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Longitudinal Associations between Overweight/Obesity and Stress Biology in Low-Income Children

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Abstract

Background/Objectives—Associations between overweight and altered stress biology have been reported cross-sectionally during childhood, but it is unclear whether overweight precedes altered stress biology or if altered stress biology predicts greater likelihood of overweight over time. The current longitudinal study investigates associations between overweight/obesity, salivary alpha amylase and cortisol morning intercept, diurnal slope, and reactivity to social stress in a cohort of low-income children during preschool and middle childhood.

Subjects/Methods—Children were recruited through Head Start and were observed and followed into middle childhood (N = 257; M = 8.0 years). Height and weight were measured at both time points. Saliva samples were collected across the day and in response to a social challenge at both ages for alpha amylase and cortisol determination.

Results—Cross-lagged panel analyses indicated that overweight/obesity at preschool predicted lower morning alpha amylase ($\beta = -0.18$, 95% CI: -0.34, -0.03; p = .023), lower morning cortisol ($\beta = -0.22$, 95% CI: -0.38, -0.06; p = .006), lower sAA diurnal slope ($\beta = -0.18$, 95% CI: -0.34, -0.03; p = .021), and lower cortisol stress reactivity ($\beta = -0.19$, 95% CI: -0.35, -0.02; p = .031) in middle childhood. Lower alpha amylase reactivity at preschool was the only biological factor that predicted higher likelihood of overweight/obesity at middle childhood ($\beta = -0.20$, 95% CI: -0.38, -0.01; p = .035).

Conclusions—These findings suggest that overweight/obesity may be driving changes in stress biology across early to middle childhood, particularly in down-regulation of morning levels of

Users may view, print, copy, and download text and data-mine the content in such documents, for the purposes of academic research, subject always to the full Conditions of use:http://www.nature.com/authors/editorial_policies/license.html#terms Corresponding Author: Jenalee Doom; University of Denver, Frontier Hall, 2155 S. Race St., Denver, CO 80210. Jena.Doom@du.edu. **Conflicts of Interest:** The authors have no conflicts of interest to disclose. stress hormones, diurnal sAA slope, and cortisol reactivity to stress, rather than stress biology driving overweight/obesity.

Keywords

Early childhood; middle childhood; cortisol; alpha amylase; overweight; BMI

Childhood and adolescent obesity rates have been increasing in the past three decades (1). In developed countries, over one in five children have overweight or obesity, and rates have also been increasing in developing countries (1). In the United States, children living in poverty are more likely to have overweight or obesity than children from higher socioeconomic groups (2). Overweight and obesity in childhood and adolescence are strong predictors of obesity in adulthood, so it is important to understand childhood factors that contribute to overweight and obesity in order to create early prevention and treatment interventions. It is likely that a combination of behavioral, biological, and environmental factors, and interactions between these factors, are involved in the increase in obesity over time, particularly for children living in poverty.

Associations between overweight/obesity and stress biology have been demonstrated in both children and adults, with overweight/obesity associated with disruptions in the hypothalamic-pituitary-adrenal (HPA) axis and the sympathetic nervous system (SNS). Typical functioning of the HPA axis involves a cascade of sequential release of corticotropin-releasing hormone (CRH) from the hypothalamus, adrenocorticotropic hormone (ACTH) from the pituitary, and the glucocorticoid cortisol from the adrenal cortex. The HPA axis is involved in the regulation of metabolism, responses to challenge, and a host of other processes critical to homeostasis. Altered HPA axis regulation has been implicated in the development of overweight and obesity. In mammals, glucocorticoids maintain and enhance energy stores (3), and in humans, too, cortisol plays a central role in regulating food intake and metabolism (4). Obesity also impacts cortisol regulation, as adipose tissue can generate cortisol, and there are interactions between adipose tissue and the HPA axis (5). Given this extensive cross-talk between systems coordinating stress responses, food intake, and metabolism (3), the dynamic patterns between cortisol regulation and obesity over time need to be investigated. Two functional measures of HPA biology, cortisol reactivity to stress and diurnal cortisol secretion, reflect the ability of the HPA axis to regulate responses to stress and to modulate circadian rhythm. These measures have been associated with overweight/obesity and will be assessed in the current study.

Associations between overweight/obesity and diurnal cortisol in children and adolescents have been mixed (6–13), and nearly all have been derived from cross-sectional studies, precluding an understanding of how associations between overweight and diurnal cortisol may be unfolding over time. One longitudinal study in adolescents provides evidence that a blunted diurnal cortisol pattern is associated with higher concurrent body mass index (BMI) and increasing BMI over time (14). In children with overweight and obesity, lower cortisol in the early and late morning and the evening have been observed compared to children with normal weight (8). Hypocortisolism has been associated with overweight for girls directly and mediated by reduced satiety responsiveness, and for boys, the association is mediated

through emotional overeating (9). In girls aged 8–13 years, heightened cortisol reactivity is associated with higher BMI (15). Likewise, cross-sectional evidence suggests that for older children (8–9 years), higher cortisol reactivity is associated with higher BMI, but this association is not present in younger children (5–7 years) (16). However, there is also evidence in preschool children that a blunted cortisol response to stress is associated with a higher BMI (17).

Associations between body mass index and SNS activity have frequently been reported (18-21). The SNS promotes secretion of norephinephrine in response to stress, which leads to increases in the enzyme salivary alpha amylase (sAA) (22); therefore, sAA has been used as a biomarker of SNS activity (23). sAA shows a diurnal pattern, with a rise across the day (24). There is evidence that chronic stress down-regulates the system, with children who have experienced chronic stress showing lower basal sAA patterns (25). Low SNS activity has been associated with low resting metabolic rate (18, 19), and medications that increase SNS activity have been demonstrated to reduce food intake (20). However, low SNS-obesity associations may be tissue-specific since high SNS activity may be more likely to promote pathogenesis in certain tissues such as the heart or blood vessels (e.g., hypertension) (26). These findings suggest that low SNS activity could be a risk factor for overweight and obesity, which could be exacerbated in children experiencing chronic stress. Alternatively, overweight could lead to greater SNS disruptions over time. For example, a higher BMI zscore at 2.5 years predicted lower cardiac reactivity to stress at age 5 years, showing a blunting of SNS reactivity over time (27). Overall, associations between basal SNS activity and overweight/obesity in children have been mixed (21, 28–31). In the studies using sAA as the marker of SNS activity, sAA output across the day was lower in school-aged girls with obesity than their normal weight counterparts (32). Lower morning sAA, a higher rise in sAA across the day, and blunted sAA reactivity were associated with increased BMI zscores in low-income preschool-aged children (33). In one longitudinal study examining cortisol, sAA, and overweight/obesity in toddlers, lower morning sAA and higher sAA slope across the day at 27 months predicted a greater likelihood of overweight at 33 months for girls. For boys, overweight at 21 months predicted lower morning sAA at 27 months (34).

As most prior work has been cross-sectional, little is known about the directionality of associations between overweight and stress biology, particularly in children. Low socioeconomic status and higher levels of stress are predictors of higher BMI; thus, investigating these associations longitudinally in low-income, highly stressed populations is a high priority for creating interventions that promote healthy weight and adaptive regulation of stress biology. The current study investigates longitudinal associations between overweight/obesity, cortisol, and sAA in low-income children, who are at higher risk for overweight/obesity or stress biology—or both—drive changes in biology and weight status is essential for identifying developmental windows for prevention and intervention efforts that can address the child overweight and obesity epidemic. Our hypotheses were that cortisol and sAA that were lower in the morning and showed lower reactivity to stress would predict later overweight/obesity. These hypotheses were

part of a secondary data analysis rather than primary hypotheses for the original data collection.

Methods

Participants

The current study uses data from the ABC Preschool and Kids cohort (9, 17, 33). Children and their parent(s) were recruited in preschool through Head Start, a federally-funded program for children from low-income backgrounds in the United States. A form was sent home to recruit children and their primary caregiver (92.6% mothers) for the study. Parents who returned the form and provided their contact information were compensated with \$10. Parents were contacted to confirm eligibility and interest in participation. Exclusion criteria included: child or parent did not speak English; primary caregiver had a 4-year college degree or greater (to target a low-income sample); child was in foster care; child had medical problems, food allergies, or perinatal complications; and gestational age < 35 weeks. Children were retained for the current analyses if they had valid data for cortisol and sAA reactivity or diurnal regulation at middle childhood. Informed consent was obtained, and the university's institutional review board approved this study.

Procedure

Children and parents participated in three assessments: two during preschool age (1st assessment age 2.9–5.2 years, N = 380; 2nd assessment age 3.2–7.1 years, N = 330) and one during middle childhood (age 7.0–10.2 years, N = 257). At the first preschool assessment, parents completed questionnaires on demographics and income, and children's height and weight were assessed. Diurnal salivary samples were collected from children at preschool for cortisol and alpha-amylase assessment 3 times per day for 3 days (morning, noon, late afternoon). At the second preschool assessment, five saliva samples were collected from the child in response to a social stressor for cortisol and alpha amylase, and children's height and weight were assessed. At the middle childhood assessment, parents completed questionnaires on demographics and income, and children's height and weight were assessed. Diurnal salivary samples were collected by parents at home 3 times per day for 3 days (morning, late afternoon, bedtime). MEMS caps were used for home data collection by parents to ensure timely home collection (92% accuracy within 15 minutes). Research assistants collected five saliva samples for sAA and cortisol determination in response to a social stressor. Details on saliva collection and stress tasks are in Supplement Sections 1.1– 1.4. Trained research assistants measured child weight and height without shoes or heavy clothing at all three assessments according to standard protocols (35) (details in Supplement Section 1.5). Overweight/obesity was defined as 85th percentile for BMI based on US Centers for Disease Control and Prevention growth charts for age and sex at each assessment (36).

Assays

Saliva samples were stored at -20° C until processing. Saliva samples were then thawed, vortexed, centrifuged for 15 minutes at 3000 rpm, separated from debris, and placed in Thermo Scientific Matrix Racks at -80° C. The same technician conducted all assays within

each assessment using the same equipment following manufacturer's instructions. An Expanded Range High Sensitivity Salivary Cortisol Enzyme Immunoassay Kit (Catalog No. 1–3002, 96-Well Kit, Salimetrics LLC, PA, USA) with a detection limit of $0.007\mu g/dL$ was used to assay cortisol. At the first preschool assessment, the intra and inter-assay coefficients of variation (CV) were 7%. The sensitivity of the assays was $0.003 \mu g/dL$. At the second preschool assessment, the intra and inter-assay CVs were 4.6% and 5.5%, respectively. At the middle childhood assessment, the average inter-assay CV was 4.0% and the intra-assay CVs were from 0.8-6.1%. Free cortisol is reported in $\mu g/dL$.

For alpha amylase, samples were assayed in duplicate with an alpha amylase kinetic reaction assay kit (Catalog No. 1–1902, 96-Well Kit, Salimetrics LLC, PA, USA). This assay uses a chromagenic substrate, 2-chloro-pnitrophenol linked with maltotriose, and the enzymatic action of alpha-amylase on this substrate produces 2-chloro-p-nitrophenol, which is measured spectrophotometrically 2 minutes after the reaction start time with a calibrated plate reader at 405 nm wavelength. The amount of alpha amylase activity is directly proportional to the increase in absorbance at 405 nm. Low, medium and high sAA controls were present in each assay. At the preschool assessments, intra-assay CVs were <6.5%, and the inter-assay CVs were <4.8%. At middle childhood, sAA intra-assay CVs averaged 4.8% and inter-assay CVs averaged 5.0%. The sensitivity (0.01 units) is determined by the lower change in absorbance reading in each assay. Any sample below the low alpha amylase control was assayed again using a dilution recommended by the manufacturer to achieve a higher concentration and a greater absorbance reading. sAA is reported in enzyme units per milliliter (U/ml).

Data analytic plan

Cross-lagged panel analysis—Three cross-lagged models were fit using Mplus 7.1.4 to assess temporal associations between overweight/obesity, sAA, and cortisol at preschool and middle childhood assessments. All three models used overweight/obesity vs. normal weight (categorical variable) at both time points. The cortisol and sAA parameters used at both time points varied in each of the models. The first model used cortisol and sAA morning intercepts (standardized), the second used cortisol and sAA diurnal slopes (standardized), and the third used cortisol and sAA reactivity (AUCi). See Supplemental Information Section 2 for intercept, slope, and reactivity calculations. For the first and second models (diurnal cortisol and sAA), the preschool data was from the first preschool assessment. For the third model (cortisol and sAA reactivity), the preschool data was from the second preschool assessment. All of the variables in the model were observed. All pathways between Time 1 and Time 2 overweight/obesity, cortisol, and sAA were estimated simultaneously, which allows for more complex models than assessing multiple linear regressions. Covariates included age, sex, race/ethnicity, parent-reported birthweight, sleep quality, and medication use theorized to affect cortisol/sAA at preschool and middle childhood (see Supplement Sections 1.6–1.9). Pubertal development was included as a covariate for the middle childhood variables. Model fit was assessed using recommended guidelines in the field (37, 38), including the comparative fit index (CFI; > .90) and the root mean square error of approximation (RMSEA; .05 denotes good fit, .08 denotes adequate fit).

Results

Descriptive statistics

A total of 257 children had salivary cortisol and sAA data available at the preschool and middle childhood assessments (see Table 1 for demographics). Children in the current study did not differ from children who participated in earlier waves of the study or from those who did not provide cortisol at the middle childhood assessment as a function of the following T1 variables: age, sex, BMIz, income-to-needs ratio, primary caregiver education level, or race/ ethnicity (all *p*s > .05). Due to missing data, 230 children were included in the diurnal cortisol/sAA analyses, and 219 were included in the cortisol/sAA reactivity analyses. Age was recorded at the first preschool assessment (M = 4.3 years, SD = 0.5), the second preschool assessment (M = 4.9 years, SD = 0.7), and the middle childhood assessment (M = 8.0 years, SD = 0.7). At the first preschool assessment, 39.7% of the sample was classified as having overweight/obesity, 40.9% as having overweight/obesity at the second preschool assessment, and 48.0% as having overweight/obesity in middle childhood. The income-to-needs ratio in preschool was 0.85 (SD = 0.68), indicating that children were generally living in low-income households.

Model fit indices

All three models demonstrated good fit according to the CFI. The intercept model had an RMSEA of 0.085, which demonstrates adequate fit, while the other two models demonstrated good fit with the RMSEA criteria.

Cortisol/sAA morning intercept and overweight/obesity

Overweight/obesity at preschool (first assessment) predicted a lower sAA morning intercept ($\beta = -0.18, 95\%$ CI: -0.34, -0.03; Table 2; Figure 1a), lower cortisol morning intercept ($\beta = -0.22, 95\%$ CI: -0.38, -0.06), and greater likelihood of overweight/obesity in middle childhood ($\beta = 0.85, 95\%$ CI: 0.75, 0.95). Higher sAA morning intercept at preschool predicted a higher sAA morning intercept at middle childhood ($\beta = 0.64, 95\%$ CI: 0.57, 0.71). There was a significant within-time association between overweight/obesity and lower cortisol morning intercept in middle childhood ($\beta = -0.25, 95\%$ CI: -0.47, -0.03). The model explained 78.2% of the variance in overweight/obesity, 50.2% of the variance in sAA morning intercept at middle childhood.

Cortisol/sAA diurnal slope and overweight/obesity

Overweight/obesity at preschool predicted a more blunted increase in sAA across the day ($\beta = -0.18$, 95% CI: -0.34, -0.03; Table 3; Figure 1b) and greater likelihood of overweight/ obesity at middle childhood ($\beta = 0.85$, 95% CI: 0.75, 0.95). A blunted cortisol slope at preschool predicted a steeper cortisol slope at middle childhood ($\beta = -0.21$, 95% CI: -0.32, -0.10). Overweight/obesity at middle childhood predicted within-time associations with a blunted rise in sAA across the day ($\beta = -0.32$, 95% CI: -0.57, -0.06) and a more negative cortisol slope across the day ($\beta = -0.40$, 95% CI: -0.62, -0.17). The model explained 78.0%

of the variance in overweight/obesity, 6.8% of the variance in sAA slope, and 13.8% of the variance in cortisol slope at middle childhood.

Cortisol/sAA reactivity and overweight/obesity

Overweight/obesity at preschool predicted more blunted cortisol reactivity ($\beta = -0.19$, 95% CI: -0.35, -0.02) and greater likelihood of overweight/obesity at middle childhood ($\beta = 0.86$, 95% CI: 0.77, 0.96; Table 4; Figure 1c). More blunted sAA reactivity at preschool predicted a greater likelihood of overweight/obesity in middle childhood ($\beta = -0.20$, 95% CI: -0.38, -0.01). The association between higher sAA reactivity at preschool and higher sAA reactivity in middle childhood was not statistically significant at p = 0.08 ($\beta = 0.12$, 95% CI: -0.01, 0.25). There were no within-time associations between overweight/obesity and sAA or cortisol reactivity at preschool or middle childhood. The model explained 82.5% of the variance in overweight/obesity, 7.4% of the variance in sAA reactivity, and 13.2% of the variance in cortisol reactivity at middle childhood.

Sensitivity analyses

We conducted sensitivity analyses removing any participants who regularly take medications known to affect cortisol or sAA regulation and the results did not change (see supplement).

Discussion

The current study was the first to examine longitudinal associations between cortisol, sAA, and overweight/obesity across the preschool and middle childhood years, providing information about the directionality of observed associations between overweight/obesity and stress biology during childhood. Overall, analyses suggested that overweight/obesity predicted greater changes in stress biology over time, from early to middle childhood, rather than stress biology predicting increased likelihood of overweight/obesity over this time period. Specifically, overweight/obesity in preschool predicted future lower morning levels of cortisol and sAA, blunted cortisol reactivity, and a lower sAA slope across the day in middle childhood. However, the exception was that blunted sAA reactivity to stress in preschool predicted higher likelihood of overweight/obesity in middle childhood.

There are well-established associations between fat accumulation and cortisol regulation, which are consistent with findings in the current analyses. The literature supports that high cortisol levels and long-term HPA axis activation promote the accumulation of visceral fat over time (39). Elevated cortisol increases appetite and disrupts the regulation of energy balance (4). Greater cortisol secretion in adults with central obesity has been consistently noted (40). Conversely, a blunted diurnal cortisol pattern and low morning and evening cortisol levels have also been linked to higher BMI (8, 14). Thus, there are likely complex, bidirectional associations that increase likelihood of fat accumulation and cortisol dysregulation over time. In the current study, overweight/obesity at preschool predicted lower morning cortisol at middle childhood, suggesting down-regulation of the HPA axis with excess adipose tissue. This finding is consistent with research in adults reporting abdominal fat is associated with lower morning cortisol, suggesting down-regulation of the HPA axis in response to the negative feedback resulting from high cortisol levels that can be

secreted from fat tissue (41, 42). Chronically high levels of cortisol act on upstream mediators of the HPA axis (e.g., CRF, ACTH) to adaptively down-regulate the basal system to prevent the effects of chronic HPA activation (43), thus resulting in low morning cortisol levels. As heightened levels of morning cortisol are needed to mobilize energy resources, children with low morning cortisol levels may lack the resources necessary to behaviorally and biologically adapt to daily challenges (44). Low cortisol levels have been associated with overweight in children (8, 9) and the process leading to low cortisol could also increase vulnerability to certain health disorders (45).

In preschoolers, blunted cortisol reactivity has been observed in children with higher BMI zscores (17); current findings suggest that overweight/obesity in preschool predicts a more blunted cortisol response to stress in middle childhood. Higher levels of adipose tissue could lead to or exacerbate metabolic problems that increase risk for cardiovascular disease and metabolic syndrome (42). Blunted cortisol reactivity may be a marker of risk in this lowincome sample, particularly because blunted cortisol reactivity has been associated with social and emotional problems in high-risk children (46). Low morning cortisol levels and blunted cortisol reactivity could predispose vulnerable children to emotional or behavioral problems and may contribute to higher rates of these problems in children with overweight/ obesity (47). There are individual differences in cortisol and sAA regulation due to genetics and epigenetics (48, 49) as well as genetic and epigenetic differences in risk for overweight/ obesity (50). Certain genetic or epigenetic profiles could have significant effects on pathways from stress biology to overweight/obesity or from overweight/obesity to stress biology. Although genetic or epigenetic factors were not examined in the current study, this is an important area for future research.

The current study provides additional evidence that low sAA activity predicts and is predicted by overweight/obesity in children. This finding addresses one of the most important avenues for research in SNS activation and overweight/obesity by assessing whether high or low sAA activity predicts greater likelihood of overweight/obesity over time (18). However, it must be noted that SNS activation may be higher or lower in individuals with overweight or obesity depending on the region of the body measured (18) and the type of measurement (e.g., hypertension in individuals with obesity). The finding of lower morning sAA and lower sAA slope across the day in middle childhood following overweight/obesity in preschool could reflect a down-regulation of the SNS as lower sAA levels have been associated with chronic stress (25), which could have implications for future behavior and physical health. Attenuated morning SNS activity and lower sAA diurnal slope could be due to down-regulation from chronically high levels of SNS activity, similar to down-regulation in the HPA axis following high HPA activity, which may reflect a failure to adequately prepare for daily challenges in the context of chronic stress, such as the stress of living in poverty for low-income children (51). This finding is consistent with a longitudinal study in toddlers finding that overweight at 21 months predicted lower morning sAA at 27 months, although that finding was specific to boys (34). Blunting of morning sAA levels in middle childhood following overweight/obesity in preschool, particularly in the context of poverty, could increase the likelihood of maladaptive biological and behavioral responses to stress in the future, which could influence future health through a number of pathways. The finding that overweight/obesity predicted lower diurnal sAA slope was not

consistent with evidence in young children that higher sAA slope is concurrently associated with and predictive of higher BMI (33, 34). Further research is needed to understand whether age is an important moderator of these associations.

The only biomarker that predicted greater likelihood of overweight/obesity in middle childhood was blunted sAA reactivity to stress in preschool. This is consistent with evidence that low SNS activity is associated with low resting metabolic rate (18, 19) and greater food intake (20). If children with blunted sAA reactivity have a lower resting metabolic rate and greater food intake over time, without offsetting this intake with physical activity, they may be more likely to develop overweight/obesity. Previous work has shown that higher sAA is associated with satiety (52), suggesting that lower sAA levels could be associated with greater hunger and less satiety, which could lead to excessive food intake. Low SNS activity has also been associated with obesity in children (30). However, the current findings are inconsistent with some work reporting heightened SNS activity in individuals with obesity (21, 28, 29, 53, 54). Most studies did not focus on SNS reactivity, however, and were conducted cross-sectionally with older children or adults. As this is the first study to examine these pathways longitudinally into middle childhood, and sAA is an indirect marker of SNS activity, this association needs to be replicated in other samples. These associations may be specific to certain individuals with a positive energy balance (18), so we need to understand whether there are genetic, psychobiological, or environmental factors that moderate the association between SNS activation and overweight/obesity in childhood. Our sample is socioeconomically high-risk, so greater experiences of psychosocial stress or exposure to obesogenic environments likely influenced overweight/obesity and stress biology compared to low-risk populations. These findings also may not generalize to developmental periods outside early-to-middle childhood. It will be important to understand whether other aspects of SNS and HPA activity, such as chronic integrated cortisol measured in hair, are associated with overweight/obesity in a similar manner over time. Measures in different tissues address unique aspects of stress regulation and may show different associations with adiposity across development.

There were limitations to the current study. The stress reactivity task in preschool differed from the stress reactivity task in middle childhood. As preschool and middle childhood are very different developmental periods, the social stress tasks were designed to include a strong, developmentally appropriate social-evaluative component known to elicit stress responses at each age tested. Future work is needed to establish social stressors that are effective and similar across childhood. Timing of the diurnal saliva samples also differed between waves, with the preschool samples occurring between 8:30am and 4:30pm, and the middle childhood samples typically between 8am and 9pm. Our analytic strategy accounted for the timing of the samples when calculating the diurnal intercept and slope of sAA, but differences in methodology could still partially contribute to the results. Pubertal development was reported by parents, and thus may be biased or inaccurate compared to a medical exam. We did not measure physical activity as a potential covariate. We also used only BMI z-score as our measure of adiposity, and future research including additional measures of adipose tissue is needed. Finally, the study was limited to a low-income population in the rural Midwest, so it may not generalize to all children. We also did not include non-English speaking families in the study, so results will need to be replicated in

non-English speaking populations. The current study did not adjust for multiple comparisons due to the nature of the pre-specified comparisons in the model (55), though future studies are needed to replicate the current findings.

Conclusions

The current study suggests that disruptions in stress biology, particularly down-regulation of morning levels of stress-mediating hormones, cortisol reactivity to stress, and lower diurnal sAA slope are more likely to follow overweight/obesity in children rather than precede overweight/obesity. A blunted sAA stress response at preschool was the only biological predictor of overweight/obesity in middle childhood. Importantly, these associations were reported in low-income children, a population with an outsized burden of the obesity epidemic. This prospective longitudinal study is the first to map associations between overweight/obesity and stress biology from preschool to middle childhood, providing insight into the directionality of observed associations and the course of overweight/obesity and disruptions in stress biology. Future research is needed to understand the mechanisms between these associations to improve prevention and intervention efforts that aim to enhance child health.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

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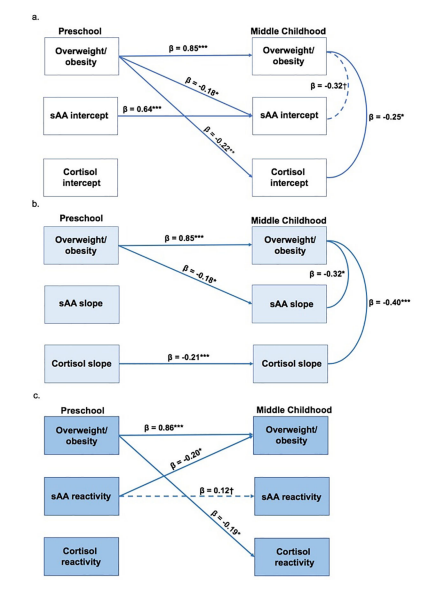


Figure 1.

Model tested for associations between overweight/obesity and 2a) sAA and cortisol intercept, 2b) sAA and cortisol diurnal slope, and 2c) sAA and cortisol reactivity. β values are standardized estimates. Statistical significance indicated by $\dagger p < 0.10$, *p < 0.05, **p < 0.01, ***p < 0.001. During preschool, sAA and cortisol morning intercept and diurnal slope were measured at the first assessment, and sAA and cortisol reactivity were measured at the second assessment.

Table 1.

Participant Characteristics.

	M	SD	%
Preschool: Age First Assessment (years)	4.25	0.52	
Preschool: Age Second Assessment (years)	4.88	0.70	
Middle Childhood: Age (years)	7.95	0.71	
Time Between Preschool Assessments (months)	7.46	5.35	
Time Between First Preschool and Middle Childhood Assessments (months)	44.40	7.89	
Female			49.0
Middle Childhood Pubertal Status	1.47	0.59	
Child Race/Ethnicity			
Non-Hispanic White			52.9
African American			16.7
Hispanic/Latino			10.1
American Indian			0.4
Asian/Pacific Islander			0.8
Multiracial			19.1
Parent Education			
Did Not Graduate High School			17.5
High School Degree or GED			30.0
Some College Courses			40.1
2-year College Degree			12.5
Preschool Income-to-Needs Ratio	0.85	0.68	
Middle Childhood Income-to-Needs Ratio	1.11	0.75	
BMIz at Preschool (First Assessment)	0.83	1.09	
Overweight/Obesity at Preschool (First Assessment)			39.7
BMIz at Preschool (Second Assessment)	0.86	1.09	
Overweight/Obesity at Preschool (Second Assessment)			40.9
BMIz at Middle Childhood	0.97	0.99	
Overweight/Obesity at Middle Childhood			47.9

Note. Means, standard deviations, and percentages of participants' demographic information and key variables. N = 257. T1 = preschool assessment, T2 = middle childhood assessment. GED = General Educational Development Test (high school equivalency test in the United States). Percentages are calculated for all participants with valid data on that measure.

Table 2.

Overweight/obesity, sAA and cortisol morning intercept cross-lagged analysis.

	β	95% CI	<i>p</i> -value
	μ	95 /0 CI	<i>p</i> -value
Within-time paths			
T1 Overweight/obesity→T1 sAA intercept	-0.09	-0.27, 0.09	.31
T1 Overweight/obesity→T1 Cortisol intercept	-0.06	-0.23, 0.12	.53
T1 Cortisol intercept \rightarrow T1 sAA intercept	0.08	-0.07, 0.23	.28
T2 Overweight/obesity \rightarrow T2 sAA intercept	-0.32	-0.67, 0.03	.069 [†]
T2 Overweight/obesity→T2 Cortisol intercept	-0.25	-0.47, -0.03	.029*
T2 Cortisol intercept \rightarrow T2 sAA intercept	0.08	-0.11, 0.27	.41
Autoregressive paths			
T1 Overweight/obesity \rightarrow T2 Overweight/obesity	0.85	0.75, 0.95	<.001 ***
T1 sAA intercept \rightarrow T2 sAA intercept	0.64	0.57, 0.71	<.001 ***
T1 Cortisol intercept \rightarrow T2 Cortisol intercept	-0.05	-0.19, 0.10	.53
Cross-lagged paths			
T1 Overweight/obesity \rightarrow T2 sAA intercept	-0.18	-0.34, -0.03	.023*
T1 Overweight/obesity \rightarrow T2 Cortisol intercept	-0.22	-0.38, -0.06	.006**
T1 sAA intercept \rightarrow T2 Overweight/obesity	0.01	-0.11, 0.13	.88
T1 sAA intercept \rightarrow T2 Cortisol intercept	-0.01	-0.14, 0.11	.83
T1 Cortisol intercept \rightarrow T2 Overweight/obesity	0.06	-0.07, 0.19	.38
T1 Cortisol intercept \rightarrow T2 sAA intercept	0.00	-0.11, 0.11	.99

Statistical significance indicated by

$$^{\dagger} p < 0.10$$

$$\bar{p} < 0.05$$

** p<0.01

*** p<0.001.

T1 = preschool (first assessment), T2 = middle childhood assessment.

Table 3.

Overweight/obesity, sAA and cortisol slope cross-lagged analysis.

	ß	95% CI	p-value
Within-time paths			
T1 Overweight/obesity→T1 sAA slope	0.09	-0.08, 0.27	.30
T1 Overweight/obesity→T1 Cortisol slope	-0.05	-0.21, 0.12	.57
T1 Cortisol slope \rightarrow T1 sAA slope	0.05	-0.09, 0.18	.52
T2 Overweight/obesity \rightarrow T2 sAA slope	-0.32	-0.57, -0.06	.016*
T2 Overweight/obesity→T2 Cortisol slope	-0.40	-0.62, -0.17	<.001 ***
T2 Cortisol slope \rightarrow T2 sAA slope	0.10	-0.02, 0.22	.10
Autoregressive paths			
T1 Overweight/obesity \rightarrow T2 Overweight/obesity	0.85	0.75, 0.95	<.001 ***
T1 sAA slope \rightarrow T2 sAA slope	0.00	-0.12, 0.13	.97
T1 Cortisol slope \rightarrow T2 Cortisol slope	-0.21	-0.32, -0.10	<.001 ***
Cross-lagged paths			
T1 Overweight/obesity \rightarrow T2 sAA slope	-0.18	-0.34, -0.03	.021 *
T1 Overweight/obesity→ T2 Cortisol slope	0.00	-0.17, 0.17	>.99
T1 sAA slope \rightarrow T2 Overweight/obesity	0.01	-0.11, 0.13	.86
T1 sAA slope \rightarrow T2 Cortisol slope	0.02	-0.09, 0.13	.72
T1 Cortisol slope \rightarrow T2 Overweight/obesity	0.04	-0.10, 0.17	.58
T1 Cortisol slope \rightarrow T2 sAA slope	-0.03	-0.15, 0.09	.63

Statistical significance indicated by

 $^{\dagger} p < 0.10$

*

** p<0.01

*** p<0.001.

T1 = preschool (first assessment), T2 = middle childhood assessment.

Table 4.

Overweight/obesity, sAA and cortisol reactivity cross-lagged analysis.

	β	95% CI	p-value
Within-time paths			
T1 Overweight/obesity→T1 sAA reactivity	0.11	-0.05, 0.28	.18
T1 Overweight/obesity→T1 Cortisol reactivity	-0.06	-0.14, 0.02	.12
T1 Cortisol reactivity \rightarrow T1 sAA reactivity	0.07	-0.06, 0.20	.27
T2 Overweight/obesity \rightarrow T2 sAA reactivity	-0.20	-0.70, 0.30	.43
T2 Overweight/obesity→T2 Cortisol reactivity	0.09	-0.31, 0.50	.65
T2 Cortisol reactivity \rightarrow T2 sAA reactivity	0.06	-0.04, 0.16	.22
Autoregressive paths			
T1 Overweight/obesity \rightarrow T2 Overweight/obesity	0.86	0.77, 0.96	<.001 ***
T1 sAA reactivity \rightarrow T2 sAA reactivity	0.12	-0.01, 0.25	.080 [†]
T1 Cortisol reactivity \rightarrow T2 Cortisol reactivity	0.02	-0.39, 0.42	.93
Cross-lagged paths			
T1 Overweight/obesity→ T2 sAA reactivity	0.16	-0.07, 0.38	.17
T1 Overweight/obesity \rightarrow T2 Cortisol reactivity	-0.19	-0.35, -0.02	.031*
T1 sAA reactivity \rightarrow T2 Overweight/obesity	-0.20	-0.38, -0.01	.035 *
T1 sAA reactivity \rightarrow T2 Cortisol reactivity	0.11	-0.03, 0.24	.12
T1 Cortisol reactivity \rightarrow T2 Overweight/obesity	0.03	-0.02, 0.09	.24
T1 Cortisol reactivity \rightarrow T2 sAA reactivity	-0.06	-0.30, 0.18	.64

Statistical significance indicated by

 $^{\dagger} p < 0.10$

** p<0.01

*** p<0.001.

T1 = preschool (second assessment), T2 = middle childhood assessment.