Global and gene-specific DNA methylation across multiple tissues in early infancy: implications for children’s health research

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ABSTRACT An increasing number of population studies are assessing epigenetic variation in relation to early-life outcomes in tissues accessible to epidemiologic researchers. Epigenetic mechanisms are highly tissue specific, however, and it is unclear whether the variation observed in one of the tissue types is representative of other sources or whether the variation in DNA methylation is distinct, reflecting potential functional differences across tissues. To assess relations between DNA methylation in various samples from newborns and children in early infancy, we measured promoter or gene-body DNA methylation in matched term placenta, cord blood, and 3–6 mo saliva samples from 27 unrelated infants enrolled in the Rhode Island Child Health Study. We investigated 7 gene loci (KLF15, NR3C1, LEP, DEPTOR, DDIT4, HSD11B2, and CEBPB) and global methylation, using repetitive region LINE-1 and ALUYb8 sequences. We observed a great degree of interlocus, intertissue, and interindividual epigenetic variation in most of the analyzed loci. In correlation analyses, only cord blood NR3C1 promoter methylation correlated negatively with methylation in saliva. We conclude that placenta, cord blood, and saliva cannot be used as a substitute for one another to evaluate DNA methylation at these loci during infancy. Each tissue has a unique epigenetic signature that likely reflects their differential functions. Future studies should consider the uniqueness of these features, to improve epigenetic biomarker discovery and translation.—Armstrong, D. A., Lesseur, C., Conradt, E., Lester, B. M., Marsit, C. J. Global and gene-specific DNA methylation across multiple tissues in early infancy: implications for children’s health research. FASEB J. 28, 2088–2097 (2014). www.fasebj.org

Key Words: cord blood • epigenetic epidemiology • placenta • saliva

The term epigenetics refers to heritable changes in gene expression without alterations in the underlying DNA sequence (1). DNA methylation of cytosine residues within CpG dinucleotides is the most commonly studied epigenetic mark (2). It is relevant during development and is thought to mediate, at least in part, epigenetic programming (3, 4). Epigenetics could provide a molecular basis for the epidemiologically derived theory of developmental origin of health and disease (DOHaD; ref. 5). In addition, recent emphasis has been placed on the role of environmental and lifestyle exposures in epigenetic programming and development (6), including prenatal socioeconomic adversity (7), nutrition (8), maternal depression (9), air and water quality (10), smoking (11), and exercise (12).

Epigenetic epidemiology is an emerging field that attempts to elucidate associations of human diseases, disorders, and conditions with epigenetic variation (4, 13, 14). For this purpose, multiple tissue sources have been examined. Peripheral blood is a common source of DNA in adults (15), due to its relative accessibility and because it offers the possibility of prospective sampling, especially attractive in the context of

Abbreviations: AGA, appropriate for gestational age; AluYb8, Arthrobacter luteus subfamily Yb8; CEBPB, CAAT-enhancing binding protein β; DDIT4, DNA-damage-inducible transcript 4 protein; DEPTOR, DEP domain-containing, mTOR-interacting protein; DOHaD, developmental origin of health and disease; HSD11B2, hydroxysteroid dehydrogenase 11 β2; HSF, heat shock factor; KLF15, Kruppel-like factor 15; LEP, leptin; LGA, large for gestational age; LINE-1, long interspersed nuclear element 1; NR3C1, nuclear receptor subfamily 3, group C, member 1; RICHS, Rhode Island Child Health Study; SGA, small for gestational age; Sp1, specificity protein 1; TF, transcription factor

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dynamic epigenetic changes (16). Birth cohort studies, on the other hand, frequently collect cord blood and placenta to assess the newborn’s epigenome (16–19), and evidence from these studies has strengthened the relation between epigenetics and the DOHaD theory. However, cord blood and placenta can be sampled at only one time point, their predictive capability beyond newborn and early-life outcomes has yet to be shown, and their collection requires somewhat complicated and time-dependent protocols that may be available only in large studies at academic medical centers. Similarly, blood sampling through venipuncture is not ideal during early infancy.

Alternatively, saliva is a more practical DNA source, because it can be collected through noninvasive protocols and can be collected more readily at repeated time points throughout childhood and beyond. The use of saliva for clinical assays dates back to 1928 (20), and today it is commonly used for various molecular assessments (21). Recently, an increasing number of epigenetic epidemiologic studies have used saliva as a DNA source. For example, in a study performed in adult twins, saliva DNA methylation at 3 CpG sites in different genes predicted age in a population ranging from 18 to 70 yr (22). Moreover, a recent study (19) found a positive correlation in TACSTD2 promoter methylation between saliva and peripheral blood in 11-yr-old children. This evidence suggests that, at least for some loci, saliva DNA methylation can act as a surrogate for leukocyte methylation, but in general, few data are available on the relationship during early infancy between DNA methylation in saliva samples and other tissue sources, such as placenta and cord blood. Given the high tissue specificity of epigenetic mechanisms, understanding the relationship of epigenetic marks, such as DNA methylation between different tissues, can have important consequences on study design and data interpretation arising from their assessment.

Therefore, we assessed global and gene-specific DNA methylation in paired tissues from newborns: umbilical cord blood and placental tissue and saliva collected in early infancy. Our goal was to investigate the extent to which these tissues could serve as surrogates for one another, particularly for genes relevant to the developmental origins of health and disease during infancy.

**MATERIALS AND METHODS**

**Study population**

Infants included in this study are part of the ongoing Rhode Island Child Health Study (RICHS; ref. 23), which enrolls mother–infant dyads after delivery at Women and Infants Hospital (Providence, RI, USA). Percentiles based on birth weight and gestational age are calculated from the Fenton growth chart (24) and used to enroll term infants born small for gestational age (SGA; <10th percentile) or large for gestational age (LGA; >90th percentile), along with an appropriate for gestational age (AGA) infant matched on sex, gestational age (±3 d), and maternal age (±2 yr). Only singleton, viable infants are included in the study. Other exclusion criteria include maternal age < 18 yr, maternal life-threatening medical complications, and infant congenital or chromosomal abnormalities. Information from the delivery medical record was collected with a structured chart review form, followed by an interviewer-administered structured questionnaire to obtain demographic, lifestyle, and exposure histories. All participants provided written, informed consent for involvement in this study. The protocols used were approved by the Institutional Review Boards of Women and Infants Hospital and Dartmouth College and were performed in accordance with the Declaration of Helsinki.

**Biological sample collection and nucleic acid extraction**

Placental biopsies (n=27) free of maternal decidua were collected within 4 h of delivery. In total, 12 fragments (~1 g), 3 from each quadrant, were excised from the maternal side of the placenta 2 cm from the umbilical cord insertion site. Immediately after collection, the samples were placed in RNAlater (Life Technologies, Carlsbad, CA, USA) and stored at 4°C. After ≥72 h, tissue segments from each placental region were blotted dry, snap-frozen in liquid nitrogen, homogenized by pulverization in a stainless steel cup and piston unit (Cellerusher, Cork, Ireland), and stored at −80°C until needed. When possible, infant cord blood (n=16) was obtained from residual samples collected for clinical purposes with a syringe from the umbilical cord after delivery and stored in blood collection tubes (4°C) until DNA extraction. From 26 of the infants, saliva samples were collected with the OraGene Saliva Collection Kit (DNA Genotek Inc., Ottawa, ON, Canada) at 1 follow-up visit between 3 and 6 mo of age. DNA was extracted from homogenized placental samples and an aliquot of whole umbilical cord blood with the DNeasy Blood and Tissue Kit (Qiagen, Inc, Valencia, CA, USA). DNA isolation from saliva samples was performed with the OraGene extraction kit (DNA Genotek). The DNA was quantified with the ND2000 spectrophotometer (Thermo Fisher Scientific Inc., Waltham, MA, USA). DNA from each sample was sodium bisulfite modified with the EZ DNA Methylation Kit (Zymo Research, Irvine, CA, USA). All tissue samples from the same individual were modified in the same plate. The procedures were performed according to the manufacturer’s instructions.

**Gene selection and DNA methylation analysis**

We selected genes for methylation analysis by using a multifaceted approach. First, we focused on several genes that are currently under study in epidemiologic investigations of DOHaD. Group 1: hydroxysteroid dehydrogenase 11 β2 (HSD11B2; 3 CpGs), nuclear receptor subfamily 3, group C, member 1 (NR3C1; 13 CpGs), and leptin (LEP; 23 CpGs) (25). Second, we included genes that are downstream targets, have been noted to interact with the genes in group 1, or have been reported to be highly expressed in placental tissue. Group 2: DNA-damage-inducible transcript 4 (DDIT4; 4 CpGs), Krüppel-like factor 15 (KLF15; 3 CpGs), CCAAT-enhancer-binding protein β (CEBPB; 7 CpGs), and DEP domain-containing, mTOR-interacting protein (DEPTOR; 4 CpGs) (27–29). Most of the CpGs localize to gene promoter regions for HSD11B2, NR3C1, LEP, CEBPB, and KLF15. However, we also assessed gene body CpG methylation for DEPTOR and DDIT4. Table 1 displays the sequence, chromosomal position, and number of CpG sites for each gene locus. Finally, we examined 2 repetitive-element regions that have been used in epidemiologic and exposure assessment
studies as surrogate measures of global DNA methylation: *Arthrobacter luteus* subfamily Yb8 (*AluYb8*), a short stretch of DNA originally characterized by the action of the *A. luteus* restriction endonuclease, and long interspersed nuclear element 1 (*LINE-1*).

Bisulfite pyrosequencing was performed to detect DNA methylation, as described elsewhere for the genes *HSD11B2* (23, 30), *NR3C1* (31), and *LEP* (26) and the repetitive sequences *LINE-1* (32) and *AluYb8* (33). PCR products were amplified from bisulfite-modified DNA with the Pyromark PCR Kit (Qiagen). PCR and sequencing primers (Integrated DNA Technologies, Coralville, IA, USA) for *HSD11B2*, *LEP*, *NR3C1*, *AluYb8*, and *LINE-1* are presented in Supplemental Table S1. Predesigned Pyromark CpG assays (Qiagen) were used to analyze DNA methylation within the following genes: *CEBPB* (PM00015855), *DDIT4* (PM00044275), *DEPTOR* (PM00018585), and *KLF15* (PM00036400). Cycling conditions for predesigned assays were 94°C for 15 min, followed by 45 cycles at 94°C for 30 s, 55°C for 1 min, and 72°C for 1 min, with a final extension of 7 min at 72°C. PCR products were sequenced with the PyroMark MD instrument (Qiagen), and the percentage of methylation at each CpG site was quantified with Pyro Q-CpG 1.0.11 software (Qiagen). All samples were run in triplicate, and average values were used in subsequent analyses. Non-CpG cytosines in each read served as internal controls to verify bisulfite DNA modification efficiency (≥95% in all samples from 3 tissues). All procedures were performed according to the manufacturer’s protocols.

### Statistical analyses

Spearman rank coefficients were calculated to evaluate correlations between CpG units within the loci. The mean methylation across the CpGs analyzed within the loci was used to obtain an overall measurement for each of the examined regions, which was used in subsequent analyses. Summary statistics, including variances, were obtained for each tissue across all loci studied. Mean methylation between tissues within a given locus was compared by using linear mixed-effects models that accounted for the presence of paired samples. Spearman rank coefficients were calculated to evaluate correlations between tissues for each locus. Statistical significance was set at *P* < 0.05; all tests were 2-sided. All analyses were conducted in RStudio 0.97.314 (RStudio, Boston, MA, USA) using R 3.0.1 (34).

### RESULTS

#### Study population characteristics

Table 2 displays the sociodemographic and clinical characteristics of the study population. The sample of infants enrolled in this study was evenly distributed between males and females and, in accordance with the RICHES cohort design, was oversampled for SGA (11%) and LGA (33%) infants. All infants were considered term (mean gestational age, 39 wk). The majority of the infants (63%) were born by Cesarean section of Caucasian mothers whose age ranged between 18 and 39 yr. Only 1 mother reported smoking during pregnancy. The average age of infant saliva sample collection was 18 wk (range, 13–24 wk).

### Putative transcription factor (TF) binding site maps

We used the TF binding site online database TFSEARCH 1.3 (http://www.cbrc.jp/research/db/TFSEARCH.html) (35) to create an overlapping map of potential TF binding sites associated with the DNA sequences and CpG sites ana-

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**Table 1. Gene loci—chromosomal position: sequence and CpGs for pyrosequencing analysis**

<table>
<thead>
<tr>
<th>Gene locus</th>
<th>Sequence, nonconverted</th>
<th>Chromosomal position</th>
<th>CpGs in assay</th>
<th>CpG chromosomal position</th>
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<tbody>
<tr>
<td><strong>KLF15</strong></td>
<td>TCTGCGTGATTGAGGGGGAAGCG</td>
<td>3:126076655–126076677</td>
<td>3</td>
<td>126076659, −669, −676</td>
</tr>
<tr>
<td><strong>NR3C1(1)</strong></td>
<td>CGGGGACCTGCGGGGGGGGGGGGAGGCTTAG</td>
<td>5:142783588–142783640</td>
<td>5</td>
<td>142783592, −599, −602, −608, −611</td>
</tr>
<tr>
<td><strong>NR3C1(2)</strong></td>
<td>CGGTGCGCCTCTTAAAGCCGGGCAGAGGAGAC</td>
<td>5:142783501–142783585</td>
<td>8</td>
<td>142783501, −503, −513, −519, −533, −555, −570, −573</td>
</tr>
<tr>
<td><strong>LEP(1)</strong></td>
<td>CCGCCGGTGTCCTGCGGGGGGCGAGGCCCTC</td>
<td>7:127881127–127881204</td>
<td>11</td>
<td>127881127, −129, −131, −143, −145, −148, −161, −169, −171, −185, −204</td>
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<tr>
<td><strong>LEP(2)</strong></td>
<td>AAATGCGGGCGGGGGGGAGGCTGGCA</td>
<td>7:127881246–127881350</td>
<td>12</td>
<td>−246, −257, −260, −269, −280, −293, −298, −312, −330, −339, −344, −350</td>
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<tr>
<td><strong>DEPTOR</strong></td>
<td>CGCGGATGCGGGGGCGACAAAGGAGCTGGCACGG</td>
<td>8:120886136–120886172</td>
<td>4</td>
<td>120886137, −146, −151, −171</td>
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<tr>
<td><strong>DDIT4</strong></td>
<td>CCCCGGCCGCTGCGGCTTTTGCTTGGAGCCCG</td>
<td>10:74034388–74034417</td>
<td>4</td>
<td>74034392, −400, −405, −416</td>
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<tr>
<td><strong>HSD11B2</strong></td>
<td>CCGGGACGGTGCTGCTCTTCTCCCGCCGCGC</td>
<td>16:67464387–67464417</td>
<td>4</td>
<td>67464389, −395, −399, −412</td>
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<tr>
<td><strong>CEBPB</strong></td>
<td>CGGGGGCGCGACTCTGCCCCGCTGGGGG</td>
<td>20:48806715–48806742</td>
<td>7</td>
<td>48806715, −720, −722, −732, −736, 738, −741</td>
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DNA methylation in multiple tissues in early infancy

Figure 1 shows diagrams of each gene with sequence position relative to the nearest exons, ATG start site, CpG sites, and potential TFs associated with each sequence. Some common TFs that overlapped CpG sites within the gene groups include alcohol dehydrogenase regulator 1 (ADR1), GATA binding protein 1 (GATA1), heat shock factor (HSF), and specificity protein 1 (Sp1).

DNA methylation correlations between CpGs

We examined DNA methylation across 68 CpG units distributed between different genes and repetitive-elements loci. With the exception of NR3C1 and LEP, the loci examined contained ≤10 CpG units and spanned short sequences. We calculated Spearman rank coefficients between CpGs of each locus across each of the examined tissues. The average coefficients and ranges are presented in Supplemental Table S2. These correlations were locus and tissue dependent, with the placenta exhibiting higher coefficients than the cord blood or saliva. Of note, the saliva samples exhibited the least amount of correlation between the examined CpG units. Given this tissue heterogeneity and according to the generally accepted method in most of the literature, we calculated mean methylation across CpG units of each of the examined regions.

Inter-tissue and interindividual variation in DNA methylation

The distribution of mean DNA methylation in each region is shown in Table 3. Overall, mean DNA methylation varied widely, ranging from 0 to 89%. As expected, global DNA methylation measured at the LINE-1 and AluY8b repetitive elements was higher (>40%) than the methylation observed at the gene-specific loci surveyed in this study. Although higher, mean methylation levels differed significantly between tissues for both the repetitive sequences LINE-1 and AluY8b (P<0.001, all comparisons; Table 3). Mean DNA methylation at the CEBPB, HSD11B2, and LEP promoters was low to moderate, with values ranging from 1.7 to 36.6% and means that differed significantly between tissues (P<0.001, all comparisons; Table 3). In contrast, DEPTOR, DDIT4, KLF15, and NR3C1 mean methylation was <5% in the tissues examined, but similar to our previous results, these 4 low-methylation regions exhibited different mean methylation values between tissues (P<0.001, all comparisons; Table 3).

Table 3 also describes the variance of the mean methylation in each region across tissues. Of the tissues analyzed in this study, the placenta showed the greatest interindividual variability when measured with a dispersion statistic, such as the variance. Of the gene loci examined, placental LEP exhibited considerably more interindividual variability than any of the other regions. Across 7 gene loci, only LEP and CEBPB displayed variability within the 3 tissues analyzed. For HSD11B2 and NR3C1, interindividual variability in DNA methylation was high only in placenta and saliva, respectively. Global DNA methylation at the LINE-1 and ALUYB8 repetitive elements displayed ample variance, especially in placental tissue.

To inquire about interindividual variation within each CpG locus in the regions examined, we estimated variances of the CpG sites of each locus. For global DNA methylation, placental samples had the greatest degree of variability in both elements, but CpG4 in
Figure 1. CpG assay/TF binding site maps. Diagram of chromosomal position for each gene, including the pyrosequencing sequence, the intra-assay CpG locations with overlapping putative TF binding sites, the nearest exon, and the gene ATG start site. CEBPB, HSD11B2, LEP, NR3C1, and KLF15 CpG assays are located in the putative gene promoter region, upstream of the ATG start site for each gene. DEPTOR and DDIT4 are examples of CpG assays located downstream of the ATG start site in the gene body.
AluY8b in saliva had the highest variance. Interestingly, these elements showed greater variation in saliva, followed by placental tissue and then cord blood (Supplemental Table S3). The largest area analyzed, \textit{LEP}, displayed higher variance values for most of the 23 CpGs in placenta and saliva, but not in cord blood (Supplemental Table S4). The 13 CpGs of \textit{NR3C1} exhibited very low variation in all placental and cord blood samples, but at the same loci, methylation variance at saliva CpGs was much higher. The \textit{CEBPB} region showed low variability in placental tissue, intermediate in saliva, and higher in cord blood. \textit{HSD11B2} methylation variability was low in cord blood and high across the loci in placenta and saliva (Supplemental Table S4). Overall, DNA methylation at the examined loci displayed high interindividual variability that was both locus and tissue dependent, and we did not observe any conserved pattern of variation across tissues.

DNA methylation correlations between tissues

Using Spearman’s rank correlations between paired infant tissues, we observed a significant correlation in DNA methylation between cord blood and placental tissue for \textit{AluY8b} ($p=0.57$; $P=0.03$) and a positive correlation for \textit{LINE-1} methylation ($p=0.39$; $P=0.14$; Table 4). Between the placental DNA methylation and saliva methylation data collected in infants between 3 and 6 mo of age, we did not observe correlations of any of the investigated regions. Correspondingly, in comparisons of saliva and cord blood methylation, we did not find significant correlations, with the exception of a significant inverse correlation in the \textit{NR3C1} promoter ($p=-0.58$; $P=0.02$). Hence, in this study, we found scarce intraindividual correlations between tissues in DNA methylation for most of the loci.

DISCUSSION

Several groups have investigated DNA methylation in paired samples from tissues of adult humans (36–39) or related maternal and fetal DNA methylation patterns (40), but only 2 have investigated DNA methylation in different fetal tissues (41, 42). We are the first to assess methylation signatures from healthy, unrelated infants in 3 paired tissues commonly collected in birth cohort studies. We focused on 7 genes that have been analyzed in the DOHaD epidemiologic literature (23, 25, 26, 43) and their potential downstream targets (27) or on genes reported to be highly expressed during fetal development (28, 29).

DNA methylation inhibits gene expression, mainly by 2 mechanisms: preventing the binding of TFs with their cognate DNA recognition sequence (44) or recruiting

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<th>Table 3. Global and gene-specific summary statistics of percentage of mean methylation in placenta, cord blood, and saliva</th>
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<td>\textit{AluY8b}</td>
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$M$, mean; var, variance; min, minimum; max, maximum.

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<th>Table 4. Spearman rank correlations of mean DNA methylation between tissues</th>
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methyl-CpG-binding proteins (MBPs) that can repress transcription (45, 46). Therefore, it is important to understand the genomic context of the regions being examined for DNA methylation by quantitative methylation assessment. In examining putative TF binding sites within the sequences examined, we commonly identified, most notably, HSFs and Sp1 binding sites in most of the genes. HSFs are transcriptional regulators that are inducible during stress responses, but also during normal development and physiology (47). Similarly, Sp1 regulates transcription of a large number of genes and is involved in a variety of processes such as growth, differentiation, immune responses, and stem cell regulation (48). The Sp1 binding region contains GC-rich motifs and has been identified in the 5′ region of several genes (49), which explains the high frequency of Sp1 binding sites within the investigated loci that are generally CpG-rich promoter regions. Although, the TF binding site analysis performed is based on putative sites, some have been confirmed. For example, Sp1 and CEBP have been confirmed to interact with the LEP promoter (50). Given the technical limitations of sequence-based TF queries, it remains to be determined whether these or other TFs are actively involved in regulation of the genes analyzed in this study, but this could be an intriguing endeavor, especially in light of the tissue-specific expression of TFs (51). The tissue-specific nature of the TFs binding to these sites may also be critical in driving the differences in tissue-specific DNA methylation observed in this study.

Within the loci examined, correlations between DNA methylation of CpG units varied across tissues. Placenta exhibited the highest degree of correlation and saliva the lowest degree. Correspondingly, the loci LINE-1, HSD11B2, CEBPB, NR3C1, and LEP displayed average coefficients >0.2 in 3 of the tissues, whereas other loci, such as DEPTOR, had low correlation between adjacent CpGs. The length of the sequence does not appear to play a role in the degree of correlation. These results highlight the dynamic nature and tissue specificity of methylation marks and suggest that correlation should be assessed and appropriate statistical modeling of the data should be used. This finding, along with the data regarding TF binding, supports, at least in some cases, examining CpG loci individually.

As expected, we observed a higher degree of methylation at the LINE-1 and Alu8b elements, but, consistent with our findings in gene-specific regions, methylation of these sequences differed between tissues. This intertissue variation may be explained by the polymorphic and ubiquitous location of the elements across the genome (52) or tissue-specific differences in the need to control their expression. Given their abundance, these sequences have been the subject of a considerable amount of study, especially in the cancer biomarker field (53). In addition, placental global DNA methylation has been shown to be associated with in utero exposures (54), highlighting the need for early-life epigenetic profiling of these elements (55–57).

Among the gene-specific loci, we observed a great degree of intertissue and interindividual variation. Most notably, placental LEP and HSD1B2 exhibited abundant variation when compared to the other loci examined. These results are consistent with studies by our group (23, 26) and others (58, 59) that have profiled placental methylation and found a negative correlation with gene expression in this tissue (23, 60). This is the only study that has assessed saliva LEP and HSD1B2 methylation in infants; we found lower methylation of LEP in saliva than in the other profiled tissues. The low level may be explained by previous observations of leptin in microglobules in whole saliva and salivary glands (61). Similarly, the lower HSD1B2 methylation in saliva could be related to the expression of this glucocorticoid antagonist in sweat and salivary glands (62). Notably, although there is likely to be greater cellular heterogeneity in blood than in the cells found in saliva, LEP had higher interindividual variation in saliva than in cord blood.

This is the first report of DNA methylation patterns in these newborn/infant tissues within CEBPB, a TF implicated in numerous cellular processes, including myeloid cell differentiation (63). We observed higher promoter methylation in cord blood, a somewhat unexpected finding, as these samples are likely to contain hematopoietic progenitors including myeloid cells (64), although these are probably a minor cell type in the samples. We did not quantify gene expression; hence, we cannot state the functional relevance of this finding in cord blood. More research is needed to explore CEBPB epigenetic variation and potential relation to gene expression, but our preliminary results showed an interesting pattern of intertissue and interindividual variation at this locus.

The promoter of KLF15, a downstream target of NR3C1 (27), and gene body of DEPTOR were also examined. Our results indicate very low methylation levels and low variability in all tissues, demonstrating that at least at the examined CpGs, methylation profiling may not be useful in larger studies. Methylation at the gene body of DDIT4 was higher, and interindividual variation was found at this locus in placental tissue in comparison to that in cord blood or saliva. Although, DDIT4 interacts with NR3C1 in skeletal muscle (27), it is unknown whether this gene is epigenetically regulated or is expressed in the tissues examined.

The glucocorticoid receptor NR3C1 is a known mediator of glucocorticoid signaling. Methylation in the examined area (exon 1F) correlates inversely with expression in placental tissue (25). Although we observed generally low levels of methylation at NR3C1, this promoter further exemplifies the tissue specificity of this epigenetic mark, and even these low levels of methylation have been linked to variability in expression. Saliva methylation was the highest and most variable, and, in this region, it varied inversely with total salivary cortisol in female adults who participated in a social stress test (65). Such findings demonstrate the potential utility of these biomarkers, even
though the intertissue correlation may be low, as these markers can still be representative of key phenotypes.

We observed a positive correlation between cord blood and placental AluYb8 methylation, and similar results were observed for LINE-1, although with limited statistical power. However, saliva global DNA methylation did not correlate with placental or cord blood. We could hypothesize that this discrepancy is due to the extemporaneousness of saliva sample collection, but we did not have saliva collected at birth to test this hypothesis. In addition, we observed a negative correlation between cord blood and saliva NR3CI methylation, but not for any of the other gene loci or repetitive elements. If the observed correlation is confirmed in future studies, it could offer the possibility of prospective saliva sampling and NR3CI DNA methylation profiling in infants and eliminate the need of venipuncture in a susceptible population. Similar correlations between blood and buccal cell methylation have been observed in a study in adults (39), but the NR3CI locus was not investigated, and a correlation for LEP methylation was found that we did not observe in our study. These contrasting results highlight the need for future studies that assess DNA methylation in different tissues at different time points in life, especially with the recognized dynamic nature of the epigenome.

This study has several advantages. We profiled methylation in 5 paired tissues from infants, and we investigated several gene candidates and global DNA methylation by using a highly quantitative technique. However, our sample size was limited, which may have limited our power to detect associations, and we did not take into account the cellular heterogeneity of these tissues that, at least in cord blood, has been shown to influence DNA methylation (18). In addition, we did not have saliva DNA samples collected at birth; thus, we could not assess how temporal changes may affect the correlations between saliva and the other tissues studied. Overall, our findings, in line with the growing literature on epigenetic mechanisms, suggest highly tissue-specific DNA methylation that would not support using these tissues as surrogates for one another at the loci examined. Each tissue’s unique epigenetic signature is likely to reflect its differential functions. At the same time, numerous studies have begun to link both environmental factors and phenotypes with DNA methylation in these various tissues. Such studies strongly support the utility of accessible epigenetic features as biomarkers of both exposure and effect in epidemiologic and clinical/translational studies. It is critical to carefully assess how various tissue epigenetic marks relate to these factors and outcomes and to interpret findings appropriately in light of this specificity (7, 25, 66–72).

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