03629447), NKX6.1 (cg22588426, cg07688460, cg17758081), CHGA (cg13332653, cg15480367, cg05700406), FABP5 (cg00696973, cg10563714, cg16128701), CTRP1 (cg19231170, cg19472078, cg0164308, cg07162665), GAS6 (N/A), ONECUT1 (cg14217069, cg02061705, cg26155897, cg06657050, cg15446043), and SLC2A8 (cg07688460, cg19021975, cg11312566, cg12281690, cg04016166, cg03804985) genes were hypermethylated in SGA infants. These genes were related to β-cell development and function, inflammation, and glucose and lipid metabolism and associated with body mass index, body composition parameters and markers of insulin resistance at 12 and 24 months.

Conclusion In conclusion, at 12 months, abnormal methylation of GPR120 and NKX6.1 persists and new epigenetic marks further involved in adipogenesis and energy homeostasis arise in SGA infants. These abnormalities may contribute to metabolic dysfunction and diabetes risk later in life.

INTRODUCTION

Low birth weight followed by rapid postnatal weight gain has been associated with increased metabolic risk in adulthood.
Significance of this study

How might these results change the focus of research or clinical practice?

- The degree of abnormal methylation of the reported genes may become a novel biomarker for the early identification of those SGA infants at high risk for developing metabolic disorders later in life.
- Follow-up of these infants with personalized lifestyle measures will possibly prevent a rapid catch-up in body weight that, besides the genetic and epigenetic background, seems to be a key driver for the development of insulin resistance and diabetes.

including for obesity, type 2 diabetes and cardiovascular disorders.\(^1\) The extent of this mismatch between the prenatal and postnatal environments—as judged by the Z-score change from weight at birth to body mass index (BMI) in childhood—appears to be critical in metabolic programming, as it relates closely to central (hepato-visceral fat) and insulin resistance.\(^2\) Along these lines, catch-up infants born small-for-gestational-age (SGA, birth weight below −2 SD), and particularly those who receive enriched formulas and have a faster weight gain, depict low levels of high molecular weight (HMW) adiponectin and are less insulin sensitive and more abdominally adipose than those born appropriate-for-gestational-age (AGA).\(^3\) Although the mechanisms underlying this sequence of events have not been elucidated in full, there is increasing evidence of the key role played by epigenetics—especially DNA methylation—in the fetal programming of adult pathologies.\(^4\) DNA methylation is modulated by environmental factors, particularly by nutrition, and it has been proved that an adverse nutritional environment during the perinatal period can lead to persistent epigenetic modifications producing stable changes in gene expression later in life.\(^5\) Methylation of cytosine bases in CpG dinucleotides induces chromatin modifications leading to repression of gene expression by recruitment of methyl-binding and chromatin remodeling proteins, or by blocking transcriptional machinery from accessing start sites on a gene’s promoter.\(^6\) We have recently reported a differential methylation and expression pattern of genes involved in the regulation of glucose and lipid metabolism in placenta and cord blood of SGA infants, which may influence fetal growth, early adipogenesis and later diabetes risk.\(^7\) Remarkably, DNA methylation levels have been shown to change more rapidly in the immediate postnatal years, tending to stabilize beyond age 7.\(^8\) However, reports on DNA methylation patterns during early childhood are scarce, and none has assessed simultaneously endocrine-metabolic parameters and/or body composition.\(^9\) Here, we tested whether differential DNA methylation patterns persist in catch-up SGA infants—as compared with AGA infants—and whether new epigenetic marks involved in the control of energy metabolism arise by age 12 months. We also assessed the associations among those differentially methylated genes and endocrine-metabolic and body composition variables at age 12 and 24 months.

RESEARCH DESIGN AND METHODS

Study population

The study cohort consisted of 51 infants born either AGA (n=30; 57% female) or SGA (n=21; 57% female) who participated in a previous study assessing DNA methylation in placenta and cord blood of SGA versus AGA infants and its association with endocrine-metabolic parameters and body composition,\(^1\) and who were subsequently followed up to age 24 months.

The specific inclusion criteria were (1) singleton pregnancy, absence of maternal pathology (hypertension, pre-eclampsia or gestational diabetes) and drug addiction; (2) term birth (37–42 weeks) with a birth weight range between −1.1 and +1.1 SD (AGA) and below −2 SD (SGA); (3) spontaneous catch-up in weight and length, defined as weight and length Z-score >−2.0 by age 12 months;\(^1\) (4) auxological (weight, length, BMI and BMI Z-score), endocrine-metabolic assessments and body composition at birth, and at 12 and 24 months (see below); and (5) written, informed consent in Spanish/Catalan language. Exclusion criteria were congenital malformations or complications at birth (need for resuscitation or parenteral nutrition). Eighty-six per cent of AGA infants were exclusively breast fed from 0 to 4 months, versus 59% of SGA infants.

Assessments

Infant’s weight and length were measured at birth and at age 15 days, as reported,\(^1\) and again at 12 and 24 months. BMI was calculated at each time point, and BMI Z-scores were derived from country-specific and sex-specific references.\(^9\)

Blood samples were obtained at birth from the umbilical cord (before placental separation)\(^1\) and in the morning state after an overnight fast at age 12 and 24 months. Whole blood collected in EDTA tubes was used for DNA extraction at age 12 months. Serum samples were also obtained at 12 and at 24 months and kept at −80°C.

The entire cohort (n=51) completed the first year of follow-up including auxological, endocrine-metabolic and body composition assessments. In the subsequent year, n=7 patients dropped out from the study, so that completion of all assessments could only be performed in n=44 infants at age 24 months (26 AGA and 18 SGA) (online supplemental figure 1, flow chart). The characteristics of the patients lost to follow-up did not differ from those who completed the study (data not shown).

Serum glucose was measured by the glucose oxidase method. Insulin, IGF-I and lipids were assessed by immunochemiluminiscence (DPC IMMULITE 2500; Siemens, Erlangen, Germany); intra-assay and inter-assay coefficients of variation were <10%. Insulin resistance was estimated with the homeostatic model assessment (fasting
insulin (mU/L) x fasting glucose (mmol/L)/22.5). HMW adiponectin was assessed by ELISA (R&D Systems, Minneapolis, MN, USA); intra-assay and inter-assay coefficients of variation were <9%.

Body composition was assessed at age 15 days and at 12 and 24 months, by dual X-ray absorptiometry with a Lunar Prodigy coupled to Lunar software (Lunar, Madison, WI, USA), adapted for infants. CVs were <3% for fat and lean mass.

**DNA extraction and bisulfite conversion**

DNA was extracted from peripheral blood samples by the phenol–chloroform method (Promega, Madison, WI, USA), following the manufacturer’s protocol. DNA quality and concentration were assessed using a UV–VIS spectrophotometer (Nanodrop 1000; Agilent Technologies, Wilmington, DE, USA). To minimize batch effect, samples were randomized per birth weight and sex. Bisulfite (BS) conversion of 300 ng DNA was performed using the Methylcode Bisulfite Conversion Kit (Invitrogen, Carlsbad, CA, USA) following manufacturer’s guidelines. Conversion quality control was conducted by quantitative PCR (qPCR) and melting curve analysis.

**DNA methylation microarray in peripheral blood samples**

DNA methylation profiling at age 12 months was performed in eight AGA and eight SGA samples with the Agilent DNA Methylation array (ID 049738; Agilent Technologies), which examines 27,800 highly informative CpG sites located within the proximal promoter regions of 14,475 genes. Nearly 100% of these CpG sites were localized within CpG islands. The process for isolating methylated DNA from purified DNA samples, and labeling and hybridization to Human DNA Methylation Array, was conducted following the manufacturer’s protocol (Agilent Microarray Analysis of Methylated DNA Immunoprecipitation V.1.1; Agilent Technologies), as previously described. 

**Pre-processing and analysis of Agilent DNA methylation microarray**

Microarray image was scanned using Agilent SureScan and raw β values were exported using Agilent’s Feature Extraction Software (V.10.7). Raw methylation data were preprocessed using Agilent Genomic Workbench v6.5 BATMAN algorithm (Bayesian Tool for Methylation Analysis) and normalized to control probes present on the array. Outlier features on the arrays were flagged by the same software package. Values were log2-transformed and logged data were used for principal component analysis (PCA) and for statistical analysis. The Welch T-test was used for identification of differentially methylated genes. An individual probe was considered differentially methylated if its p value was <0.05 (not corrected for multiple testing) and if the effect size (Cohen’s d) of t-tests was ≥0.25 (range 0.25–0.8). The resulting p values for each gene were then adjusted for multiple testing using the Benjamini-Hochberg method with a false discovery rate threshold of 10%.

**GO enrichment and KEGG pathway analysis**

The identified differentially methylated CpGs between AGA and SGA subgroups were assigned to linked genes using the Agilent DNA methylation microarray annotation file. To investigate the biological relevance of these genes, we performed a Gene Ontology (GO) analysis (http://www.geneontology.org). Significantly enriched (p<0.01) GO terms in SGA versus AGA infants were obtained and subdivided into three categories “Biological Processes”, “Molecular functions” and “Cellular components”. Gene networks and canonical pathways representing key genes were identified using the KEGG (Kyoto Encyclopedia of Genes and Genomes) software.

**Selection of candidate genes for validation by bisulfite pyrosequencing**

Genes showing a differential DNA methylation pattern between the AGA and SGA subgroups were ranked according to their significance level, and those with the lowest adjusted p value (≤10⁻³) that were related to energy metabolism were validated in the entire population (30 AGA, 21 SGA) by pyrosequencing. Candidate genes were GPR120 (G protein–coupled receptor 120), NFKX6.1 (NFκx6 homeodomain 1), IGFBP4 (Insulin-like growth factor binding protein 4), FABP5 (Fatty acid binding protein 5), CTRP1 (Complement Clq tumor necrosis factor–related protein 1), GAS6 (Growth arrest specific 6), ONECUT1 (One cut homeobox 1), CHGA (Chromogranin A), SLC2A8 (Solute carrier family 2 member 8), CPTIA (Carnitine palmitoyltransferase 1A), SIK2 (Sirtuin 2), and TRAF2 (TNF receptor associated factor 2).

**Amplification of differentially methylated genes and bisulfite pyrosequencing**

Specific pyrosequencing primers and pre-designed PCR primers flanking all CpG islands from promoter regions of the selected genes were used for validation: GPR120 (PM00044758), chromosome 10, bp 95325316–95327446 (4 CpG sites); NFKX6.1 (PM00019299), chr 4, bp 8517332–8519573 (5 CpG sites); IGFBP4 (PM000181258), chr 17, bp 83599248–83601371 (4 CpG sites); FABP5 (PM00036694), chr 8, bp 82192382–82194533 (5 CpG sites); CTRP1 (PM00182245), chr 17, bp 77019906–77021302 (7 CpG sites); GAS6 (PM00055216), chr 13, bp 11534252–11536397 (4 CpG sites); ONECUT1 (PM00059976), chr 15, bp 53082181–53084445 (5 CpG sites); CHGA (PM00056070), chr 14, bp 93388198–93390313 (4 CpG sites); SLC2A8 (PM00042065), chr 9, bp 130158218–130160383 (6 CpG sites); CPTIA (PM00152285), chr 11, bp 86610165–86612396 (6 CpG sites); SIK2 (PM00154672), chr 11, bp 111471855–111473975 (6 CpG sites); TRAF2 (PM00014302), chr 9, bp 139779587–139781726 (5 CpG sites).
Bisulfite-converted DNA (1 µL) was used as template for each qPCR in a final volume of 25 µL. Hot-start PCR was performed using the PyroMark PCR kit (Qiagen, Germantown, USA) and the correct size of the PCR product was verified by gel electrophoresis. PCR amplimers were purified and analyzed using the PyroMark Q96 ID system and evaluated using the PyroMark software V2.5 (Qiagen). The methylation level of each CpG site was estimated by the proportion of C (%) in each region. The methylation status of each gene was assessed as the percentage of average methylation at targeted CpG sites.

Statistics and ethics
Statistical analyses were performed using SPSS software V23.0. Unpaired t-test was used to study differences between AGA and SGA subgroups. Correlation and stepwise multiple regression analysis were used to study the associations between methylation status at age 12 months, and auxological, endocrine-metabolic and body composition parameters. Covariance analysis was used to adjust for sex and BMI Z-score. The level of significance was set at p < 0.05.

The study was approved by the Institutional Review Board of Hospital Sant Joan de Déu at Barcelona University; written informed consent was obtained before delivery.

RESULTS
Clinical, endocrine-metabolic and body composition variables
Table 1 summarizes the main clinical, endocrine-metabolic and body composition variables of the studied population. At birth, SGA infants displayed lower circulating levels of IGF-I, and less fat, lean mass and abdominal fat, as reported. At age 12 months, SGA infants normalized their circulating IGF-I levels, as well as their lean mass and abdominal fat, although they remained lighter. The pattern was similar at age 24 months.

DNA methylation patterns in SGA versus AGA infants
We identified a total of 129 genes differentially methylated in SGA versus AGA infants at age 12 months. Among those, 51 were hypermethylated and 78 were hypomethylated (online supplemental tables 1–4). There were no differences in methylation profile according to early nutrition (breast vs formula feeding 0–4 months), and thus the results within each the AGA and SGA subgroups were pooled for analysis.

The PCA score plot showed a clear separation between the AGA and SGA subgroups, with distinct DNA methylation profiles aggregating together within the same group (figure 1). The three principal components explained 86.3% of the variance observed between subgroups.

GO functional analysis and KEGG enrichment analysis
To explore potential molecular mechanisms related to birth weight, differentially methylated genes (DMG) were analyzed using GO functional analysis and KEGG enrichment analysis. The enrichment analyses of GO are summarized in table 2. A total of 84 biological processes, 55 molecular functions and 17 cellular components were enriched among the DMG. Regarding biological processes and molecular function, regulation of metabolic processes and organ development together with DNA binding and transcription regulation were the most enriched terms in the respective categories.

The top 10 KEGG pathways of the DMG were enriched in lipid and glucose metabolism, cell development and function, cellular signaling, DNA binding, transcription regulation, regulation of immunity, neural differentiation, and regulation of apoptosis and potassium channel (table 3).

Validation of methylation status by pyrosequencing in SGA versus AGA infants
Bisulfite pyrosequencing confirmed that 10 of the 12 candidate genes were differentially methylated in SGA versus AGA infants; specifically, GPR120 (cg14582356, cg01272400, cg23654127, cg03629447), NKX6.1 (cg22598426, cg07688460, cg17444738, cg12076463, cg10457539), CPT1A (cg14073497, cg00941258, cg12778395) and IGFBP4 (cg15471812) were hypermethylated (all p<0.001), whereas CHGA (cg1332653, cg15480367, cg05700406), FABP5 (cg00696973, cg10563714, cg16128701), CTRP1 (cg19231170, cg19472078, cg0164309, cg07162665, cg17758081, cg18996910, cg06709099), GAS6 (N/A), OECUT1 (cg14217069, cg02061705, cg26158897, cg06657050, cg15446043) and SLC2A8 (cg20758474, cg19021975, cg11312566, cg12281690, cg04016166, cg03804985) were hypomethylated (p=0.002 for FABP5; p>0.0001 for the other five genes) in SGA infants (figure 2). These differences were maintained after adjusting for sex and BMI Z-score. Pyrosequencing did not confirm a differential methylation pattern in SIK2 and TRAF2.

Correlation analyses
Bivariate correlations between gene methylation levels at age 12 months and selected clinical, endocrine-metabolic and body composition parameters are summarized in online supplemental table 5.

Hypermethylation of GPR120, NKX6.1 and CPT1A, as well as hypomethylation of CHGA, CTRP1, GAS6, OECUT1 and SLC2A8 associated with lower BMI Z-scores, and with less fat and lean mass at age 12 months.

At age 24 months, methylation status of GPR120, NKX6.1, CTRP1 and CPT1A still associated with lower fat and lean mass whereas SLC2A8 was positively correlated with lean mass. The methylation pattern of all these genes (except for CPT1A) together with OECUT1 associated with lower BMI Z-score at 24 months. The changes in fat mass between 0–12 and 0–24 months showed a negative correlation with GPR120 and NKX6.1 methylation levels.

The methylation profile of GPR120, NKX6.1, CHGA, GAS6, OECUT1, SLC2A8 and FABP5 associated positively with HOMA-IR at 12 and 24 months, and with 0–12 month and 0–24 month changes in HOMA-IR and IGF-I levels. Hypermethylation of IGFBP4 directly associated with IGF-I levels at 12 and 24 months and with IGF-I changes between 0 and 24 months.
Table 1: Clinical, endocrine-metabolic and imaging data from infants born appropriate-for-gestational-age (AGA, N=30) or small-for-gestational-age (SGA, N=21)

| Anthropometry | Baseline | 12 months | Δ0–12 months | 24 months | Δ0–24 months |
|---------------|----------|-----------|--------------|-----------|--------------|
|               | AGA 30   | SGA 21    | AGA          | SGA       | AGA          | SGA          | AGA          | SGA          | AGA          | SGA          |
| Sex (% female) | 56.70%   | 57.10%    | –            | –         | –            | –            | –            | –            | –            | –            |
| Gestational age (weeks) | 40.2±0.2 | 38.7±0.3† | –            | –         | –            | –            | –            | –            | –            | –            |
| Weight Z-score | 0.1±0.1  | −2.3±0.1† | −0.1±0.3     | −1.5±0.2† | −0.2±0.3     | 0.8±0.2*     | 0.0±0.3      | −1.4±0.3†   | −0.1±0.3     | 0.9±0.3*     |
| BMI Z-score    | 0.5±0.1  | −1.3±0.2† | −0.1±0.3     | −1.2±0.3‡ | −0.6±0.3     | 0.1±0.4      | 1.3±0.8      | −0.1±0.8    | −0.8±0.3     | 1.2±0.3      |
| Placental weight (kg) | 0.6±0.02 | 0.5±0.03* | –            | –         | –            | –            | –            | –            | –            | –            |

Endocrine-metabolic variables

| HOMA-IR       | 1.2±0.3  | 1.0±0.3  | 0.7±0.2     | 0.9±0.3   | −0.5±0.4     | −0.1±0.5     | 1.0±0.4      | 1.0±0.4      | −0.2±0.5     | 0.0±0.6      |
| HMW adiponectin (mg/L) | 40±2     | 39±3     | 20±3        | 21±2      | −20±3       | −18±3        | 14±2         | 13±1         | −26±3       | −26±4        |
| IGF-1 (nmol/L) | 70±10    | 34±2‡    | 62±7        | 66±7      | −8±13       | 32±7‡        | 82±9         | 86±10        | 12±9        | 52±10*       |
| TG (mmol/L)   | –        | –        | 1.2±0.2     | 1.2±0.1   | –           | –           | 0.9±0.1      | 0.8±0.1      | –           | –           |
| HDL-C (mmol/L) | –       | –        | 0.9±0.1     | 1.1±0.1   | –           | –           | 1.2±0.1      | 1.1±0.1      | –           | –           |
| LDL-C (mmol/L) | –       | –        | 2.3±0.1     | 2.2±0.2   | –           | –           | 2.6±0.2      | 2.6±0.2      | –           | –           |

Body composition (DXA)

| Age at DXA (days) | 14±1    | 13±1    | 374±6      | 389±11   | 360±6       | 375±11      | 760±10       | 759±14       | 746±11      | 746±14      |
| Fat mass (kg)     | 0.7±0.1 | 0.5±0.1†| 3.6±0.1    | 3.0±0.1‡ | 2.9±0.1     | 2.6±0.1     | 3.9±0.2      | 3.2±0.2*     | 3.2±0.2     | 2.8±0.2     |
| Abdominal fat (kg) | 0.03±0.00 | 0.02±0.00† | 0.19±0.01 | 0.17±0.02 | 0.16±0.01 | 0.15±0.02 | 0.18±0.02 | 0.14±0.01 | 0.16±0.02 | 0.13±0.01 |
| Lean mass (kg)    | 3.1±0.1 | 2.3±0.1† | 6.9±0.1    | 5.9±0.1   | 3.8±0.1     | 3.7±0.1     | 8.5±0.2      | 8.0±0.2      | 5.9±0.2     | 5.3±0.2     |

Data are mean±SEM. *p<0.05, ‡p<0.01 and †p<0.001 between subgroups. The assessments at 24 months were performed in 26 AGA and 18 SGA infants. The bold values highlight the statistically significant differences.
Circulating levels of LDL-C were associated with the methylation pattern of FABP5, GAS6 and CPT1A genes at age 24 months.

**Multivariate linear models**

In linear regression analysis adjusted for sex and BMI Z-score, methylation status of NKX6.1 and GPR120 were independent predictors of total fat at age 12 months and 24 months, respectively (online supplemental table 6). Lean mass was independently explained by SLC2A8 methylation at the age of 12 months and by ONECUT1 methylation at 24 months (online supplemental table 6). Moreover, SLC2A8 methylation independently explained 42% of the 0–24 month changes in BMI Z-score variability (online supplemental table 6).

**DISCUSSION**

Here, we report for the first time a differential DNA methylation profile in peripheral blood of SGA infants aged 12 months, its association with BMI, body composition parameters and markers of insulin resistance at 12 and at 24 months, and its potential contribution to an altered fat distribution on postnatal catch-up in weight. We identified n=41 differentially methylated CpG sites in gene promoter regions; 13 CpG were hypermethylated and located in GPR120, NKX6.1, CPT1A, IGFBP4 genes and 28 CpG were hypomethylated along the CHGA, FABP5, CTRP1, GAS6, ONECUT1 and SLC2A8 genes.

GPR120 is a functional receptor for n−3 fatty acids and a key regulator of adipogenesis, as well as of energy metabolism, insulin secretion and inflammation.\(^{20-22}\) GPR120 deficiency leads to obesity, insulin resistance and hepatic steatosis in mice fed a high-fat diet.\(^{23}\) In contrast, activation of GPR120 increases insulin secretion, protects pancreatic β cells from inflammation,\(^ {24}\) and reduces fat mass and body weight through activation of brown adipose tissue (BAT) thermogenesis\(^ {25}\); GPR120 is also required for neonatal adaptive thermogenesis in mice.\(^ {26}\) We have previously reported that GPR120 is hypermethylated in cord blood of SGA infants and that it associates with birth weight and reduced fat mass across
ongoing at age 12 months, together with the hypermethylation of ONECUT1, may negatively affect β-cell number, as suggested by the association between the methylation status of these genes and HOMA-IR, fat mass and BMI Z-scores. At age 24 months, SGA infants still have normal levels of IGF-I and HMW adiponectin, less fat mass and are insulin sensitive; however, by age 3 years, SGA children develop high IGF-I levels, a thicker carotida and lower concentrations of HMW adiponectin, supporting the notion that the impairment of insulin action may occur from age 2 years onwards. This phenotype aligns well with the abnormal methylation by age 12 months of several genes related to the control of glucose and lipid metabolism. For example, FABP5, which is involved in the regulation of adipose tissue function and inflammation, was hypomethylated in SGA infants, and transgenic mice models have shown that FABP5 overexpression impairs glucose tolerance, which is in turn reverted in FABP5 knockout mice; FABP5 also plays a role in the development of carotid atherosclerosis. In addition, CTRP1 was also hypomethylated in SGA infants, and is reported to increase in obesity, fatty liver disease, atherosclerosis and type 2 diabetes, and to be associated with major adverse cardiovascular events.

At age 6 years, SGA infants are more insulin resistant and have more pre-peritoneal and hepatic fat than AGA infants. Disruption of lipid homeostasis and reduced mitochondrial function are among the mechanisms that could contribute to this sequence of events. CPT1A, a key regulatory enzyme of β-oxidation required for transport of long-chain fatty acids into mitochondria, was hypermethylated in SGA infants. A decrease in fat oxidation may be followed by fat accumulation; for example, in rodents, the inhibition of fat oxidation results in an increase in intracellular lipids and a decrease in insulin action, whereas in humans an increased respiratory quotient (RQ), indicative of decreased fat oxidation, predicts weight gain and ectopic fat storage and is associated with a deterioration of insulin sensitivity. Inversely, over-expression of CPT1A in the liver of obese mice reduces inflammation and improves insulin signaling. Recently, epigenome-wide association studies have disclosed the causal role of CPT1A methylation in type 2 diabetes and the association between intron 1 CPT1A methylation and gestational BMI.

SLC2A8, also known as GLUT8, is a glucose and fructose transporter highly expressed in oxidative tissues and required for the development of fructose-induced hepatic steatosis. SLC2A8 hypomethylation could additionally contribute to the higher hepatic fat fraction in SGA infants since overexpression of SLC2A8 in hepatocytes represses PPARγ and impairs fatty acid metabolism.

CHGA is a prohormone secreted by neuroendocrine tissues serving as precursor of biologically active peptides including PST (pancrestatin) that interfere with insulin action. PST-treated adipocytes show a decrease in insulin-stimulated lipogenesis, whereas CHGA null mice display increased insulin sensitivity, even after a diet-induced

| Table 2 | Gene Ontology (GO) analysis of differentially methylated genes (n=129) |
|---------|---------------------------------------------------------------|
|         | Observed genes | P value |
| (a) Biological processes | | |
| Regulation of metabolic processes | 39 | 8.30E−05 |
| Glucose metabolism | 4 | 4.70E−05 |
| Lipid metabolism | 4 | 3.40E−05 |
| Organ development | 24 | 7.70E−04 |
| Cellular signaling | 3 | 4.50E−04 |
| Regulation of immunity | 3 | 1.30E−04 |
| Cell adhesion | 5 | 2.40E−03 |
| Cell differentiation | 2 | 1.70E−03 |
| (b) Molecular functions | | |
| Transcription regulation | 23 | 3.70E−04 |
| DNA binding | 32 | 6.30E−03 |
| (c) Cellular components | | |
| Integral to membrane organization | 10 | 1.80E−03 |
| Intrinsic to plasma membrane | 7 | 1.50E−03 |

(a) Biological Processes, (b) Molecular function and (c) Cellular components. Significant enriched components are grouped according to three categories.

early infancy, suggesting that GPR120 could be among the mediators of early fat mass accretion via modulating adipogenesis and lipogenesis. Here, we show that this methylation pattern is maintained in late infancy, and associates with lower BMI Z-scores and less fat mass, and with higher HOMA-IR. IGFBP4, which controls adipose tissue expansion by suppressing IGF-I signaling and angiogenesis promotion, was also hypermethylated in SGA infants. Overall, the combined abnormal methylation of GPR120 and IGFBP4 and ensuing deregulation could result in an impaired adipogenesis in SGA infants.

NKX6.1 is a transcription factor playing a key role in prenatal β-cell differentiation and in postnatal β-cell proliferation and function. This gene was found to be hypermethylated and downregulated in SGA infants at birth, and here we show that this pattern is still detectable at age 12 months. Cells lacking NKX6.1 are unable to express PDX1 and MAFA, both transcription factors needed for the maintenance of β-cell identity and function. Interestingly, a recent report showed that GPR120 prevents lipid-induced β-cell damage through regulation of PDX1 expression. Moreover, ONECUT1—a hepatic transcription factor with a programmed downregulation during pancreas development—acts as MAFA suppressor, and the inappropriate reactivation of this transcriptional regulator occurs in diabetes. It is thus tempting to speculate that the abnormal methylation of GPR120 and NKX6.1 in SGA infants, already detectable at birth and
Table 3  Differentially methylated genes involved in KEGG pathways

| Pathway                        | P value     | Methylation status | Gene symbol                                 |
|--------------------------------|-------------|--------------------|---------------------------------------------|
| Lipid metabolism              | 1.80E−05    | Hyper              | GPR120, CPT1A                                |
|                               |             |                    | FABP5, LMFI, CTRP1, TM6SF2, ELOVL5           |
| Cell development and function | 3.70E−05    | Hyper              | NKP6.1, SMAD7, GATA2B, ADAMTS1, NR2F2, POUSF2, TTC30B, RUNX2, NBW1R1 |
|                               |             | Hypo               | IGFBP4, DNA2, PDZPRN3, AFTPH, EFNAR5, NAPAS3, KCN3C, MICAL2, SLC30A9, SLC22A15, NAPAS3 |
| Glucose metabolism            | 4.20E−05    | Hyper              | RAB4B, *SIK2                                 |
|                               |             |                    | **TRAFO2, GAS5, ONECUT1, MIDN, SLC2A8, CTRP1, GPR4, CHGA |
| Cellular signaling            | 4.30E−04    | Hyper              | ITPRIP, DUSP4, CHRM4, ATP2C1                 |
|                               |             | Hypo               | PPP4R4, PCDHB2, ILK, SLC22A15, FGD5, CLSTN1, GNB1, DUSP7, INSRR, IL17RD, SLC2A8 |
| DNA binding                   | 5.10E−04    | Hyper              | H2AFY, SMAD7, HIST1H2AJ, XRCC5               |
|                               |             |                    | ZNF558, ETV3, ZNF141, ZNF7, EN2, ZNZ3, ZNF462, ZNF7 |
| Transcription regulation      | 8.70E−04    | Hyper              | HLX, BCL6, PATZ1, POUSF2, FOXB1, NR2F2, RUNX2, C1orf113, POUSF1, C7orf64, DDX17 |
|                               |             | Hypo               | JDP2, ETV7, ONECUT1, ZNF132, NAPAS3, HBP1, EOMES, NEUROG2, ZFHX4, RNF4, HBP1, UTF1, SLC30A9, TAF4 |
| Regulation of immunity        | 2.10E−03    | Hyper              | TNFAP8L1, LMBR1                              |
|                               |             |                    | ETV3, SUGT1, UBE2F, LONRF1                    |
| Neural differentiation        | 2.80E−03    | Hyper              | POUSF2, FZD2, RUNX2                           |
|                               |             |                    | NEUROG2                                     |
| Regulation of apoptosis       | 4.10E−03    | Hypo               | TRAF2, ROBO4, PPP1R13B, GAS6                 |
| Potassium channel             | 8.80E−03    | Hyper              | KCN4                                         |
|                               |             |                    | KCN3C, TMEM38A, CLIC6                        |

Pathways are arranged (top to bottom) according to p value. The genes in bold (n=12) were selected for pyrosequencing validation in all the study subjects (n=30 AGA, n=21 SGA).

*Failed genes in pyrosequencing validation.
AGA, appropriate-for-gestational-age; KEGG, Kyoto Encyclopedia of Genes and Genomes; SGA, small-for-gestational-age.

Figure 2  Methylation levels of validated genes in peripheral blood from infants born appropriate-for-gestational-age (AGA, n=30) or small-for-gestational-age (SGA, n=21) at age 12 months. Left panel (A): GPR120, NKP6.1, CPT1A and IGFBP4 were hypermethylated in SGA infants. Right panel (B): CHGA, FABP5, CTRP1, GAS6, ONECUT1 and SLC2A8 were hypomethylated in SGA infants. *p<0.05; **p<0.01; ***p<0.0001.
obesity, highlighting the importance of CHGA–PST interaction in the development of insulin resistance. 47 The hypomethylation of CHGA in SGA infants could hamper further subcutaneous adipogenesis favoring ectopic fat storage on catch-up in weight by inducing adipose tissue dysfunction.

The relevance of GAS6 hypomethylation in SGA infants remains unclear since its role in cancer, obesity, inflammation and insulin resistance remains controversial. 48–50 GAS6 is a member of the vitamin K–dependent protein family that binds to TAM (Tyro3, Axl and Mer) receptors; and high and low levels of GAS6 have been reported, respectively, in overweight and obese adolescents, 49 and in patients with type 2 diabetes. 50

The limitations of the present study include the relatively small size of the studied population, the absence of gene expression assessments, the lack of methylation analysis at age 24 months, the lack of adjustment for cell composition in peripheral blood and the absence of methylation/expression assessments in insulin-target tissues due to ethical restraints. The strengths include the strict inclusion criteria, the use of the same methods over time and the prospective design of the study, allowing to assess the associations of the methylation patterns with endocrine-metabolic and body composition markers over the first 2 years of life.

Overall, our results strengthen the notion that an adverse intrauterine environment can produce long-term changes in gene expression through epigenetic mechanisms, which in turn can predispose to metabolic disorders.

In conclusion, we identified altered epigenetic marks in peripheral blood of SGA infants in genes involved in the control of adipogenesis and energy homeostasis that may exert long-term programming effects and thus increase the risk for obesity and diabetes in this population.

Author affiliations
1Endocrinology Department, Institut Pediàtric Hospital Sant Joan de Déu, University of Barcelona, Esplugues, Barcelona, Spain
2Centro de Investigación Biomédica en Red de Diabetes y Enfermedades Metabólicas Asociadas (CIBERDEM), ISCIII, Madrid, Spain
3Department of Pediatrics, Dr. Josep Trueta Hospital and Girona Institute for Biomedical Research, Girona, Spain
4Department of Development & Regeneration, University of Leuven, Leuven, Flanders, Belgium

Contributors MD contributed to study design, researched data, wrote the manuscript and reviewed/editors manuscript. EG researched data. AL-B and FdZ reviewed/editors the manuscript. LI contributed to study design, wrote the manuscript and reviewed/editors the manuscript. LI and MD are Clinical Investigators of CIBERDEM (Centro de Investigación Biomédica en Red de Diabetes y Enfermedades Metabólicas Asociadas, Instituto de Salud Carlos III, Madrid, Spain). AL-B is a Clinical Investigator of the IS Fund for Scientific Research (Ministry of Science and Innovation, Spain). LI is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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