Early-Life Socioeconomic Disadvantage and Metabolic Health Disparities

Camelia E. Hostinar, PhD, Kharah M. Ross, PhD, Edith Chen, PhD, and Gregory E. Miller, PhD

ABSTRACT

Objective: A quarter of the world’s population have metabolic syndrome (MetS). MetS prevalence is stratified by socioeconomic status (SES), such that low SES is associated with higher MetS risk. The present study examined the relative roles of early-life SES and current SES in explaining MetS risk.

Methods: Participants (N = 354; ages = 15–55 years, M[SD] = 36.5[10.7] years; 55% female; 72.9% white, 16.9% Asian, 10.2% others) were evaluated for SES and MetS. All were in good health, defined as free of chronic medical illness and acute infectious disease. Using occupational status as a proxy for SES, we recruited roughly equal numbers of participants with low-low, low-high, high-low, and high-high combinations of early-life and current SES. We used the International Diabetes Federation definition for MetS using race- and sex-specific cutoffs for waist circumference, triglyceride levels, high-density lipoprotein cholesterol, blood pressure, and glycosylated hemoglobin levels.

Results: Analyses revealed a main effect of low early-life SES on increased MetS risk according to the three separate definitions. They included the traditional MetS diagnosis (odds ratio [OR] = 1.53, confidence interval [CI] = 1.01–2.33, p = .044), the number of MetS components for which diagnostic thresholds were met (OR = 1.61, CI = 1.10–2.38, p = .015), and a continuous indicator of metabolic risk based on factor analysis (F(1,350) = 6.71, p = .010, partial η² = .019). There was also a significant interaction of early-life SES and current SES in predicting MetS diagnosis (OR = 1.54, CI = 1.02–2.34). The main effects of current SES were nonsignificant in all analyses.

Conclusions: These findings suggest that MetS health disparities originate in childhood, which may be an opportune period for interventions.

Key words: early-life adversity, metabolic syndrome, social mobility, socioeconomic status.

INTRODUCTION

Approximately a quarter of the world’s population have metabolic syndrome (MetS) (1–3), a clustering of abdominal obesity, insulin resistance, dyslipidemia, and elevated blood pressure (1,4). MetS is associated with a two-fold increased risk of cardiovascular events or death (5) and a five-fold increased risk of incident type 2 diabetes mellitus (1). With MetS prevalence rates rising globally (6,7), it is increasingly important to understand its distribution and pathogenesis. Better insights around these issues could improve risk stratification and identify intervention targets.

Most research on the pathogenesis of MetS has focused on interrogating proximal causal factors such as energy imbalance, adiposity, insulin resistance, and physical inactivity (3,4). Much less is known about the distal social and economic contexts that shape these proximal causes across the lifespan. For instance, one observation that has been replicated in several countries is that the prevalence of MetS patterns by concurrent adult socioeconomic status (SES) measured through several indices (e.g., educational attainment, income, subjective social status), such that individuals with lower SES have a higher likelihood of diagnosis (1,8), above and beyond the effects of race/ethnicity and life-style factors (2,8–10).

The stratification of MetS risk by current SES has been noted not only in adults but also in children and adolescents (11,12). These socioeconomic disparities spark a number of questions that have been so far largely overlooked in studying the association between low SES and risk of MetS. For instance, is the increased MetS risk in low SES populations a result of exposures in adulthood? Or, because low SES is relatively stable across the life course, is it the accumulation of living in disadvantaged environments from childhood onward? Does the timing of exposure to low SES (i.e., childhood versus adulthood) matter in explaining the risk of MetS? And for those who experience socioeconomic mobility—either upward or downward—does the risk of MetS change accordingly?

Epidemiological studies are revealing that the global prevalence of MetS is not only rising in adults but also steadily increasing in children and adolescents worldwide (13). Furthermore, children and adolescents who experience socioeconomic disadvantage...
and other psychosocial adversities are at an increased risk of metabolic dysfunction later in life (14–19). These findings raise questions about the developmental origins of adult MetS and challenge us to find ways to understand when demographic and psychosocial risk factors for MetS might begin to influence health. Early-life conditions are thought to shape the risk of obesity and MetS in adulthood through a number of pathways (13,15,20). Chronically stressful environments during fetal life and childhood are associated with low birth weight, altered childhood growth patterns, and earlier pubertal onset (20) and with endocrine, autonomic, and metabolic patterns that promote visceral adiposity (13,20). In addition, proinflammatory signaling (21) and altered threat, reward, and executive control processes (22) that shape health behaviors such as food intake and physical exercise likely further exacerbate metabolic risk. Together, all these sources of influence are thought to contribute to enhancing metabolic risk in adulthood.

However, what is not well understood is whether the MetS risk conferred by early-life conditions in general and low early-life SES in particular is independent of adult experiences. This is important to consider, given that most poor children grow up to be poor adults (23). Empirically, the association between childhood SES and risk of type 2 diabetes and obesity in adulthood is attenuated when adjusting for adult SES (15), but there is limited evidence on this issue available, and the results are inconclusive. Indeed, in studies where childhood SES and adult SES are modeled together, the findings are mixed. Some studies observe that childhood but not adult SES predicts metabolic risk (16,17,24,25). Others find the reverse (26) or independent roles for both childhood and adult SES (27–29). This question is inherently difficult to address in the general population because childhood and adult SES are typically correlated because of a high degree of continuity in SES across the lifespan for many individuals (15). The innovative aspect of the present study is that it explicitly addressed this challenge by disentangling the roles of early-life and current SES by study design.

Studying upward and downward social mobility can also shed light on the independent contributions of early-life and adult SES. There is very limited research on associations between social mobility and MetS risk, but emerging patterns suggest that low childhood SES is associated with greater odds of MetS in later life, and that upward social mobility does not offset this risk (28,30,31). These results contrast with findings showing similar risk of diabetes for those with low-high combinations of childhood and adult SES compared with individuals with continuously high SES (32), suggesting that more work is needed to clarify these patterns. Proposing clearer life-course models of links between SES and MetS could inform interventions regarding the ideal developmental stages when intervening would have maximal health benefits.

The Present Study

The present study aimed to examine the independent and interactive roles of early-life and current adult SES in explaining variability in adult metabolic risk, indexed by the three following types of outcomes: (a) diagnosis of MetS according to the criteria specified by the International Diabetes Federation (IDF) definition, (b) a count of the number of MetS components on which the participant met the IDF criteria, and (c) a continuous factor score indexing the common variance of the following five standardized MetS components: waist circumference, blood pressure, triglycerides, high-density lipoprotein (HDL) (reversed), and glycosylated hemoglobin (HbA1c). The rationale for using several indices of metabolic risk is that even though IDF’s MetS diagnosis is clinically useful and has generalizability by providing information that is comparable globally (4), there is also some recognition in the literature that continuing to examine the individual risk factors embedded in MetS and the way they cluster together is a useful avenue for understanding subclinical disease processes and their etiology (33,34). Furthermore, there has been some debate about whether MetS is best viewed and analyzed as a dichotomous diagnosis or a spectrum of risk, including continuous numeric components for each of the risk factors (34). For this reason, the present study examined not only a dichotomous indicator of MetS diagnosis according to the IDF criteria but also sought to descriptively characterize the prevalence of individual risk factors and to additionally examine metabolic risk across a continuum, captured in two ways (count of components and factor scores), as described in more depth hereinafter.

METHODS

Participants

The study recruited 360 participants from Vancouver, British Columbia, Canada, through postings in local media and public transit. Recruitment and testing were conducted between February 2009 and May 2012. Only six participants had missing data for MetS components; thus, primary analyses were conducted on 354 participants. These 354 participants were between the ages of 15 and 55 years (M[SD] = 36.5[10.7] years, 55% female) and recruited to fit into one of the four groups defined by early-life (low versus high) and adulthood (low versus high) SES (see operational definitions under Measures and participant characteristics in Table 1). To minimize confounding by health status, participants had to be (a) free of infectious disease in the 2 weeks before testing, as evidenced by self-report and a normal complete blood count, and (b) without a history of serious and chronic medical illnesses including, but not limited to, cancer, diabetes mellitus, heart disease, stroke, autoimmune disease, human immunodeficiency virus/acquired immunodeficiency syndrome, hepatitis, chronic obstructive pulmonary disorder, asthma, schizophrenia, bipolar disorder, and dementia. The participants who presented with acute infections were rescheduled after the signs had resolved. The candidates were also screened out if they were not fluent in English and if they were pregnant or had been pregnant in the previous year.

Of the 2880 individuals who responded to the study advertisements, 417 met all of the eligibility criteria described previously and volunteered to participate after learning about the study details. Three hundred sixty subsequently attended their scheduled laboratory session. The project was approved by the University of British Columbia’s research ethics board, and all participants gave written informed consent.

Procedure

All participants completed laboratory sessions between 8:00 AM and 10:00 AM after an overnight fasting period. After the study details were explained, the participants gave written consent and had blood drawn by antecubital venipuncture to measure MetS components (see hereinafter). Later in the session, the participants completed a battery of self-report measures and behavioral tasks and had their height, weight, and waist circumference measured (see details in Measures).
TABLE 1. Characteristics of the Sample (N = 354)

<table>
<thead>
<tr>
<th>Life-Course SES Trajectory</th>
<th>Low-Low SES (n = 97)</th>
<th>Low-High SES (n = 93)</th>
<th>High-Low SES (n = 72)</th>
<th>High-High SES (n = 92)</th>
<th>Differs by Early SES</th>
<th>Differs by Current SES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y</td>
<td>35.5 (10.7)</td>
<td>39.5 (9.5)</td>
<td>34.9 (10.6)</td>
<td>35.6 (11.5)</td>
<td>.054</td>
<td>.04</td>
</tr>
<tr>
<td>Sex, female, n (%)</td>
<td>47 (48.5)</td>
<td>55 (59.1)</td>
<td>42 (58.3)</td>
<td>50 (54.3)</td>
<td>.65</td>
<td>.44</td>
</tr>
<tr>
<td>Descent, European, n (%)</td>
<td>65 (67)</td>
<td>59 (63.4)</td>
<td>59 (81.9)</td>
<td>75 (81.5)</td>
<td>&lt;.001</td>
<td>.84</td>
</tr>
<tr>
<td>Parental education (household max), y</td>
<td>11.8 (2.7)</td>
<td>12.4 (2.6)</td>
<td>15.4 (2.1)</td>
<td>16.1 (2.6)</td>
<td>&lt;.001</td>
<td>.01</td>
</tr>
<tr>
<td>Current education (household max), y</td>
<td>15.1 (2.8)</td>
<td>15.7 (2.5)</td>
<td>14.1 (2.3)</td>
<td>16.6 (2.5)</td>
<td>.03</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Current annual household income</td>
<td>$25–$34.9 K</td>
<td>$50–$74.9 K</td>
<td>$25–$34.9 K</td>
<td>$50–$74.9 K</td>
<td>.39</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Waist circumference, cm</td>
<td>89.7 (16.3)</td>
<td>86.9 (15.9)</td>
<td>84.4 (13.3)</td>
<td>83.4 (12.6)</td>
<td>.005</td>
<td>.17</td>
</tr>
<tr>
<td>Systolic blood pressure</td>
<td>108.9 (13.3)</td>
<td>107.8 (11.7)</td>
<td>105.02 (11.2)</td>
<td>108.1 (10.8)</td>
<td>.21</td>
<td>.60</td>
</tr>
<tr>
<td>Diastolic blood pressure</td>
<td>69.0 (10.8)</td>
<td>70 (9.5)</td>
<td>67.53 (9.6)</td>
<td>69.7 (7.9)</td>
<td>.44</td>
<td>.16</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>1.2 (.74)</td>
<td>1.1 (.60)</td>
<td>.97 (.51)</td>
<td>1.01 (.59)</td>
<td>.073</td>
<td>.61</td>
</tr>
<tr>
<td>HDL cholesterol</td>
<td>1.4 (.34)</td>
<td>1.5 (.39)</td>
<td>1.46 (.37)</td>
<td>1.5 (.42)</td>
<td>.32</td>
<td>.085</td>
</tr>
<tr>
<td>HbA1c</td>
<td>5.4 (.39)</td>
<td>5.4 (.39)</td>
<td>5.33 (.29)</td>
<td>5.3 (.33)</td>
<td>.26</td>
<td>.79</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>26.7 (6.3)</td>
<td>26.6 (6.8)</td>
<td>25.4 (5.8)</td>
<td>24.4 (3.9)</td>
<td>.003</td>
<td>.28</td>
</tr>
<tr>
<td>Heavy smoker, ≥10 cigarettes/d, n (%)</td>
<td>14 (15.9)</td>
<td>2 (2.3)</td>
<td>10 (14.5)</td>
<td>3 (3.3)</td>
<td>.75</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Alcohol user, ≥10 drinks/wk, n (%)</td>
<td>9 (10.2)</td>
<td>10 (11.2)</td>
<td>12 (17.4)</td>
<td>16 (17.6)</td>
<td>.07</td>
<td>.78</td>
</tr>
<tr>
<td>Physical activity, h/wk</td>
<td>2.5 (3.0)</td>
<td>2.6 (2.4)</td>
<td>3.1 (2.9)</td>
<td>2.99 (2.9)</td>
<td>.11</td>
<td>.97</td>
</tr>
</tbody>
</table>

SES = socioeconomic status; HDL = high-density lipoprotein; HbA1c = glycosylated hemoglobin; BMI = body mass index.

Data are presented as M (SD), unless otherwise indicated.

* The study included four groups of participants with low-low, low-high, high-low, and high-high combinations of early-life and current SES.

**Measures**

**Socioeconomic Status**

Participants were recruited on the basis of their early-life and current SES, as defined by the occupational status ratings derived from the United Kingdom’s National Statistics Socio-economic Classification. This system is used widely in epidemiology and has been updated regularly since 1911, allowing for comparability with previous research. It is also well suited to the Canadian social structure. The National Statistics Socioeconomic Classification system codes occupational status using industry, occupation title, occupation description, self-employment status, supervisor status, and number of employees in the company/organization. It also provides specific procedures in case one piece of information among these is unavailable. All recruiters received extensive training before using this system. Coding consistency was regularly checked and discrepancies were resolved by consensus. If applicants did not clearly fall into one of the designated categories, they were not invited to participate in the study. Candidates whose life-course SES fell into one of the four categories, as defined by early-life and current circumstances, were enrolled in the study. The categories were low early/low current (n = 97), low early/high current (n = 93), high early/low current (n = 72), and high early/high current SES (n = 92). The participants were recruited into the four groups on the basis of occupational status because it is often a more visible aspect of SES than educational attainment and because people can more reliably recall their parents’ occupations than income or education. Occupations were graded on an eight-point scale, which was reduced into the three superordinate categories of low, middle, and high SES. Volunteers from either low or high SES categories for both childhood and current SES were enrolled, whereas those from middle categories were ineligible (e.g., small employers and own account workers, lower supervisory, and technical occupations). Low SES included routine and manual occupations, those who never worked and the long-term unemployed. This included positions such as cleaners, laborers, and transportation operatives. High SES included higher managerial and professional occupations, such as architects, engineers, and medical practitioners. These definitions were used to categorize both early-life and current SES. For early-life SES, we coded parental occupational status during their first 5 years of life, using the higher of mother’s versus father’s ratings. To classify the current SES of prospective participants, we coded their occupational status for the past 5 years, as well as that of their romantic partner (the higher of the two ratings was used). A small minority of the participants in our study were in the ages of 15 to 23 years (10.5%) and were full-time students, financially supported by their families. For these cases, we used parental occupation to categorize current SES, unless the participants were financially independent. However, results did not differ when including or excluding this age group.

**Metabolic Outcomes**

A trained phlebotomist collected the overnight fasting blood samples via antecubital venipuncture into serum separator and EDTA-treated tubes (Becton-Dickinson, Oakville, Ontario, Canada). The serum separator tubes were left standing for 60 minutes before being centrifuged at 1000g for 10 minutes, as per manufacturer's instructions. After the serum was harvested, it was frozen at −30°C until assayed in batch at the St. Paul’s Hospital Clinical Trials Laboratory, Vancouver, Canada. Fasting HDL was assayed in serum. Standard enzymatic techniques using cholesterol esterase and cholesterol oxidase were used after low density, intermediate density, and very low density lipoproteins had been precipitated through centrifugation. Assays were performed on a Hitachi 911 instrument (Kyowa Medex, Japan). The interassay coefficient of variation was 5.1%, and the lower limit of detection was 0.189 mmol/L. Glycosylated hemoglobin was measured in whole blood from the EDTA-treated tubes, at the same laboratory, via a Bio-Rad D-10 ion exchange chromatography method (Bio-Rad Laboratories Inc, Hercules, Calif). Assays yield the percentage of hemoglobin that is glycosylated out of total hemoglobin, with greater values indicating higher blood glucose exposure for the past 3 months. The lower range of detection was 1.2%, with an interassay coefficient of variation of 1.2%. Triglycerides were determined enzymatically.
physical activity was measured with the well-validated Paffenbarger Activity Scale (40), which estimates weekly hours of brisk physical activity.

Data Analysis

Data Preparation

Variables were examined for outliers and their approximation of the normal distribution before analyses. Values that exceeded four standard deviations from the mean were Winsorized and replaced with the value at the 99.9th percentile (waist circumference, $n = 3$; triglyceride levels, $n = 4$; HbA1c, $n = 1$; diastolic blood pressure, $n = 1$). Most variables approximated the normal distribution. In contrast, variables indexing the count of MetS components, daily cigarette smoking, alcohol use, and weekly physical activity had a pronounced right skew. MetS components count, cigarette use, and alcohol consumption variables were too skewed to be corrected with mathematical transformations and were, therefore, converted into ordinal scales. For smoking, the new variable was coded as 0 for nonsmoker, 1 for less than 10 daily cigarettes, and 2 for 10 or more cigarettes per day. For alcohol use, it was 0 for zero drinks per week, 1 for less than 10 drinks per week, and 2 for 10 or more drinks per week. For MetS component counts, meeting criteria for three or more symptoms was coded as 3, and ordinal regression was used instead of linear regression in analyses that had this measure as an outcome. For the Paffenbarger index of physical activity, a square-root transformation of the total number of hours of brisk physical activity per week was sufficient to reduce skewness and kurtosis and normalize the distribution; thus, the square-root of the variable was used in analyses.

Missing Data

Only 1.7% ($n = 6$) of the participants were missing data in analyses without covariate adjustment and 8.89% ($n = 32$) in covariate-adjusted analyses. Thus, data imputation was not necessary given that estimates are unlikely to be biased when the rate of missingness is less than 10% (41).

Statistical Analyses

To examine SES disparities in the prevalence of MetS diagnoses, we conducted logistic regression analyses, predicting the binary diagnosis outcome. Effect coding was used for estimating early-life and current SES main effects and their interaction in initial analyses. To probe interactions and test social mobility effects, follow-up logistic regressions using dummy coding were used to examine the differences between the four SES subgroups (low-low, low-high, high-low, and high-high). Then, we used ordinal regression analyses to test the effects of early-life SES, current SES, and their interaction on the count of MetS components, with follow-up ordinal regression analyses used to compare the low-low, low-high, high-low, and high-high groups for probing social mobility effects. Finally, analyses of covariance were used to test the main effects of early-life and current SES and their interaction on MetS factor scores. To examine the role of social mobility, analyses of covariance examining the main effect of SES group (low-low, low-high, high-low, or high-high groups) were also tested in these analyses. If the main effect of SES was significant, the analysis was followed up by the comparisons of marginal means for the low-low versus low-high SES groups (corresponding to the effect of upward social mobility) and the comparisons of the high-high versus high-low groups (corresponding to downward social mobility). All results are presented with and without adjustment for covariates (age, sex, race/ethnicity, alcohol consumption, cigarette smoking, and physical activity levels). Age did not moderate the effects of early-life SES, current SES, or their interaction in any of the analyses (all $p > .22$); thus, these interaction terms were not included in the findings reported here.

RESULTS

Table 1 presents the sample characteristics. As can be seen, the recruitment strategy resulted in the expected differences in parental...
education, current education, and current income based on early-life SES and current SES. The participants with high current SES were also somewhat older and less likely to be heavy cigarette smokers than those with low current SES; thus, we statistically adjusted for these covariates in all analyses while also presenting unadjusted results. The participants who experienced high early-life SES were more likely to be of European descent than those with low early-life SES, thus again, we also used this measure as a covariate for each of the questions we addressed. Interestingly, in these preliminary analyses, there was a main effect of early-life SES on waist circumference and body mass index, with higher values for those with low early-life SES, but no main effect of current SES (Table 1).

Descriptive statistics revealed that metabolic risk disparities by SES were starker when considering early-life SES versus current SES. As Figure 1 shows, the participants with low early-life SES were more likely to meet the IDF criteria for central adiposity ($\chi^2(1) = 5.3, p = .021$), blood pressure ($\chi^2(1) = 7.58, p = .006$), triglycerides ($\chi^2(1) = 2.58, p = .11$), and HbA1c ($\chi^2(1) = 3.28, p = .070$) compared with the participants with high early-life SES. These groups had similar rates of low HDL ($\chi^2(1) = 0.006, p = .94$). By contrast, none of the MetS components varied significantly according to current SES ($\chi^2(1) > 1.22, p > .27$), although the pattern of values favored those with higher current SES. The bivariate correlations among early-life SES, current SES, and each of the MetS summary measures or individual components (Table S1, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A375) also revealed stronger associations of these MetS risk indices with early-life SES compared with current SES.

Using logistic regression, we next examined whether the prevalence of MetS diagnoses varied according to early-life SES, current SES, and/or their interaction. Table 2 presents the results of these analyses, in both crude and covariate-adjusted models. In both cases, there was a significant main effect of early-life SES ($p = .015$), such that the participants from low SES backgrounds had 1.83 greater odds (confidence interval [CI] = 1.12–2.96) of meeting the IDF criteria for MetS diagnosis compared with those with high SES backgrounds. In both models, the main effect of current SES was nonsignificant (odds ratio [OR] = 0.96, CI = 0.59–1.57), and there was a significant early by current SES interaction (OR = 1.78, CI = 1.1–2.87). To decipher the interaction, we conducted follow-up logistic regression analyses using dummy coding for the four SES subgroups (low-low, low-high, high-low, and high-high combinations of early-life and current SES). When using the low-low SES group as the reference, analyses revealed that the other three groups all had significantly lower odds of meeting the diagnostic criteria (low-high group: OR = 0.34, CI = 0.13–0.94; high-low group: OR = 0.10, CI = 0.02–0.45; and high-high group: OR = 0.33, CI = 0.12–0.89). When the high-high group was used as the reference, MetS prevalence rates did not differ statistically from the low-high and high-low groups (OR = 1.05, CI = 0.34–3.26 and OR = 0.29, CI = 0.06–1.54, respectively). Together, these analyses suggest that consistent exposure to low SES across the life course is associated with the highest likelihood of adult MetS, and upward mobility may offset the risk of MetS diagnosis (Fig. 2).

The next sets of analyses explored metabolic risk using ordinal and continuous outcomes. We started by defining risk as a simple count of the number of MetS components each participant had.

**FIGURE 1.** MetS components prevalence by early-life SES and current SES. Color image is available only in online version (www.psychosomaticmedicine.org).
Table 3 presents the results of these analyses in both crude- and covariate-adjusted ordinal regression models. In the unadjusted model, there was a significant main effect of early-life SES (OR = 1.61, CI = 1.10–2.38), such that the participants from low SES backgrounds exhibited a higher number of MetS components (on average, M[SE] = 1.15 [0.08]) compared with the participants from high SES backgrounds (on average, M[SE] = 0.87 [0.07]) (Fig. S1, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A375, for prevalence). As with the binary diagnostic outcome, the main effect of current SES was nonsignificant (OR = 1.31, CI = 0.89–1.92), and in these models, so was the interaction (OR = 0.78, CI = 0.53–1.14). The same pattern emerged in the adjusted model, where there was a main effect of early-life SES (OR = 1.51, CI = 0.99–2.29) but not current SES (OR = 1.22, CI = 0.80–1.85) or an interaction (OR = 0.74, CI = 0.50–1.12). Given our interest in social mobility, we conducted a follow-up ordinal regression comparing the four SES groups using dummy coding. When using the low-low SES group as the reference, analyses revealed that the high-low and high-high SES groups had significantly lower counts of MetS components (high-low group: OR = 0.49, CI = 0.27–0.90; high-high group: OR = 0.54, CI = 0.31–0.96), whereas the low-high group had marginally (p = .091) fewer symptom counts than the low-low group (OR = 0.61, CI = 0.35–1.08), suggesting a trend-level beneficial effect of upward social mobility. When the high-high group was used as the reference, the count of MetS components in this group did not differ statistically from that in the low-high and high-low groups (OR = 1.12, CI = 0.64–1.96 and OR = 0.91, CI = 0.50–1.65, respectively)—that is, there was no evidence for the effects of downward social mobility.

Next, we used factor analysis to derive a continuous metabolic risk score for all participants on the basis of the severity of their signs. These factor scores reflected the common variance of the five continuous metabolic measures (see details in Methods). As Table 4 shows, the pattern of results mirrored those for MetS components count. There was a significant main effect of early-life SES (p = .010) in the crude model, such that the participants from

---

**TABLE 2. Logistic Regression Results Predicting MetS Diagnosis (Binary Outcome)**

<table>
<thead>
<tr>
<th>Model</th>
<th>B</th>
<th>SE</th>
<th>Wald</th>
<th>p</th>
<th>Exp (B)</th>
<th>95% CI (Lower–Upper)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model 1 (N = 354)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant</td>
<td>−2.29</td>
<td>0.19</td>
<td>138.63</td>
<td>&lt;.001*</td>
<td>0.10</td>
<td></td>
</tr>
<tr>
<td>Early-life SES</td>
<td>0.45</td>
<td>0.19</td>
<td>5.27</td>
<td>.022*</td>
<td>1.56</td>
<td></td>
</tr>
<tr>
<td>Current SES</td>
<td>0.30</td>
<td>0.18</td>
<td>2.81</td>
<td>.094</td>
<td>1.35</td>
<td></td>
</tr>
<tr>
<td>Model 2 (N = 354)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant</td>
<td>−2.20</td>
<td>0.18</td>
<td>149.59</td>
<td>&lt;.001*</td>
<td>0.11</td>
<td></td>
</tr>
<tr>
<td>Early-life SES</td>
<td>0.43</td>
<td>0.21</td>
<td>4.06</td>
<td>.044*</td>
<td>1.53</td>
<td></td>
</tr>
<tr>
<td>Current SES</td>
<td>0.11</td>
<td>0.21</td>
<td>0.29</td>
<td>.59</td>
<td>1.12</td>
<td></td>
</tr>
<tr>
<td>Early-life SES by current SES</td>
<td>0.43</td>
<td>0.21</td>
<td>4.17</td>
<td>.041*</td>
<td>1.54</td>
<td></td>
</tr>
<tr>
<td>Model 3 (N = 354)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant</td>
<td>−2.39</td>
<td>0.21</td>
<td>126.33</td>
<td>&lt;.001*</td>
<td>0.09</td>
<td></td>
</tr>
<tr>
<td>Early-life SES</td>
<td>0.43</td>
<td>0.21</td>
<td>4.26</td>
<td>.041*</td>
<td>1.53</td>
<td></td>
</tr>
<tr>
<td>Current SES</td>
<td>0.11</td>
<td>0.21</td>
<td>0.29</td>
<td>.59</td>
<td>1.12</td>
<td></td>
</tr>
<tr>
<td>Early-life SES by current SES</td>
<td>0.43</td>
<td>0.21</td>
<td>4.17</td>
<td>.041*</td>
<td>1.54</td>
<td></td>
</tr>
<tr>
<td>Model 4 (N = 328)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant</td>
<td>−4.79</td>
<td>1.05</td>
<td>2.87</td>
<td>.001*</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>0.05</td>
<td>0.02</td>
<td>6.72</td>
<td>.010*</td>
<td>1.05</td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td>0.13</td>
<td>0.04</td>
<td>0.11</td>
<td>.74</td>
<td>1.14</td>
<td></td>
</tr>
<tr>
<td>Race/ethnicity</td>
<td>0.56</td>
<td>0.51</td>
<td>1.21</td>
<td>.27</td>
<td>1.76</td>
<td></td>
</tr>
<tr>
<td>Alcohol use</td>
<td>0.00</td>
<td>0.28</td>
<td>0.00</td>
<td>1.00</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>Cigarette smoking</td>
<td>0.51</td>
<td>0.27</td>
<td>3.61</td>
<td>.058</td>
<td>1.66</td>
<td></td>
</tr>
<tr>
<td>Physical activity</td>
<td>−0.04</td>
<td>0.03</td>
<td>1.95</td>
<td>.16</td>
<td>0.96</td>
<td></td>
</tr>
<tr>
<td>Early-life SES</td>
<td>0.60</td>
<td>0.25</td>
<td>5.93</td>
<td>.015*</td>
<td>1.83</td>
<td></td>
</tr>
<tr>
<td>Current SES</td>
<td>−0.04</td>
<td>0.25</td>
<td>0.03</td>
<td>.87</td>
<td>0.96</td>
<td></td>
</tr>
<tr>
<td>Early-life SES by current SES</td>
<td>0.58</td>
<td>0.24</td>
<td>5.58</td>
<td>.018*</td>
<td>1.78</td>
<td></td>
</tr>
</tbody>
</table>

SE = standard error; CI = confidence interval for exp (B); SES = socioeconomic status.

Early-life SES and current SES are effect coded such that coefficients represent the effect of being low SES.

* p < .05.
low SES backgrounds had higher composite scores \(M \pm SE = 0.11 \pm 0.06\) than the participants from high SES backgrounds \(M \pm SE = -0.13 \pm 0.07\). This difference became marginal \(p = .063\) with the introduction of covariates (see Fig. 3 for estimated marginal means in this adjusted analysis). Neither the current SES main effect nor the interaction was significant \(F(1,350) = 0.37, p = .54\) and \(F(1,350) = 1.03, p = .31\), respectively. With respect to social mobility effects, the main effect of SES in a one-way analysis of variance was statistically significant in unadjusted analyses \(F(3,350) = 2.74, p = .043\) but was no longer significant with covariate adjustment \(F(3,318) = 1.34, p = .26\); thus, we did not probe pairwise comparisons between the four SES groups any further.

**DISCUSSION**

The prevalence rates of MetS have been rising globally (6), and its distribution and pathogenesis need to be better understood (1,42). Epidemiological studies are increasingly showing socioeconomic stratification of MetS risk (1), which may in turn contribute to explaining social gradients in coronary heart disease morbidity and mortality (10,43). The present study aimed to disentangle the roles of early-life and current SES in explaining MetS risk in a sample of healthy Canadian adults. Secondly, we aimed to test whether there was any evidence consistent with the effects of upward or downward social mobility on MetS risk.

We found a consistent pattern across all the following three metabolic outcomes used: diagnosis according to the IDF criteria, count of MetS components present, and continuous factor scores. In each case, there was a main effect of early-life SES, indicating greater metabolic risk among participants from disadvantageous backgrounds and a nonsignificant main effect of current SES; in addition, there was a significant interaction of early-life SES and current SES in predicting MetS diagnosis. On average, the participants with low childhood SES had 1.83 greater odds of meeting the IDF criteria for MetS.

**TABLE 3. Results of Ordinal Regression Predicting Count of MetS Components**

<table>
<thead>
<tr>
<th>Model 1 (N = 354)</th>
<th>B</th>
<th>SE</th>
<th>Wald</th>
<th>p</th>
<th>Exp (B)</th>
<th>95% CI (Lower–Upper)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Early-life SES</td>
<td>0.48</td>
<td>0.20</td>
<td>5.88</td>
<td>.015*</td>
<td>1.61</td>
<td>1.10–2.36</td>
</tr>
<tr>
<td>Current SES</td>
<td>0.31</td>
<td>0.19</td>
<td>2.48</td>
<td>.12</td>
<td>1.36</td>
<td>0.93–1.99</td>
</tr>
<tr>
<td>Model 2 (N = 354)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early-life SES</td>
<td>0.48</td>
<td>0.20</td>
<td>5.90</td>
<td>.015*</td>
<td>1.61</td>
<td>1.10–2.38</td>
</tr>
<tr>
<td>Current SES</td>
<td>0.27</td>
<td>0.20</td>
<td>1.90</td>
<td>.17</td>
<td>1.31</td>
<td>0.89–1.93</td>
</tr>
<tr>
<td>Early-life SES by current SES</td>
<td>-0.25</td>
<td>0.20</td>
<td>1.63</td>
<td>.20</td>
<td>0.78</td>
<td>0.53–1.14</td>
</tr>
<tr>
<td>Model 3 (N = 354)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>0.03</td>
<td>0.01</td>
<td>11.29</td>
<td>.001</td>
<td>1.04</td>
<td>1.01–1.06</td>
</tr>
<tr>
<td>Sex</td>
<td>0.16</td>
<td>0.21</td>
<td>0.57</td>
<td>.45</td>
<td>1.17</td>
<td>0.78–1.76</td>
</tr>
<tr>
<td>Race/ethnicity</td>
<td>0.04</td>
<td>0.25</td>
<td>0.03</td>
<td>.87</td>
<td>1.04</td>
<td>0.64–1.69</td>
</tr>
<tr>
<td>Alcohol use</td>
<td>-0.25</td>
<td>0.15</td>
<td>2.74</td>
<td>.098</td>
<td>0.78</td>
<td>0.58–1.05</td>
</tr>
<tr>
<td>Cigarette smoking</td>
<td>0.33</td>
<td>0.18</td>
<td>3.39</td>
<td>.066</td>
<td>1.39</td>
<td>0.98–1.98</td>
</tr>
<tr>
<td>Physical activity</td>
<td>-0.02</td>
<td>0.02</td>
<td>1.22</td>
<td>.27</td>
<td>0.98</td>
<td>0.95–1.01</td>
</tr>
<tr>
<td>Early-life SES</td>
<td>0.41</td>
<td>0.21</td>
<td>3.72</td>
<td>.054</td>
<td>1.51</td>
<td>0.99–2.29</td>
</tr>
<tr>
<td>Current SES</td>
<td>0.20</td>
<td>0.21</td>
<td>0.84</td>
<td>.36</td>
<td>1.22</td>
<td>0.80–1.85</td>
</tr>
<tr>
<td>Early-life SES by current SES</td>
<td>-0.30</td>
<td>0.21</td>
<td>2.03</td>
<td>.15</td>
<td>0.74</td>
<td>0.50–1.12</td>
</tr>
</tbody>
</table>

Model 4 (N = 328)

| Age                        | 0.03 | 0.01 | 11.29| .001 | 1.04    | 1.01–1.06           |
| Sex                        | 0.16 | 0.21 | 0.57 | .45  | 1.17    | 0.78–1.76           |
| Race/ethnicity             | 0.04 | 0.25 | 0.03 | .87  | 1.04    | 0.64–1.69           |
| Alcohol use                | -0.25| 0.15 | 2.74 | .098 | 0.78    | 0.58–1.05           |
| Cigarette smoking          | 0.33 | 0.18 | 3.39 | .066 | 1.39    | 0.98–1.98           |
| Physical activity          | -0.02| 0.02 | 1.22 | .27  | 0.98    | 0.95–1.01           |
| Early-life SES             | 0.41 | 0.21 | 3.72 | .054 | 1.51    | 0.99–2.29           |
| Current SES                | 0.20 | 0.21 | 0.84 | .36  | 1.22    | 0.80–1.85           |
| Early-life SES by current SES | -0.30 | 0.21 | 2.03 | .15  | 0.74    | 0.50–1.12           |

SE = standard error; CI = confidence interval for \(\exp (B)\); SES = socioeconomic status.

* \(p < .05\).

**TABLE 4. The Association Between Early-Life and Adult SES With Continuous MetS Factor Scores**

<table>
<thead>
<tr>
<th>Model 1 (N = 354)</th>
<th>(F)</th>
<th>(df)</th>
<th>(p)</th>
<th>Partial (\eta^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Early-life SES</td>
<td>6.73</td>
<td>1; 352</td>
<td>.010*</td>
<td>0.019</td>
</tr>
<tr>
<td>Current SES</td>
<td>0.76</td>
<td>1; 352</td>
<td>.39</td>
<td>0.002</td>
</tr>
<tr>
<td>Model 2 (N = 354)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early-life SES</td>
<td>6.71</td>
<td>1; 350</td>
<td>.010*</td>
<td>0.019</td>
</tr>
<tr>
<td>Current SES</td>
<td>0.37</td>
<td>1; 350</td>
<td>.54</td>
<td>0.001</td>
</tr>
<tr>
<td>Early-life SES by current SES</td>
<td>1.03</td>
<td>1; 350</td>
<td>.31</td>
<td>0.003</td>
</tr>
</tbody>
</table>

Model 3 (N = 354)

| Age                        | 26.5  | 1; 318| <.001*| 0.077            |
| Sex                        | 33.54 | 1; 318| <.001*| 0.101            |
| Race/ethnicity             | 0.009 | 1; 318| .92   | <0.001           |
| Alcohol use                | 0.70  | 1; 318| .40   | 0.002            |
| Cigarette smoking          | 1.35  | 1; 318| .25   | 0.004            |
| Physical activity          | 4.19  | 1; 318| .041* | 0.013            |
| Early-life SES             | 3.49  | 1; 318| .063  | 0.011            |
| Current SES                | 0.02  | 1; 318| .88   | <0.001           |
| Early-life SES by current SES | 0.60  | 1; 318| .44   | 0.002            |

\(df =\) degrees of freedom; SES = socioeconomic status.

Partial \(\eta^2\) is reported as a measure of effect size.

* \(p < .05\).
diagnosis compared with those with high early-life SES. Is this difference clinically meaningful? One way to address this question is to compare its magnitude with that of other established risk factors. In this regard, the 83% higher likelihood of MetS diagnosis in those from low early-life SES backgrounds is comparable with the MetS risks associated with cigarette smoking in this sample and the risks associated with 16 years of aging. Furthermore, those meeting the criteria for MetS diagnosis have a two-fold increased risk of cardiovascular events or death (5) and a five-fold increased risk of diabetes (1), which would translate to substantially higher mortality events or death (5) and a five-fold increased risk of diabetes diagnosis have a two-fold increased risk of cardiovascular of aging. Furthermore, those meeting the criteria for MetS di-
agnosis compared with those with high early-life SES. Is this difference clinically meaningful? One way to address this question is to compare its magnitude with that of other established risk factors. In this regard, the 83% higher likelihood of MetS diagnosis in those from low early-life SES backgrounds is comparable with the MetS risks associated with cigarette smoking in this sample and the risks associated with 16 years of aging. Furthermore, those meeting the criteria for MetS diagnosis have a two-fold increased risk of cardiovascular events or death (5) and a five-fold increased risk of diabetes (1), which would translate to substantially higher mortality levels at the population level.

The nonsignificant main effect of current SES may seem puzzling in light of the previous literature revealing concurrent associations between adult SES and MetS risk (1). However, many individuals who experience low SES in adulthood are exposed to socioeconomic disadvantage throughout their lifespan (23), so it is unclear whether adult SES in any of these studies is an independent risk factor or reflects cumulative disadvantage. Only a few studies have explicitly tested the relative roles of childhood and adult SES against each other, and among these reports, the evidence is quite mixed. When modeling childhood and adult SES together, some studies with large samples have found that only childhood SES retains explanatory power (16,17,24,25), consistent with our findings. Some have found that only adult SES is significant (26) or that both childhood and adult SES are significant (27–29). Prospective studies found a prevailing role of childhood SES over adult SES (17,24,25), whereas retrospective studies (26) tended to find a stronger role for concurrent SES, suggesting that when precisely measured, childhood SES might play a stronger role in shaping MetS risk than adult SES. The novel contribution of the present study is that it isolates the effects of early-life and current SES from each other. From a scientific perspective, this is a useful approach. However, because life-course continuity in SES is the norm in many countries, particularly the United States, the patterns seen here may not reflect trends at the population level. Upwardly and downwardly mobile people are overrepresented in our sample relative to the population at large.

The lack of a main effect of current SES in the present study could also be interpreted in light of the statistical interaction observed. Low current SES was only associated with increased risk of MetS diagnosis for individuals who also experienced low early SES, and there was no evidence of metabolic risk associated with downward mobility. These patterns suggest the possibility that current low SES shapes MetS risk but may require sensitization by adverse early-life experiences. There is certainly some suggestive evidence from nonhuman animals that early-life conditions may have developmental programming effects that affect lifelong metabolism (13,15,20). Perhaps with the closure of putative sensitive periods for shaping neuroendocrine and metabolic function, the role of environmental adversity becomes progressively weaker across the lifespan. Or perhaps, it is simply an issue of duration of exposure, and the metabolic risk associated with current low SES is contingent on a sufficient previous experience of disadvantage. Future studies will have to test these competing explanations.

An additional consideration in interpreting the findings is that the present sample was Canadian. Socioeconomic gradients in MetS components such as obesity are less steep in Canada, when compared with countries that are culturally similar such as the United States (44). It is possible that in societies with less pronounced health disparities, adverse experiences need to occur during sensitive or vulnerable periods for these effects to manifest, but more cross-national research will be needed to test this hypothesis.

Turning to the interaction between early-life and current SES, we found varying patterns when examining the three MetS outcomes. Namely, there was a significant interaction between early-life and current SES in predicting the binary MetS variable, such that diagnosis rates were higher among the participants with a low-low pattern relative to the individuals with other trajectories. This interaction was nonsignificant in predicting the ordinal and continuous metabolic outcomes. This is unlikely to be an artifact of differential statistical power, because binary outcomes generally require greater power to detect than continuous outcomes. One possibility is that the MetS diagnosis may reflect an underlying pathological process, which is more than the sum of its parts, as has been discussed (33,34). This putative disease state might be less well measured by the component count and factor scores, thus diluting some of the associations with SES indices.

Nevertheless, the conflicting patterns of findings make it difficult to draw firm conclusions about the importance of mobility. Based on the binary diagnostic outcome, it seems that upward mobility can offset the MetS risks associated with low early SES. Indeed, in the upwardly mobile group (low-high trajectory), the prevalence of MetS was statistically identical to the high-high and high-low groups, a pattern that is consistent with at least one previous study (32). However, the nonsignificant interactions in
the models with ordinal and continuous metabolic outcomes sug-
gest that these offsetting influences of mobility are not especially
robust. These patterns are consistent with an emerging develop-
mental literature, which suggests that upward mobility may entail
some cost to health, particularly for individuals from highly disad-
vantaged backgrounds (45–47). Replication will also be needed to
rule out the possibility of type 1 error.

The present study also found no evidence that downward mo-
bility was associated with detrimental effects on metabolic health.
This suggests that high early-life SES may confer resilience
against later disruptions, supporting the notion that intervening
early in the lifespan may be beneficial. Nevertheless, we cannot
rule out the possibility that the negative effects of downward
mobility in adulthood might simply take longer to manifest,
as discussed previously, or that we had limited statistical power
to detect these effects given that the sample size for individuals
in the high-low group was slightly smaller than the others.
Furthermore, given the correlational nature of these analyses,
conclusions regarding social mobility are merely suggestive of
patterns that should be explored using experimental (e.g., cash
transfer interventions) or quasi-experimental (e.g., adoption
studies) methodology.

The present study’s design provides a novel contribution to our
inferences about the relative roles of early-life versus later SES by
disentangling two SES dimensions that typically covary in the
population. The inevitable trade-off of the recruitment strategy in
this study was that the sample was not designed to be a represen-
tative section of the Canadian population, which may limit the
generalizability of the findings. The participants studied here
were also younger than the general adult population (only those 55
years or younger were recruited) and healthier (no history of chronic
disease), and the participants who experienced upward or downward
mobility were oversampled, which allowed us to have enough
participants to begin to model these effects that have generated
mixed findings in the literature (28,30–32). It is not surprising
then that the average rate of MetS in our sample (10.2%) was
slightly lower than that of the adult Canadian population
(19.1%, Riediger and Clara 48), suggesting that the effects we
report here might be even stronger in the general population.
Other limitations include the retrospective assessment of early-
life SES and the absence of measures of childhood metabolic
health. Given that poor health in childhood is associated with
lower occupational status in adulthood (49), multivariate prog-
nostic longitudinal studies that monitor changes in both SES and health
are an important next step for testing social causation versus
social selection hypotheses about the links between low SES and health. In addition, these prospective studies should shed
light on potential mediators and moderators explaining the
association between low SES and health. Given that poor health in childhood is associated with lower occupational status in adulthood (49), multivariate past
longitudinal studies that monitor changes in both SES and health
are an important next step for testing social causation versus
social selection hypotheses about the links between low SES and health. In addition, these prospective studies should shed
light on potential mediators and moderators explaining the
relation between low SES and current MetS risk, which were
not examined here—for example, positive affect (50), sleep
(51), work stress (52), practice of relaxation techniques (53), and
others.

Despite these limitations, the current study provides new ins-
sights into the associations between socioeconomic disadvantage
and risk of MetS, which seemed to be more strongly associated
with early-life SES than current adult SES. If this observation is
supported by future studies, one implication may be that early
childhood is an opportune period for targeting interventions to re-
duce the risk of MetS across the lifespan.

We thank the participants for their contribution to this project.

Source of Funding and Conflicts of Interest: This research was
supported by National Institutes of Health Grants R01 HD058502
and F32 HD078048. The authors report no conflicts of interest.

REFERENCES

777–822.

2. Park YW, Zhu S, Palaniappan L, Heshka S, Carnethon MR, Heymsfield SB. The
metabolic syndrome: prevalence and associated risk factor findings in the US
population from the Third National Health and Nutrition Examination Survey,

364–73.

4. Alberti KG, Zimmet P, Shaw J. Metabolic syndrome—a new world-wide defi-
nition. A consensus statement from the International Diabetes Federation. Diabet

5. Gami AS, Witt BJ, Howard DE, Erwin PJ, Gami L, Somers VK, Monori VM.
Metabolic syndrome and risk of incident cardiovascular events and death: a sys-
tematic review and meta-analysis of longitudinal studies. Am Coll Cardiol


7. Hossain P, Kawar B, El Nabah M. Obesity and diabetes in the developing

8. Manuck SB, Phillips JE, Gianaros PJ, Flory JD, Muldoon MF. Subjective socio-
economic status and presence of the metabolic syndrome in midlife community

9. Ortiz MS, Myers HF, Schetter CD, Rodriguez CJ, Seeman TE. Psychosocial pre-
dictors of metabolic syndrome among Latino groups in the Multi-Ethnic Study

ities in the metabolic syndrome and coronary heart disease among middle-aged

11. Chun S, Lee S, Son HJ, No H, Oh HY, Jung HB, Lee HJ, Kang JH, Song HJ,
Park YJ, Park KH. Clinical characteristics and metabolic health status of obese


between childhood and adulthood predictors of obesity and metabolic syndrome.

14. Gustafsson PE, Persson M, Om AH. Life course origins of the metabolic syn-
drome in middle-aged women and men: the role of socioeconomic status and obe-
sive risk factors in adolescence and early adulthood. Am J Epidemiol 2011;21:
103–10.

15. Tamayo T, Christian H, Rathmann W. Impact of early psychosocial factors (child-
hood socioeconomic factors and adversities) on future risk of type 2 diabetes,
metabolic disturbances and obesity: a systematic review. BMC Public Health
2010;10:525.

16. Lehman B, Taylor S, Kiefe C, Seeman T. Relation of childhood socioeconomic
status and family environment to adult metabolic functioning in the CARDIA

17. Non AL, Rewak M, Kawachi I, Gilman SE, Lowks EB, Appleton AA, Roman
JC, Buka SL. Childhood social disadvantage, cardiometabolic risk, and chronic

and childhood maternal education level, job status. Findings from the Korean Na-
35:207–15.

19. Gustafsson PE, Hammarstroem A. Socioeconomic disadvantage in adolescent
women and metabolic syndrome in mid-adulthood: an examination of pathways

20. Perunidou P, Chrousos GP. Metabolic consequences of stress during childhood

21. Miller GE, Chen E, Parker KJ. Psychological stress in childhood and susceptibil-
ity to the chronic diseases of aging: moving toward a model of behavioral and bi-

22. Nusslock R, Miller GE. Early-life adversity and susceptibility to the chronic
diseases of aging: moving toward a model of behavioral and biological mecha-

23. Wagener R, Adelman R. Childhood and intergenerational poverty: the long-
term consequences of growing up poor. New York, NY: National Center for Chil-
dren in Poverty; 2009.
Early-Life Socioeconomic Disadvantage


