Original Contribution

Accounting for Life-Course Exposures in Epigenetic Biomarker Association Studies: Early Life Socioeconomic Position, Candidate Gene DNA Methylation, and Adult Cardiometabolic Risk


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Recent studies suggest that epigenetic programming may mediate the relationship between early life environment, including parental socioeconomic position, and adult cardiometabolic health. However, interpreting associations between early environment and adult DNA methylation may be difficult because of time-dependent confounding by life-course exposures. Among 613 adult women (mean age = 32 years) of the Jerusalem Perinatal Study Family Follow-up (2007–2009), we investigated associations between early life socioeconomic position (paternal occupation and parental education) and mean adult DNA methylation at 5 frequently studied cardiometabolic and stress-response genes (ABCA1, INS-IGF2, LEP, HSD11B2, and NR3C1). We used multivariable linear regression and marginal structural models to estimate associations under 2 causal structures for life-course exposures and timing of methylation measurement. We also examined whether methylation was associated with adult cardiometabolic phenotype. Higher maternal education was consistently associated with higher HSD11B2 methylation (e.g., 0.5%-point higher in 9–12 years vs. ≤8 years, 95% confidence interval: 0.1, 0.8). Higher HSD11B2 methylation was also associated with lower adult weight and total and low-density lipoprotein cholesterol. We found that associations with early life socioeconomic position measures were insensitive to different causal assumption; however, exploratory analysis did not find evidence for a mediating role of methylation in socioeconomic position-cardiometabolic risk associations.

epigenetic biomarker; life course; marginal structural models; methylation; time-dependent confounding

Abbreviations: ABCA1, adenosine triphosphate binding cassette subfamily A member 1 gene; BMI, body mass index; HSD11B2, hydroxysteroid (11-β) dehydrogenase 2 gene; INS-IGF2, insulin–insulin-like growth factor 2 readthrough gene region; LEP, leptin gene; NR3C1, nuclear receptor subfamily 3 group C member 1 (“glucocorticoid receptor”) gene.

Early life exposure to stressful environments such as poverty, famine, and war may increase adult susceptibility to mortality (1), cardiovascular disease (2, 3), and poor pregnancy outcomes (4–6). Programming of the fetal epigenome through changes in DNA methylation (7–10) is a putative mechanism. The response of methylation in gene promoter regions to intrauterine environment (11–13) and persistence into adulthood (6–10) may mediate the association of early life exposures with adult health. Importantly, DNA methylation of growth, metabolism, and stress response genes (8, 11–14) may be programmed by prenatal stressors, including those related to socioeconomic position (7, 15, 16).

Several studies have found associations between severe maternal stressors and candidate gene DNA methylation in cardiometabolic and stress response genes in neonates (17), adolescents (18), and adults (12, 19). Early life socioeconomic position as measured by parental education and other factors has also been associated with systemic, differential offspring methylation (20–23) and, recently, with functionally relevant candidate genes (16). However, studies investigating associations...
between parental socioeconomic position and offspring methylation in specific cardiometabolic and stress response genes, such as the nuclear receptor subfamily 3 group C member 1 gene (NR3C1) or the insulin–insulin-like growth factor 2 readthrough gene region (INS-IGF2) (24, 25), may provide evidence for a mechanistic role of DNA methylation in early life socioeconomic position–adult health relationships (26–29). Moreover, investigating associations among women may be of particular interest, because their own early life environment may affect the intrauterine environment for their offspring (6, 26, 27).

However, a major identified gap in the empirical literature is identifying whether observed associations between early life factors and adult DNA methylation may be due to life-course experiences (16, 30), rather than strictly early life programming. For example, adolescent adiposity or obesity may both mediate the early life–adult methylation relationship through developmental programming (30), as well as confound relationships between adult socioeconomic position (due to health selection (31)) or smoking (due to desire for weight loss (32)) and adult methylation. To avoid overcorrecting for such mediators in estimating the direct effects of early life environment, we may find that indirect adjustment for these variables through marginal structural models may be necessary (3).

We used several alternative life-course models and modeling techniques to investigate associations between measures of early life socioeconomic position and DNA methylation in commonly studied cardiometabolic (33) and stress-related (7, 34, 35) genes among young adult women (mean age = 32 years). Properly accounting for complex relationships among life-course variables by using marginal structural models may provide stronger evidence that developmental programming is responsible for early life socioeconomic position–adult methylation associations. Additionally, we investigated adult DNA methylation–adult health associations and explored potential mediation of early life socioeconomic position–adult health associations by DNA methylation.

**METHODS**

**Study setting and population**

Our investigation was conducted in the Jerusalem Perinatal Study that included all 17,003 births to Jerusalem residents between 1974 and 1976 (36, 37). Around the time of birth, maternal medical history, pregnancy course, and offspring birth weight were abstracted from birth certificates or maternity ward logs. Additionally, mothers were interviewed 1 or 2 days postpartum. In the subsequent Jerusalem Perinatal Study Family Follow-up Study, a sample of 1,400 mother–offspring dyads was drawn from the Jerusalem Perinatal Study, oversampled by maternal prepregnancy body mass index (BMI, weight (kg)/height (m)²) ≥27 and offspring birth weight ≤2,500 g or ≥4,000 g, and restricted to singletons born ≥36 weeks without overt congenital malformations. Between 2007 and 2009, offspring (mean age = 32 years) were interviewed by telephone about sociodemographic, behavioral, and health history. In later physical examinations, peripheral blood samples were collected, assayed, and stored (36–38). All female Jerusalem Perinatal Study Family Follow-up Study offspring with stored blood (n = 613) were included in the current study. Jerusalem Perinatal Study Family Follow-up Study protocols were approved by the University of Washington (Seattle, Washington) and Hadassah-Hebrew University Medical Center (Jerusalem, Israel) institutional review boards.

**Data collection**

*Early life socioeconomic position.* We utilized 4 measures for early life socioeconomic position: 1) 6-category paternal occupational class (i.e., 6 = lowest/manual and 1 = highest/professional) as reported during the maternal postpartum interview; 2) binary paternal occupational class (high = 1–3 and low = 4–6); and total years of 3) maternal and 4) paternal education (36–39). The class measure was derived from ranking 108 occupations by mean years of education (37). Representative occupations include teachers (class 1), rabbis (class 2), accountants (class 3), electricians (class 4), skilled agricultural workers (class 5), and unskilled laborers (class 6) (Web Figure 1 available at http://aje.oxfordjournals.org/). This measure has been shown to be associated with a number of outcomes in the Jerusalem Perinatal Study population (37). Maternal occupation was not included, as a majority of women in the study (59%) were not employed outside the home (37). We also categorized parental education as ≤8, 9–12, and ≥13 years.

*Candidate gene selection and methylation profiling.* Five genes were selected on the basis of prior literature and putative cardiometabolic (adenosine triphosphate binding cassette subfamily A member 1 gene (ABCA1), INS-IGF2, and leptin gene (LEP)) and stress response (11β-dehydrogenase 2 gene (HSD11B2) and NR3C1) functions. These genes are related to cholesterol homeostasis (ABCA1) (24, 33), insulin production and growth (INS-IGF2) (24, 25, 33), energy balance (LEP) (33), cortisol inactivation (HSD11B2) (40–42), and cortisol signaling (NR3C1) (17, 18, 43), respectively. Using stored peripheral blood samples obtained at the physical examination, we performed quantitative assessment of methylation by use of the MassARRAY procedure (Sequenom, Inc., San Diego, California) (44, 45) at Roswell Park Cancer Institute’s Genomics Shared Resource (Buffalo, New York). Briefly, 1 μg of genomic DNA was bisulfite converted by using an EZ DNA Methylation Kit (Zymo Research, Orange, California). Converted DNA was then amplified by polymerase chain reaction under standard conditions (44; Web Figure 2) using primers designed in MethPrimer (45) to coincide with prior literature (17, 18, 24, 25, 33, 39–44; Web Figure 2, Web Appendix 1). Matrix-assisted laser desorption/ionization time-of-flight mass spectrometry was conducted to analyze cleavage products, with methylation calls performed by EpiTyper v1.0 (Sequenom) and written to an Oracle 8i database. Each cleavage product corresponding to 1 CpG (cytosine-phosphate-guanine site of methylation) unit, with the output value being proportion methylated among several proximal CpG sites (Web Appendix 1). Region-specific methylation was calculated as the arithmetic mean of all CpG units in the region. Control runs of 0%, 50%, and 100% methylation were included for quality control. CpG units with >25% failed methylation calls were excluded (3 for ABCA1, 2 for LEP, and 4 for NR3C1). No subjects were excluded on the basis of these criteria.

*Adult cardiometabolic and pregnancy outcomes.* At the subject’s physical examination, height, weight, waist and
hip circumference, and systolic and diastolic blood pressures were measured by study staff (36). Additionally, a concomitantly drawn fasting (≥8 hours since last meal) peripheral blood sample was assayed for total cholesterol and lipids on the VITROS 5,1 FS Chemistry System (Ortho Clinical Diagnostics, Raritan, New Jersey), and high- and low-density lipoprotein cholesterol concentrations were calculated (36).

We constructed 3 binary outcomes: obesity (BMI ≥30); metabolic syndrome defined on the basis of International Diabetes Federation criteria (46); and, among women reporting any children (n = 451), whether they had any term, low birth weight offspring (<2,500 g; yes/no). We were interested in the latter because our previous work suggested that grandmaternal education is positively associated with grandchild birth weight (47).

Early life confounders. Data on maternal prepregnancy overweight status (prepregnancy BMI ≥27 based on oversampling criteria), age at birth, any prenatal smoking, and parity from postpartum interview, as well as subject’s birth weight from medical records (36–38), were included. Previous studies have shown maternal prepregnancy BMI, age, ethnicity (39), and prenatal smoking and offspring birth weight (25) to be confounders of early life socioeconomic position-methylation associations. Although all mothers in this study reported Jewish ethnicity, we included whether a mother reported immigrating from the West as a confounder (38). Because birth order (48) and age (9) may be related to methylation, we also adjusted for maternal parity and subject’s age at blood draw.

Life-course measures. Although methylation patterns are established early (8, 9, 21), they may also be influenced by a subject’s life course. We included the following variables from the interview at age 32 years: self-reported adolescent overweight status (yes/no) in grades 4–6 (~10 to 12 years), total years of education, religiosity (secular, traditional, religious, or ultra-Orthodox), marital status, parity, and alcohol or tobacco use. In particular, overweight status may be a time-dependent confounder (i.e., both a consequence of early programming and a confounder of later mediator-outcome relationships) (47). Because recall may be biased, we included an objectively measured BMI at age 17 years available for some participants (n = 374) (37, 38) in sensitivity analyses.

We constructed a traditional family role factor score and dummy for any drinking/smoking (yes/no). The traditional role factor score was estimated by principal components factor analysis of: religiosity, marital status, parity, and years of education. Higher scores corresponded to greater religiosity and parity, being married, and fewer years of education. These characteristics have been shown to be related to each other as well as drinking, smoking, and adiposity among Israeli (49, 50) and non-Israeli women (51). In turn, drinking, smoking, and adiposity may affect adult methylation (23, 30). We dichotomized this score at the median.

Hypothesized causal models. Because variation of DNA methylation over the life course is understudied (30, 52, 53), we must make certain causal structure assumptions to assess associations between early life socioeconomic position and DNA methylation. We used 2 alternate models: Under an “early programming” model, we assumed that methylation is set in early life and does not change in adulthood (Figure 1). Under a “cumulative effects” model, we assumed that some variation in methylation is due to life course (Figure 2).

Statistical analysis

First, we described the study population by using means or proportions and methylation by means and quartiles. To estimate early life socioeconomic position-adult methylation associations, we fit multivariable linear regression models predicting average methylation at each gene region (ABCA1, HSD11B2, INS-IGF2, LEP, NR3C1) by each of the 4 socioeconomic position measures, adjusted for the following:

1. Subject’s age at blood draw and stratification variables (maternal prepregnancy BMI ≥27 and subject’s birth weight ≤2,500, 2,501–3,999, or ≥4,000 g) (“Crude” model);
2. Also all early life confounders ("Early Programming" Model);
3. Also all life-course measures, except adolescent overweight ("Cumulative Effects" Model).

Under "Cumulative Effects," adolescent overweight status is an important biological mediator for early life socioeconomic position, but it may also confound relationships between adult risk factors (such as smoking) and methylation. Consequently,
Table 2. Detailed Methylation Distribution of Candidate Gene Promoter Regions by Paternal Occupational Class, Jerusalem Perinatal Family Follow-up Study, 1974–2009

<table>
<thead>
<tr>
<th>Gene</th>
<th>Chromosome</th>
<th>Location</th>
<th>No. of CpGs</th>
<th>Mean % (SD)</th>
<th>Overall (n = 619)</th>
<th>High Class (n = 296)</th>
<th>Low Class (n = 256)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABCA1</td>
<td>9</td>
<td>16:27,3–117,680,921</td>
<td>27</td>
<td>20.0 (7.3)</td>
<td>19.6 (7.2)</td>
<td>20.7 (7.4)</td>
<td>19.6 (7.2)</td>
</tr>
<tr>
<td>HSD11B2</td>
<td>11</td>
<td>12:172,622,124–123,058,622</td>
<td>11</td>
<td>6.6 (3.1)</td>
<td>6.3 (2.9)</td>
<td>6.3 (2.9)</td>
<td>6.3 (2.9)</td>
</tr>
<tr>
<td>INS-IGF2</td>
<td>7</td>
<td>11:2,182,336–2,182,640</td>
<td>8</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
</tr>
<tr>
<td>LEP</td>
<td>7</td>
<td>11:2,182,336–2,182,640</td>
<td>8</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
</tr>
<tr>
<td>NR3C1</td>
<td>11</td>
<td>17:52,783,056–174,283,905</td>
<td>5</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
<td>6.5 (1.7)</td>
</tr>
</tbody>
</table>

Abbreviations: ABCA1, adenosine triphosphate-binding cassette A1; HSD11B2, hydroxysteroid-11-β dehydrogenase 2 gene; INS-IGF2, insulin insulin-like growth factor 2 readthrough gene region; LEP, leptin gene; NR3C1, nuclear receptor subfamily 3 group C member 1 (glucocorticoid receptor) gene. **Location** refers to Genome Reference Consortium human genome build 37 (GRCh37).

RESULTS

Overall, our subjects (n = 613) did not differ substantially from the total female population of the Jerusalem Perinatal Study Family Follow-up Study (n = 715) (36). High paternal economic position measures and binary life course factors to also fit marginal structural models, indirectly adjusting for adolescent overweight by inverse probability weighting (3, 54), using the outcome model:

\[ E[Y_i | X_i = x, T_i = t, S_i = s] = \beta_0 + \beta_1 x + \beta_2 t + \beta_3 s, \]

where \( Y_i \) is mean methylation, \( X_i \) is early life socioeconomic position category, \( T_i \) is high traditional role score, and \( S_i \) is any drinking/smoking. For each predictor of \( Y_i \), stabilized weights were estimated by fitting multinomial logistic regression models to predict the probability of a given level of exposure by hypothesized causal parents. Each individual was then assigned an overall weight \( (W_i^{\text{overall}}) \) by the following set of equations:

- **Early life SEP:**
  \[ w_i^S = \frac{P(X = x_i)}{P(X = x_i|C = c_i)}. \]

- **High traditional role:**
  \[ w_i^T = \frac{P(T = t_i|C = c_i)}{P(T = t_i|C = c_i, X = x_i, B = b_i, O = o_i)}. \]

- **Any drinking/smoking:**
  \[ w_i^S = \frac{P(S = s_i|C = c_i)}{P(S = s_i|C = c_i, X = x_i, B = b_i, O = o_i, T = t_i)}. \]

Overall weight: \( W_i^{\text{overall}} = w_i^S \times w_i^T \times w_i^S \).

Here, \( C \) is a vector of confounders: maternal Western immigrant status, any prenatal smoking, parity, prepregnancy BMI, and subject’s age at blood draw; \( B \) is subject’s birth weight; and \( O \) is adolescent overweight. By weighting, the functional causal order is \( C, X, B, O, T, S \). We also fit linear regression models using these predictors (i.e., \( X, T, \) and \( S \)), to investigate whether any differences in marginal structural models estimates may be due to different parameterizations. Multiple imputation was used to estimate the influence of missingness (n = 19 with any missing values).

Secondary investigation of methylation-adult health associations

We were also interested in whether observed DNA methylation was related to adult health. However, methylation was measured from the same samples used to assay other biomarkers. Nonetheless, we estimated cross-sectional associations between methylation and adult phenotype by fitting multivariable regression models, adjusting for all confounders, and life-course variables. Continuous biomarkers (e.g., total cholesterol) were normally distributed and left untransformed (36). We also conducted exploratory analyses to see if DNA methylation mediated any indirect effects of early life socioeconomic position on adult phenotypes.

We conducted all data analyses using Stata MP 13.1 software (StataCorp LP, College Station, Texas). Given our a priori socioeconomic position measures and specific gene regions, we performed no multiple testing corrections.
Table 3. Associations Between Early Life Socioeconomic Position and Percent Candidate Gene Methylation, Jerusalem Perinatal Family Follow-up Study, 1974–2009

<table>
<thead>
<tr>
<th>Exposure</th>
<th>ABCA1</th>
<th>HSD11B2</th>
<th>INS-IGF2</th>
<th>LEP</th>
<th>NR3C1</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>β</td>
<td>95% CI</td>
<td>P Value</td>
<td>β</td>
<td>95% CI</td>
</tr>
<tr>
<td></td>
<td>Coefficient</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crude Model(^b)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High paternal occupational class(^c)</td>
<td>0.5</td>
<td>-0.9, 1.9</td>
<td>0.5</td>
<td>-0.04</td>
<td>-0.4, 0.3</td>
</tr>
<tr>
<td>Increasing paternal occupational class(^d)</td>
<td>0.5</td>
<td>0.04, 0.9</td>
<td>0.03</td>
<td>-0.02</td>
<td>-0.1, 0.1</td>
</tr>
<tr>
<td>Mother's years of education</td>
<td>0.2</td>
<td>-0.02, 0.4</td>
<td>0.09</td>
<td>0.07</td>
<td>0.01, 0.1</td>
</tr>
<tr>
<td>Father's years of education</td>
<td>0.1</td>
<td>-0.05, 0.3</td>
<td>0.2</td>
<td>0.03</td>
<td>-0.01, 0.08</td>
</tr>
<tr>
<td>&quot;Early Programming&quot; Model(^e)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High paternal occupational class(^c)</td>
<td>0.4</td>
<td>-1.1, 1.8</td>
<td>0.6</td>
<td>0.0002</td>
<td>-0.4, 0.4</td>
</tr>
<tr>
<td>Increasing paternal occupational class(^d)</td>
<td>0.5</td>
<td>0.004, 0.9</td>
<td>0.05</td>
<td>-0.008</td>
<td>-0.1, 0.1</td>
</tr>
<tr>
<td>Mother's years of education</td>
<td>0.2</td>
<td>-0.003, 0.5</td>
<td>0.05</td>
<td>0.08</td>
<td>0.02, 0.1</td>
</tr>
<tr>
<td>Father's years of education</td>
<td>0.1</td>
<td>-0.1, 0.3</td>
<td>0.2</td>
<td>0.04</td>
<td>-0.003, 0.1</td>
</tr>
</tbody>
</table>

Abbreviations: ABCA1, adenosine triphosphate binding cassette subfamily A member 1 gene; CI, confidence interval; HSD11B2, hydroxysteroid (11-β) dehydrogenase 2 gene; INS-IGF2, insulin–insulin-like growth factor 2 readthrough gene region; LEP, leptin gene; NR3C1, nuclear receptor subfamily 3 group C member 1 ("glucocorticoid receptor") gene.

\(^a\) β coefficients represent percentage-point difference in methylation per unit exposure. Two-tailed P values are given (H\(_0\): β = 0).

\(^b\) Adjusted for maternal prepregnancy overweight (≥27 kg), daughter’s birth weight category (≤2,500 g, 2,501–3,999 g, ≥4,000 g), and daughter’s age at blood draw.

\(^c\) High early life socioeconomic position (father’s occupational classes 1–3) versus low early life socioeconomic position (classes 4–6).

\(^d\) Per unit father’s occupational class increase (e.g., class 5 vs. class 6 or class 1 vs. class 2).

\(^e\) In addition to variables from footnote b, also adjusted for maternal age at birth, any maternal smoking during pregnancy, maternal Western origin, and mother’s parity.
Table 4. Associations Between Early Life Socioeconomic Position and Candidate Gene Methylation by Cumulative Effects Model, Jerusalem Perinatal Family Follow-Up Study, 1974–2009

<table>
<thead>
<tr>
<th>Model</th>
<th>Value Coefficient</th>
<th>95% Cl</th>
<th>P</th>
<th>Value Coefficient</th>
<th>95% Cl</th>
<th>P</th>
<th>Value Coefficient</th>
<th>95% Cl</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABCA1</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
</tr>
<tr>
<td>HSD1B2</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
</tr>
<tr>
<td>INS-IGF2</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
</tr>
<tr>
<td>LEP</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
</tr>
<tr>
<td>NR3C1</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
<td>0.4</td>
<td>−1.19</td>
<td>0.04</td>
</tr>
</tbody>
</table>

Abbreviations: ABCA1, adenosine triphosphate binding cassette subfamily A member 1 gene; CI, confidence interval; HSD1B2, hydroxysteroid (11-) dehydrogenase 2 gene; INS-IGF2, insulin-like growth factor 2 readthrough gene region; LEP, leptin gene; NR3C1, nuclear receptor subfamily 3 group C member 1 ("glucocorticoid receptor") gene. Adjusted for maternal prenatal characteristics (age at delivery, Western origin, any maternal smoking during pregnancy, parity, and prepregnancy overweight), daughter’s birth weight category, and daughter’s life-course mediators as continuous or binary variables. Marginal structural model estimate for the controlled direct effect of higher early life socioeconomic position by each measure (i.e., maternal occupation or maternal educational level) on mean percent methylation. Adolescent overweight was controlled by weighting only, while "high" traditional family role and any alcohol or cigarette use were also included in the outcome model as adjustment variables.

In brief, the table shows associations between early life socioeconomic position and candidate gene methylation. The data are from the Jerusalem Perinatal Family Follow-Up Study, spanning from 1974 to 2009. The table includes models for different genes, with coefficients and confidence intervals provided for each. The coefficients represent the percentage-point difference in methylation per unit of exposure, with two-tailed P-values given (H0: β = 0).
class was associated with more years of parental education, Western-origin mothers, and less maternal smoking (Table 1). Overall, peripheral blood methylation was tightly controlled; interquartile ranges were generally less than 10% (Table 2). Compared with subjects who had low occupational class fathers, those with high class fathers had slightly higher mean methylation at *ABCA1* (20.2% vs. 19.6%), *INS-IGF2* (77.4% vs. 77.2%), *LEP* (22.7% vs. 21.5%), and lower methylation at *NR3C1* (6.4% vs. 6.8%) (Table 2).

In the crude “early programming” (controlling for parental confounders; Figure 1) and “cumulative effects” (also controlling for offspring life course; Figure 2) models estimated by standard linear regression, each higher category of paternal occupational class was associated with 0.5%-points higher *ABCA1* methylation (Tables 3 and 4). However, when marginal structural models were used to account for time-dependent confounding, this association was weaker (Table 6). Notably, this was not due to different covariate parameterizations, as demonstrated by the linear regression using the same predictors (Table 4). Greater maternal education was associated with higher *HSD11B2* methylation in all models (e.g., β = 0.5 per category, 95% confidence interval: 0.1, 0.8) (Table 4). A sensitivity analysis using the 374 individuals with BMI measured at age 17 (37) found even stronger evidence for association (Web Table 1). No evidence of associations was found between socioeconomic position and *LEP*, *INS-IGF2*, or *NR3C1*. Missingness had a minimal influence on any estimates (Web Table 2).

A 1%-point higher *HSD11B2* promoter methylation was associated with lower weight, total cholesterol, and low-density lipoprotein cholesterol as well as 12% greater odds (odds ratio = 1.1, 95% confidence interval: 1.0, 1.3) of having any low birth weight offspring (Web Table 3). Each 1%-point higher *NR3C1* methylation was associated with 0.3 mm Hg higher (95% confidence interval: 0.1, 0.5) diastolic blood pressure (Web Table 3). No significant association was found for *ABCA1*, *LEP*, or *INS-IGF2*.

In exploratory mediation analyses, we used multivariable regression and marginal structural models to estimate the direct and indirect effects of early life socioeconomic position on adult outcomes using our 2 models: 1) adjusted for continuous methylation only, partitioning all indirect effects to the methylation pathway (“early programming”) and 2) adjusted for methylation and life-course measures (“cumulative effects”). We estimated indirect effects using the product of coefficients approach (55, 56) and bootstrapped confidence intervals. We explored whether for methylation, the 10th, 50th, 75th, and 95th percentiles were “functional” thresholds, in the sense that they mediated a relationship between socioeconomic position and adult phenotype. Although we observed an overall association between maternal education on metabolic syndrome and low birth weight (Web Table 4), we did not find any evidence for mediation (Web Tables 4–7).

### DISCUSSION

Overall, we found some evidence for a positive association between early life socioeconomic position and adult peripheral blood DNA methylation at *ABCA1* and *HSD11B2*. We also found associations between *HSD11B2* methylation and several adult cardiometabolic and pregnancy outcomes. Moreover, associations were fairly consistent across models, suggesting that measured life-course trajectory did not substantially explain the relationship between early life and adult methylation and supporting an early programming hypothesis. To our knowledge, this study is the first to examine these associations among young adult women while comparing different life-course causal models.

Although studies of early life environment commonly assess methylation globally (20–22, 57), several have investigated candidate genes. Appleton et al. (39) found an association between maternal ≥ high school attainment and a 9%-point higher (β = 8.8; P < 0.05; n = 444) maternal-side placental *HSD11B2* methylation, adjusted for maternal age, prepregnancy BMI, race, infant sex, and birth weight. Using similar adjustments, we found a positive association between maternal education and adult female *HSD11B2* peripheral blood methylation. Our findings complement several studies extending associations with *HSD11B2* methylation to offspring tissues (40–42), supporting the plausibility of a persistent effect of offspring early life environment. Importantly, reduced *HSD11B2* activity may be related to hypertension (58), and a study of *HSD11B2* methylation in adult peripheral blood found that essential hypertension was associated with a 20% higher promoter methylation (59). Although we found several associations between increased *HSD11B2* methylation and adult phenotype, they did not include blood pressure. However, it is unclear thus far if peripheral blood methylation itself is causal, as expressions in renal and vascular cell walls are more functionally implicated in inflammatory pathways (58). Consequently, it is possible that observed associations with peripheral blood methylation may be noncausal proxies for systemic changes in methylation produced in early development (8–10).

Nonetheless, whether such changes persist into adulthood is a separately important question, as recent investigations have suggested that early differences in peripheral blood DNA methylation related to intrauterine experience may revert to “normal” levels by late childhood (60). Even without a direct functional role, adult candidate gene DNA methylation in peripheral blood can serve as an easily assayed biomarker for early social disadvantage. However, alternative explanations are possible, such as the tendency of offspring to have environments similar to those of their parents. Our study attempted to clarify whether or not differences in adult peripheral blood DNA methylation can be explained by life-course events, or whether they are independently related to early life environment. The concordance between models supports early programming and suggests that life-course events do not completely explain observed associations. Interestingly, although we found consistent associations between *HSD11B2* methylation and maternal education across models accounting for life-course factors, we did not find as strong evidence for an association with occupational class or years of paternal education (Tables 3 and 4). While there are several potential explanations for this, including the role of the mother in child rearing, the findings are at least consistent with an in utero exposure hypothesis (61).

We also found evidence for a positive association between paternal occupational class and adult methylation at *ABCA1* (0.5%-points per level). Tobi et al. (33) found a similar association (0.7%-points, relative to unexposed siblings) between...
early gestation famine exposure and \textit{ABCA1} methylation in older adults (~58 years). Unlike Tobi et al. (33), we did not find associations with \textit{INS-IGF2} or \textit{LEP} methylation. It is possible that famine exposure may influence adult methylation through different pathways than socioeconomic position (12, 19, 33) or that they may be more sensitive to life-course influences (62). Although it has been hypothesized that \textit{ABCA1} may be related to risk of myocardial infarction, our study was in line with that of Talens et al. (24) in finding no substantial association with cardiometabolic phenotype.

Finally, we found little evidence for mediation by methylation. This may be due to several reasons including the aforementioned potential for noncausal methylation differences, the cross-sectional nature of our biomarkers, and the weak relationships between early life socioeconomic position and adult outcomes in our sample. Additionally, a mediating role of methylation in other regions and tissues cannot be excluded (29). Importantly, the exploratory analysis illustrates the difficulty of drawing conclusions about potential mediation strictly from binary associations between methylation and either exposures or outcomes (30).

There are several notable strengths to our study. First, we used state-of-the-art methods to quantify adult methylation at candidate genes previously related to early life conditions. Additionally, our study of 613 women is one of the largest to relate early life environment and candidate gene methylation, enabling us to detect small differences. Finally, our study was conducted among a population with extensive information on maternal demographics and pregnancy course, as well as offspring biomarkers at 32 years (36). Additionally, this population has a unique context for studying intergenerational health. Since 1948, Israel has had large influxes of immigrants (47% of mothers in our study) driven by religious and political pressures and exposed to multiple wars (36, 63). In contrast, it has developed rapidly, with female education averaging 16 years and a gross domestic product per capita exceeding $33,000 (US dollars) (64). Parental adversity may operate intergenerationally through persistent alterations in DNA methylation (22, 27, 28). Consequently, recent improvements in material conditions may mask susceptibility to morbidity and mortality (65). We show that various ways of adjusting for offspring life-course dynamics, including time-dependent confounding by health selection or risk behavior adoption, do little to change associations between early life socioeconomic position and adult methylation.

Several technical limitations to our study deserve mention. First, no replications of polymerase chain reaction amplification were performed (44). Because observed methylation may vary across amplification runs (66), we cannot exclude chance findings or the possibility that this imprecision prevented detection of other relationships. We note again, however, that DNA methylation at our target genes was tightly regulated. The interquartile range for \textit{HSD11B2} methylation was 1.75%-points (4.5%–6.25%) with a majority of the data within 1 standard deviation (2.1%). Additionally, we were unable to quantify cell composition. We attempted application of an adjustment method developed for epigenome-wide association studies (67). Due to the limited number of CpG units (n = 50), however, we found associations between number of maternal years of education and methylation at individual CpG units to be weak and dependent on modeling parameters, also implying that associations may only be at the aggregate level. Consequently, any observed differences in methylation may be mediated by differences in cell compositions of the samples. This is relevant as inflammation, and therefore peripheral cell composition, is likely to be an important mediator of socioeconomic position-cardiometabolic health relationships (16, 68, 69).

Relatedly, we did not conduct any primary analysis of CpG site-specific associations. There are 3 reasons for this: First, we wished to remain consistent with prior candidate gene studies from which we derived primer regions (17, 18, 24, 25, 33, 39–43; Web Appendix 1). Second, we wished to provide a straightforward analysis in the candidate gene setting, which often is an adjunct to exploratory epigenome-wide association studies (60). Third, given the dearth of literature identifying target CpGs, the small number of CpGs available for an agnostic approach would be underpowered to survive necessary multiple testing corrections. As it is, if we had chosen to correct for testing across 4 related socioeconomic position measures, only the marginal structural model-estimated association between maternal education and \textit{HSD11B2} (\(P = 0.008\)) would fall under a Bonferroni \(P = 0.0125\). Overall, these limitations reemphasize the need for replication (both within and between studies) and consideration of other sequencing techniques that may improve detection of small means and differences in candidate gene methylation (43, 66).

Additionally, caution is needed when interpreting and generalizing our findings. Israeli women are required to serve in the military following completion of 12 years of compulsory education. Women may defer service if they are admitted to university or other training opportunities, identify as ultra-Orthodox, or are pregnant. As a result, approximately half of all women serve in the military (36) and differ from those who don’t serve with respect to education, religiosity, and childbearing. Although our models account for these factors, the life-course dynamics of Israeli women may be unique.

In summary, we found some evidence of early life socioeconomic position-candidate gene methylation associations among young adult women. Specifically, we found similar associations using various causal models to account for life-course trajectory, supporting an early programming hypothesis. We also found that methylation of identified genes to be associated with adult cardiometabolic phenotype. However, exploratory mediation analyses were inconclusive, highlighting the challenging nature of epigenetic mediation analyses (30). Nonetheless, we believe that the approach we have described can add to the understanding of life-course methylation dynamics when applied to existing cohorts where blood samples are limited or in follow-up studies to epigenome-wide association studies.

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