The relationship of early-life adversity with adulthood weight and cardiometabolic health status in the 1946 National Survey of Health and Development

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Word: 5569
Tables: 6

Conflicts of Interest and Source of Funding

RH is Director of the CLOSER consortium which is supported by funding from the Economic and Social Research Council (ESRC) and the Medical Research Council (MRC). An initial grant for this work was awarded to CLOSER in 2012 and extended, by the ESRC, in 2017 (award reference: ES/K000357/1). RH was supported by the UK Medical Research Council (Programme code: MC_UU_12019/2). The UK Medical Research Council provides core funding for the MRC National Survey of Health and Development.

WJ is supported by a UK Medical Research Council (MRC) New Investigator Research Grant (MR/P023347/1). WJ & MH acknowledge support from the National Institute for Health Research (NIHR) Leicester Biomedical Research Centre, which is a partnership between University Hospitals of Leicester NHS Trust, Loughborough University, and the University of Leicester.
This work was supported by a UK Medical Research Council (MRC) New Investigator Research Grant (MR/P023347/1).

All authors declare no conflicts of interest.
SUMMARY

Childhood adversity might cause obesity via the adoption of unhealthy behaviours. It might also cause poor cardiometabolic health, either via obesity or another pathway (e.g., altered stress response). Robson et al found that childhood adversity was associated with poor cardiometabolic health at ages 60-64 years, in both normal weight and overweight/obese adults. Further research is needed to understand the mechanisms by which childhood adversity might affect cardiometabolic health without necessarily increasing body weight.
ABSTRACT

Objectives
Evidence linking early-life adversity with an adverse cardiometabolic profile in adulthood is equivocal. This study investigates early-life adversity in relation to weight and cardiometabolic health status at age 60-64 years.

Methods
We included 1,059 individuals from the 1946 National Survey of Health and Development (NSHD). Data on adversity between ages 0-15 years were used to create a cumulative childhood psychosocial adversity score and a socioeconomic adversity score. Cardiometabolic and weight/height data collected at ages 60-64 years were used to create four groups: metabolically healthy normal weight (MHNW), metabolically unhealthy normal weight (MUNW), metabolically healthy overweight/obese (MHO), and metabolically unhealthy overweight/obese (MUO). Associations between the two exposure scores and weight/health status were examined using multinomial logistic regression, with adjustment for sex and age at the outcome visit.

Results
62% of normal weight individuals were metabolically healthy, whereas only 34% of overweight/obese individuals were metabolically healthy. In a mutually adjusted model including both exposure scores, a psychosocial score of ≥ 3 (compared to 0) was associated with increased risk of being metabolically unhealthy (compared to healthy) in both normal weight adults (RR 2.49; 95% CI 0.87, 7.13) and overweight/obese adults (1.87; 0.96, 3.61). However, the socioeconomic adversity score was more strongly related to metabolic health status in overweight/obese adults (1.60; 0.98, 2.60) than normal weight adults (0.95; 0.46, 1.96).
Conclusions

Independently of socioeconomic adversity, psychosocial adversity in childhood may be associated with a poor cardiometabolic health profile, in both normal weight and overweight/obese adults.

Key words

Childhood adversity, socioeconomic adversity, cardiometabolic profile, weight status, cohort study

Adverse childhood events (ACEs)
Avon Longitudinal Study of Parents and Children (ALSPAC)
Body mass index (BMI)
Glycosylated haemoglobin (HbA1c)
High-density lipoprotein-cholesterol (HDL-C)
Metabolically healthy normal weight (MHNW)
Metabolically healthy obesity (MHO)
Metabolically unhealthy normal weight (MUNW)
Metabolically unhealthy obesity (MUO)
National Survey for Health and Development (NSHD)
Percentage body fat (%fat)
INTRODUCTION

Obesity is a major public health concern, in part because it increases risk of cardiometabolic complications such as cardiovascular disease and type two diabetes.[1] Not all obese individuals, however, develop such conditions and there is heterogeneity in cardiometabolic health among individuals with obesity, and indeed within any weight status group.[2, 3] Categorising people based on their weight and health status is a common way to capture this heterogeneity. A lot of research has focused on the group with “so-called” metabolically healthy obesity (MHO), which is most commonly defined as obesity in the presence of fewer than two metabolic abnormalities, normally including dyslipidaemia, hypertension, and diabetes.[4]

Most of the existing literature on MHO has investigated its relationship with risk for future disease outcomes or death.[5, 6] While MHO is certainly not a benign condition, meta-analysed risks of outcomes like cardiovascular disease and type two diabetes are lower in this group compared against individuals with metabolically unhealthy obesity (MUO).[2, 3] Very few studies, however, have investigated which early-life risk factors might explain such heterogeneity in health prospects among individuals with obesity.[5, 6] The strongest evidence suggests that greater severity and duration of exposure to obesity is related to increased risk of MUO compared to MHO, but evidence on modifiable or targetable early-life risk factors is lacking.[7-9] Understanding what contributes to the risk of developing a metabolically unhealthy profile in different weight categories is important as it could inform prevention and intervention programmes to delay the onset of cardiometabolic complications, even among those individuals who are obese. While an exposure of interest (e.g., smoking) might be related to increased risk of MUO compared to MHO, it might not necessarily be related to health status among normal weight individuals. Such effect modification by weight status is equally as important to understand for prevention and
intervention programmes and might implicate obesity as part of the exposure-disease mechanism.

Previous studies have reported measures of psychosocial stress, such as adverse childhood events (ACEs) (e.g., abuse, neglect, and parental divorce) and socioeconomic adversities (e.g., low head of household social class, low income, household overcrowding, and low parental education), to be associated with increased risk of diseases like cardiovascular disease and type two diabetes.[10-13] The association of childhood adversity with a sub-clinical cardiometabolic profile appears, however, to be equivocal. Previous analyses in two birth cohort studies in the UK found no consistent evidence that childhood adversity is related to continuous measures of cardiometabolic disease risk.[14, 15] In a study in Dunedin, New Zealand, however, Danese et al found an association between greater cumulative childhood adversity and increased risk for cardiometabolic marker clustering at age 32 years in a sample of 1,037 normal weight, overweight, and obese individuals.[16] To our knowledge, this finding has not been further examined at a later age in adulthood and separately within different weight status groups.

Using data from the Medical Research Council 1946 National Survey for Health and Development (NSHD), this study investigates the relationships of early-life psychosocial and socioeconomic adversity with weight and cardiometabolic health status at age 60-64 years. We hypothesise that greater adversity will be associated with increased risk of MUO compared to MHO, and also of metabolically unhealthy normal weight (MUNW) compared to metabolically healthy normal weight (MHNW).

METHODS

Study Sample
The NSHD is an on-going study. The cohort initially consisted of 5,362 singleton births that occurred during one week in March 1946 in England, Wales, and Scotland.[17] Between 2006-2011, eligible participants (those known to be alive and living with an address in either England, Scotland, or Wales) were invited to attend a clinical assessment at one of six clinical research facilities or by a research nurse at home. A total 2,229 participants out of the 2,856 invited (78%) underwent assessment: 1,690 attended a clinical research facility and the remaining 539 were seen in their homes.[18] Invitations were not sent to those who had died (N = 778), who were living abroad (N = 570), had previously withdrawn from the study (N = 594), or had been lost to follow-up (N = 564). The study obtained ethical approval from Greater Manchester Local Research Ethics Committee and the Scotland Research Ethics committee.

To be included in the present study participants had to have available data on childhood adversity and body mass index (BMI) and cardiometabolic data at ages 60-64 years. Participants classified with BMI ≤ 18.5 kg/m² were also excluded. Of the 2,229 eligible individuals, 1,059 individuals met these criteria. Briefly, of the 2,229 assessed at ages 60-64 years, three did not have age at outcome assessment, 13 had a BMI ≤ 18.5 kg/m², an additional 348 did not have the data to define weight/health status, an additional 349 did not have the data to define childhood SEP adversity, and an additional 457 did not have the data to define childhood psychosocial adversity. Supplementary Table 1 presents differences between the sample (N = 1,059) and individuals excluded from analyses due to missing data (N = 2,229 – 1,059 = 1,170).

**Exposures**

Data on psychosocial and socioeconomic adversities between zero and 15 years of age were used. The majority of data were assessed by maternal or guardian report, augmented by health visitor and teacher reports. Psychosocial adversities included the following: maltreatment (retrospectively assessed at age 43 years), low parental concern for child’s
education, parental psychiatric history, parental divorce, mother affectionless control (retrospectively assessed at age 43 years), father affectionless control (retrospectively assessed at age 43 years), parental death, maternal separation. Socioeconomic adversities included: low maternal education, low father social class, renting a house from a private landlord, poor household amenities, overcrowding, poorly repaired house, unclean child, poorly cleaned house, poor state of child’s clothes and shoes. Supplementary Table 2 provides a more detailed description of these variables. For each individual psychosocial or socioeconomic adversity, the variable was coded as 0 (no adversity) or 1 (presence of adversity). These numbers were then summed together to create two exposures: a composite psychosocial adversity score and a composite socioeconomic adversity score, each of which was further categorised as 0, 1, 2, or ≥ 3 adversities. These scores were created in order to capture repeated exposure to stressful events, as has been previously used by Caleythrop et al.[19]

Outcomes
At ages 60-64 years, weight (kg) and height (m) were measured. Diastolic and systolic blood pressures (mmHg) were assessed and a fasting blood sample was taken to assay glycosylated haemoglobin (HbA1c), high-density lipoprotein-cholesterol (HDL-C) (mmol/L), and triglycerides (mmol/L). Use of antihypertensives, antidiabetics, lipid-lowering drugs, and insulin was self-reported. Detailed description of the methods used to collected these data is provided by Kuh et al.[18]

Overweight or obesity was defined as a BMI ≥ 25 kg/m²; these groups were combined for sample size reasons. Individuals were classified as being metabolically unhealthy if they had two or more of the following complications: 1) impaired glucose metabolism, defined as HbA1C ≥ 5.7%, or use of antidiabetic drugs or insulin;[20] 2) elevated blood pressure, defined as systolic blood pressure ≥ 130 mmHg and diastolic blood pressure ≥ 85 mmHg,[21] or use of antihypertensive medications; 3) low HDL-C, based on a cut-off of 1.03
mmol/L in men and 1.29 mmol/L in women.[21] Hypertriglyceridemia, defined as triglyceride level ≥ 1.70 mmol/L.[21] or use of lipid-lowering drugs, the majority of which are known to reduce triglycerides levels by 50-60%.[22]

Individuals were categorised as either MHO, MUO, MHNW, or MUNW.

Covariates

Age in months at the 60-64 years of age assessment was recorded, as was sex at birth.

Statistical analysis

Differences between the four weight/health status groups were examined using chi-square for categorical variables and ANOVA for continuous variables (log transformed if not normally distributed). The psychosocial adversity score and socioeconomic adversity score were tabulated and their relationship tested using chi-square.

Firstly, separate associations of the psychosocial adversity score and the socioeconomic adversity score with both a binary health status outcome (unhealthy vs healthy) and a binary weight status outcome (overweight/obese vs normal weight) were examined using logistic regression, with adjustment for age and sex. Secondly, associations of the psychosocial adversity score and the socioeconomic adversity score with the composite weight/health status outcome were examined using multinomial logistic regression, with adjustment for age and sex. Each adversity score was examined separately, before putting them together in a mutually adjusted model. Post-estimation commands were used to obtain the relative risks of MUO compared to MHO and of MUNW compared to MHNW. P-values for trends (i.e., linear effect across all categories of an exposure) were obtained by entering the exposures into the models as continuous variables.
In order to test whether our results were biased by complete case analysis, we re-ran the final mutually adjusted model on five multiply imputed datasets. We imputed back up to the 1,865 individuals who were missing data on the exposures but not the outcome. Briefly, we used chained equations to impute the two adversity scores based on 10 individual adversities (with no or minimal missing data), age, sex, weight/health status and all of its components (e.g., systolic blood pressure and HDL-C), and an overall social class variable from ages 26, 36, and 43 years. The analytical model was applied to the multiple-imputed data, using Rubin’s rules to combine estimates across the five datasets.

Because BMI is a limited indicator of adiposity,[23] a sensitivity analysis was conducted in which weight status was defined according to percentage body fat (%fat) assessed by a Hologic (Bedford, MA, USA) QDR 4,500 dual-energy x-ray absorptiometry scanner, as previously described.[18] Of the 1,059 individuals in the analysis sample, 783 individuals had available DXA data. Using NHANES 1999-2004 cut-offs, overweight or obesity was defined as %fat > 39.99% for females and > 19.00% for males.[24]

All analyses were carried out using Stata MP version 14 (StataCorp LP, College Station, TX, USA).

RESULTS

Table 1 shows descriptive statistics, by weight/health status. 62% of the 312 normal weight individuals were metabolically healthy, whereas only 34% of the 747 overweight or obese individuals were metabolically healthy. The MUO group contained approximately twice the percentage of people with ≥ 3 adversities (psychosocial or socioeconomic) compared to the MHO group. Supplementary Table 3 shows the prevalence of individual adversities by weight/health status.
The psychosocial adversity score and the socioeconomic adversity score demonstrated a positive relationship, with a chi-square test reporting $p = 0.017$ (Table 2).

Table 3 shows the relationship of each adversity score with health status (unhealthy vs healthy) and weight status (overweight/obese vs normal weight) considered separately. While the psychosocial adversity score and the socioeconomic adversity score were both related to increased risk of being unhealthy, only the socioeconomic adversity score was related to increased risk of being overweight/obese.

Multinomial logistic regression analyses for each exposure considered separately demonstrated associations of the psychosocial adversity score and the socioeconomic adversity score with increased risk of MUO compared to MHO (Table 4). For example, individuals with a psychosocial adversity score of $\geq 3$ (compared to 0) had a 2.01 (95% CI 1.04, 3.88) times higher risk of MUO than MHO. Similarly, individuals with a socioeconomic adversity score of $\geq 3$ (compared to 0) had a 1.65 (1.01, 2.67) times higher risk of MUO than MHO. Among normal weight individuals, however, only psychosocial adversity was related to increased risk of being metabolically unhealthy.

In a mutually adjusted model, estimates were only slightly attenuated (Table 5). Individuals with a psychosocial adversity score of $\geq 3$ (compared to 0) had a 1.87 (0.96, 3.61) times higher risk of MUO than MHO; individuals with a socioeconomic adversity score of $\geq 3$ (compared to 0) had a 1.60 (0.98, 2.60) times higher risk of MUO than MHO; and again only psychosocial adversity was related to increased risk of being metabolically unhealthy among normal weight individuals. Further, the $p$-values for trends provided evidence that increasing psychosocial adversity had a linear relationship with greater risk of MUNW (compared to MHNW) and that increasing socioeconomic adversity had a linear relationship with greater risk of MUO (compared to MHO). Similar results were obtained using multiple imputation to increase the sample size (Supplementary Table 4).
When outcome groups were defined using %fat, the associations for psychosocial adversity score were generally of a similar magnitude but with slightly wider confidence intervals (Table 6). The relationship of a socioeconomic adversity score of \( \geq 3 \) with MUO (compared to MHO) was, however, attenuated more drastically from a 60% increased risk to only a 22% increased risk when weight status was defined according to %fat.

**DISCUSSION**

This study found that, independently of socioeconomic adversity, children exposed to the highest level of psychosocial adversity had increased risk of being metabolically unhealthy at ages 60-64 years in both normal weight and overweight/obese groups. Conversely, however, the highest level of socioeconomic adversity was only associated with increased risk of being metabolically unhealthy in the overweight/obese group.

Socioeconomic adversity and ACEs are known to be related to increased risk of diseases like cardiovascular disease, cancer, respiratory disease, and type two diabetes.[7, 10, 12, 25, 26] The associations of these exposures with sub-clinical outcomes such as an adverse cardiometabolic profile are equivocal and necessary to investigate further, in order to better understand the potential mechanisms.[14, 15, 27, 28] Also using data from the NSHD, and a nearly identical psychosocial adversity score (0, 1, 2, or 3+), Anderson et al did not find consistent evidence of any relationship with mean-levels of continuous cardiometabolic disease risk marker outcomes (e.g., blood pressures, cholesterols, triglycerides, glucose, insulin, c-reactive protein, carotid intima media thickness, and pulse wave velocity).[14] Similarly, Anderson et al found no evidence of associations in a sample of females aged 51 years at follow-up in the Avon Longitudinal Study of Parents and Children (ALSPAC).[15] The results of these two studies may have been null for a number of reasons, one being that adversity may have greater effect at higher parts of a risk factor distribution than at the mean.
(e.g., adversity might be related to higher systolic blood pressure among a sub-group of individuals who are hypertensive, while having no or minimal effect on the mean in the full sample). By using a composite cardiometabolic health outcome, which identifies individuals with multiple risk factors, we were able to reveal some of the possible long-term consequences of early-life adversity. This finding is in agreement with the Danese et al study, which founds relationships between socioeconomic adversity and social isolation with composite cardiometabolic health status at age 32 years in a sample of 1,037 normal weight, overweight, and obese individuals in Dunedin, New Zealand.[16]

Multiple mechanisms may explain the associations observed in the present study. The relationship between both adversity scores and poor cardiometabolic health status in the overweight/obese group could operate via greater life-time exposure to BMI (e.g., earlier-onset of overweight/obesity), because socioeconomic adversity and ACEs are related to unhealthier BMI trajectories.[29, 30] Similarly, socioeconomic adversity and ACEs are both related to the adoption of unhealthy lifestyle behaviours, which could partly mediate the associations we observed (and themselves operate via deleterious BMI trajectories).[10, 12, 31] There is, however, some evidence that ACEs are related to poor cardiometabolic health independently of intervening behaviours.[16] ACEs and socioeconomic adversity are proxies of stress and extreme, repetitive exposure may potentially dysregulate the stress response.[32] Early-life adversity has been linked to abnormal cortisol profiles and inflammation,[33, 34] both of which are known to be related to, for example, accumulation of visceral fat, impaired fasting glucose, and cardiovascular disease.[35-37]

We observed that socioeconomic adversity was only related to cardiometabolic health status in the overweight/obese group. Socioeconomic adversity tracks across the life course, but because poor health outcomes are observed in children who experience socioeconomic adversity irrespective of their later life socioeconomic environment and conditions, this finding is probably only partly explained by socioeconomic adversity later in adulthood.[38]
This finding also raises the question of whether the two adversities are working via different pathways, one via and one not via obesity. For example, socioeconomic adversity may be operating through a conventional series of events in which adversity leads to unhealthy behaviours which leads to obesity which leads to poor cardiometabolic health. Conversely, psychosocial adversity may be operating more through biological programming of the stress response in childhood, which leads to poor cardiometabolic health regardless of weight status. This is supported by the fact that we found no association between psychosocial adversity and overweight/obesity in our sample.

A recent survey in a healthcare setting in the UK found that 35% of adults have experienced two or more ACEs.\textsuperscript{[39]} Our finding that such individuals may have elevated cardiometabolic disease risk is further evidence on the needs to safeguard children and protect them against childhood adversity. Further, to mitigate the effects of childhood adversity, healthcare systems could screen children at risk for ACEs via questionnaires, as is being done in the USA. This could allow for stratification of treatment and intervention programmes to provide the best possible health outcomes, and importantly mitigate the cardiometabolic risks associated with carrying excess adiposity.

The strengths of this study include the utilisation of a large prospective cohort study with follow-up spanning 64 years, most adversities exposures were prospectively collected therefore limiting recall bias, and we were able to re-define weight status groups using DXA %fat, thereby overcoming a major criticism of the MHO literature. We only adjusted analyses for assessment age and sex to allow comparison to the Anderson et al \textsuperscript{[14]} paper and because the NSHD has no other measures that we think would have strong causal effects on both our exposures and outcome, and thus meet the requirements to be considered a covariate. Nonetheless, residual confounding is a possibility. Other limitations include small numbers in some sub-groups (e.g., MHNW individuals with 3+ psychosocial adversities), no psychometric evaluation (i.e., reliability and validity) of our adversity scores which included
maternal-reported and participant-reported retrospective data, and potential underreporting of maltreatment due to its sensitive nature. Retrospective measures of adversity, such as maltreatment, can suffer from low agreement with prospective measures,[40] which might have increased measurement error in our composite adversity scores and resulted in regression dilution bias.[41] Due to sample attrition and eligibility, our analysis sample did not include the full original birth cohort, which may limit generalisability to the population of Great Britain. The meaning and cultural understanding of childhood adversity in the NSHD cohort born in 1946 will be different from contemporary cohorts. Analyses in modern-day cohorts will, therefore, likely be based upon different adversity measures/scores and potentially find different health consequences compared to those reported in the present paper.

CONCLUSION

Independently of socioeconomic adversity, psychosocial adversity in childhood may be associated with a poor cardiometabolic health profile, in both normal weight and overweight/obese adults.
REFERENCES


