Are dietary patterns in early childhood associated with alcohol consumption at the age of 17 years? Analysis of data from the Avon Longitudinal Study of Parents and Children (ALSPAC) prospective cohort study

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Abstract
Objective: To examine the relationship between a posteriori dietary patterns in early childhood and alcohol consumption in adolescence.

Design: Data were obtained from the Avon Longitudinal Study of Parents and Children (ALSPAC) prospective cohort study. Dietary information was obtained using FFQ at the age of 3 and 7 years. The association between dietary patterns, derived using principal components analysis and the Alcohol Use Disorders Identification Test (AUDIT) scores (to assess harmful intake) and frequency of alcohol consumption at the age of 17 years were examined. Secondary analysis considered sugar intake as a percentage of total energy intake.

Setting: Women who gave birth between 1 April 1991 and 31 December 1992 in the Avon area in southwest England were eligible for the ALSPAC cohort study. Participants: Totally, 14 541 pregnancies were enrolled in ALSPAC during its initial recruitment phase. For this analysis, complete data were available for between 3148 and 3520 participants.

Results: Adherence to the ‘healthy’ dietary pattern at both 3 and 7 years of age was positively associated with consuming more than one alcoholic drink per week at 17 years of age, whilst adherence to the ‘traditional’ dietary pattern at both ages was protective of harmful alcohol intake at 17 years of age. Sugar intake was not associated with either alcohol outcome after adjustment for ethnicity, maternal level of education, parental social class and maternal AUDIT score.

Conclusions: For the population studied, changes to diet in early childhood are unlikely to have an impact on harmful alcohol use in adolescence given the lack of consistency across the results.

Keywords
ALSPAC
Principal component analysis
Adolescence
Alcohol
Sugar

Alcohol use is a leading cause of mortality and morbidity worldwide(1–3). The 2016 Global Burden of Disease identified alcohol as the leading risk factor in male and female deaths aged 15–49 years in 2016, and the seventh leading risk factor for all ages for deaths and disability-adjusted life years(4). Alcohol use also places a high burden on healthcare systems, costing the National Health Service (NHS) in the UK an estimated £3·5 billion per year(4,5). Alcohol consumption most commonly begins in adolescence in countries with a high prevalence of alcohol use(6,7). A recent study in the UK identified that 90% of young people within their study population had their first taste of alcohol by the age of 17 years(6). In addition, it has been shown that young people who binge drink in adolescence are more likely to report binge drinking as young adults(8) and to be exposed to greater alcohol harm as they grow older(7,9). Understanding the factors that contribute to higher alcohol consumption in the adolescent population is a key part of longer-term strategies for reducing population level alcohol harm in adults(8). Identifying sensitive periods where early intervention may have an impact is important for the development of appropriate interventions both in childhood and in adolescence(8,10). There is no single contributory factor to a young person’s propensity to over-consume alcohol(7). However, each of the factors that do contribute merit attention as they could
Children dietary patterns and adolescent alcohol intake prove to be influential in identifying points for intervention.

There is evidence to suggest that sugar and alcohol addiction may be related\(^{(11)}\). Studies in animals have identified that higher consumption of sugar, and possibly fat, may mimic some properties of addictive substances\(^{(12,13)}\). In particular, binging of sugar and fat becomes more pronounced in rodents the longer that they are exposed to increased levels of sugar and fat (i.e. levels in excess of what would normally be fed in a laboratory environment) on a regular basis\(^{(13)}\). Animal studies have also shown that exposing rats to cycles of increased sugar in the diet followed by complete withdrawal of sugar causes a similar neurochemical response to opiate withdrawal\(^{(12,14)}\), suggesting that sugar may be addictive. Understanding how this research may apply to humans is complex. Firstly, if certain foods are found to be addictive, whether a human becomes addicted to them is likely to be multifactorial\(^{(10,15)}\). In addition, linking this to other addictive substances or behaviours, such as patterns of alcohol use or addiction, adds further complexity and identifying causality is very challenging. Some addictive behaviours have been studied alongside diet and, for example, it has been identified that people with high scores on the Yale–Brown Obsessive Compulsive Scale – modified for Pathological Gambling (YPG-YBOCS) – also had higher total fat intake than others included in the study\(^{(16)}\). Biological children of parents with alcohol dependence are more likely to over-consume sugar, which indicates an association in the opposite direction\(^{(12)}\).

Recent research has identified that dietary intake in childhood may be a contributing factor to alcohol use in adolescence. A 2018 study by Mehlig et al.\(^{(10)}\) using data, the Identification and prevention of Dietary-and lifestyle-induced health Effects in Children and infantS (IDEFICS)\(^{(10)}\), showed that children with a high propensity to consume sugar and fat were at 2.46 times greater risk (RR: 2.46; (95% CI 1.47, 4.12)) of alcohol consumption in adolescence compared to those with low propensity, after adjustment for confounders\(^{(10)}\).

The literature on children’s dietary intake and later alcohol consumption is limited. To contribute to research in this area, we used data from the Avon Longitudinal Study of Parents and Children (ALSPAC), a prospective UK cohort study. Given the fact that nutrients are not consumed in isolation\(^{(17)}\), we wanted to examine the association between dietary patterns, looking at the diet as a whole in early childhood compared to later, after adjustment for confounders\(^{(10)}\).

Participants

Pregnant women resident in Avon, UK, with expected dates of delivery 1 April 1991 to 31 December 1992 were invited to take part in ALSPAC. The initial number of pregnancies enrolled was 14,541. Of these, there were 14,062 live births and 13,988 children alive at 1 year of age\(^{(18,19)}\). Data were primarily collected using self-completed questionnaires which were administered during pregnancy and at set ages once the child was born\(^{(20,21)}\). The ALSPAC participants included in this study were those who had dietary intake data available at both 38 months and 81 months, as well as alcohol use data at the age of 17 years. Ethical approval for the study was obtained from the ALSPAC Ethics and Law Committee and the Local Research Ethics Committees. Informed consent for the use of data collected via questionnaires and clinics was obtained from participants following the recommendations of the ALSPAC Ethics and Law Committee at the time. Please note that the ALSPAC study website contains details of all the data that is available through a fully searchable data dictionary and variable search tool: http://www.bristol.ac.uk/alspac/researchers/our-data/.

Measurement of exposures

Dietary intake was measured using a FFQ when the children were aged 38 months and 81 months. The main carer was asked to provide information on how regularly their child consumed a wide range of items, including everyday basic foods as well as snacks and drinks (thirty-four items at 38 months and four-one items at 81 months). The FFQ gave the following options for response: (i) never or rarely; (ii) once in 2 weeks; (iii) 1–3 times a week; (iv) 4–7 times a week or (v) more than once a day\(^{(20)}\). Portion sizes were not asked about. For everyday items, such as milk and bread, more detailed questions were asked, for example, the number of slices of bread per d and the type of bread\(^{(20,22)}\). Dietary patterns were obtained using principal component analysis (PCA) at each time point\(^{(20–22)}\). In brief, frequency of consumption options were converted to times per week as follows: (i) 0; (ii) 0-5; (iii) 2; (iv) 5-5 and (v) 10 times per week. All items were standardised by subtracting the mean and dividing by the standard deviation for each variable. The data were then entered into a PCA with a Varimax rotation\(^{(23,24)}\), and the number of components that best represented the data were chosen using a scree plot. A score for each child was calculated for each component identified at each time point, each score has a mean of 0 and a standard deviation of 1 in the population in which...
it was derived and a higher score indicates a higher adherence to that dietary pattern.

Non-milk extrinsic sugar (NMES) (free sugars in the diet not including milk) intake was estimated from the FFQ as grams per week based on the fifth edition of McCance & Widdowson’s The Composition of Foods[25]. The estimated intake was adjusted to account for overall energy intake for each participant and is presented as the percentage of overall energy intake. We have previously shown that estimated sugar intake was strongly correlated with the ‘processed’ pattern at 3 (previously labelled ‘junk’) and 7 years of age[22] (r = 0.475 and 0.637, respectively).

Measurement of outcomes
There were two outcome measures used in the study. Alcohol consumption was measured at the age of 17 years using a computerised questionnaire, which was completed by the young person in a clinic setting. The questionnaire included the Alcohol Use Disorders Identification Test (AUDIT)[26] and AUDIT score was used as the first outcome measure. Clinically, a score of 8 or more in the AUDIT test is considered to indicate an increased risk of harm from alcohol[26,27], and we therefore chose to use this dichotomy.

The second outcome was a measure of the frequency of alcohol consumption and was based on responses to a question that asked how often the young person had a drink containing alcohol. The options were never, monthly or less, 2 or 4 times a month, and 2 or 3 times a week or 4 or more times a week. We collapsed the five categories into ‘at least weekly’ (‘2 or 3 times a week’ and ‘4 or more times a week’) compared with less than weekly: (‘never’, ‘monthly or less’ and ‘2 or 4 times a month’) in order to facilitate comparison with the IDEFICS study[10].

Statistical methods
STATA v. 15.1 was used to perform logistic regression to assess the relationship with each of the exposure variables and both outcomes. OR and 95% CI are presented. The exposure variables were categorised into quintiles due to our interest in the extremes of each dietary pattern. This also enhanced interpretation of any associations (i.e. making comparisons to the lowest quintile rather than a unitless continuous variable).

A Direct Acyclic Graph (DAG) diagram was used to identify potential confounders from the available data. Relevant literature was consulted while developing the DAG to inform the decision on which variables could be potential confounders and should be included. For example, alcohol harm[28] and poor nutrition[29] are more prevalent in deprived communities and therefore social class was included as a potential confounding variable. Based on this approach, four confounding factors were included in the adjusted analysis which were the child’s ethnicity, mother’s highest educational qualification, household social class and the mother’s AUDIT score. There is evidence that there is a difference in the reporting of levels of alcohol consumption between girls and boys, and therefore an interaction test for gender was completed[10,30].

Multiple imputation by chained equation was used to impute missing data with the use of the ice command in STATA[31]. Twenty-five datasets were generated and ten switching procedures were undertaken. The results can be seen in Supplemental Tables A–D. The following variables were used to impute all outcomes, main predictors, and other variables included in the adjusted analyses plus the Family Adversity Index score, which is comprised of early parenthood, housing (adequacy, basic living facilities (e.g. hot water) and defects), financial difficulties, partner (present/absent), relationship with partner (affection/cruelty/support), family size, major family problems (child in care, not with natural mother or on at risk register), maternal depression/anxiety, substance abuse and crime (trouble with police or convictions). The analyses of associations between dietary patterns and sugar intake with AUDIT and number of drinks consumed were then repeated with the imputed data.

Results
PCA was completed on all participants with dietary data at each time point (n 10139[20] and n 8286[21] at 38 and 81 months, respectively). AUDIT score data were available for 4148 at the age of 17 years. Scores ranged from 0 to 40 with a median of 6 and a mean of 7 (sd 4.8). Alcohol consumption data were available for 3969 young people at the age of 17 years. Data on both dietary intake and alcohol consumption were available for between 3148 and 3520 of the original cohort in the unadjusted analysis.

PCA identified four major dietary components from the FFQ at 38 months, which accounted for 23.5% of the variation within the sample[20]. Each child was given a score for the four components to indicate their predominant dietary pattern or patterns[20]. The four components identified were ‘processed’ (note that in previous papers this was referred to as ‘junk’, high positive loadings for snack foods and foods high in fat such as crisps, sweets, biscuits, chocolate, sausages, burgers, chips and takeaway foods), ‘healthy’ (high positive loadings for vegetables, fruit, rice, pasta, pulses and meat substitutes), ‘traditional’ (high positive loadings for meats, poultry, potatoes and vegetables) and ‘snack’ (high positive loadings for finger foods such as bread, fruit, biscuits, crisps and cheese). Analysis at 81 months identified three dietary patterns, ‘processed’, ‘healthy’ and ‘traditional’, which had very similar loadings for foods as the 3-year patterns and explained 18.2% of variation within the sample[21]. The data available on NMES intake was adjusted to account for overall energy intake for each participant and is presented as a percentage of total energy intake. The percentages for NMES ranged...
Childhood dietary patterns and adolescent alcohol intake

from 0.6% to 39.8% at 38 months, and 1.1% to 40.3% at 81 months.

Associations were evident between each of the confounders considered and at least one of the outcomes (see Table 1). An AUDIT score $\geq 8$ was more likely in children of White ethnicity and those whose mothers also reported an AUDIT score $\geq 8$. Adolescents who reported consuming more than one drink per week were more likely to have lower educated mothers, be from White backgrounds and have mothers with a high AUDIT score.

There were no differences in mean dietary pattern scores at 3 or 7 years of age with AUDIT score. However, children consuming more than one alcoholic drink per week had lower mean scores for the ‘processed’ dietary pattern at 3 and higher mean scores for the ‘healthy’ dietary pattern at both 3 and 7. Sugar intake was slightly higher in those children with an AUDIT score $\geq 8$ (although this was $< 0.5\%$), but there was no difference in mean sugar intake according to frequency of alcohol consumption.

Supplemental Table A outlines the baseline differences in participants who did and did not provide alcohol consumption data at the age of 17 years. Mean maternal age at delivery was 1.3 years higher on average in mothers of participants who provided alcohol data. Participants who provided alcohol data were also more likely to have mothers with a higher level of education and to be from a higher social class, as well as less likely to have mothers with an AUDIT score $\geq 8$. Following multiple imputation, the descriptive characteristics of the participants who did and did not have alcohol data were similar to the non-imputed data (see online Supplemental Table A).

**Dietary patterns and harmful alcohol consumption**

Table 2 presents the associations between dietary pattern scores at 3 and 7 years of age and AUDIT scores $\geq 8$ (i.e. harmful alcohol consumption). Children in the highest quintile for the 3-year ‘processed’ pattern were more likely to have harmful alcohol drinking compared to those in the lowest quintile (unadjusted OR: $1.41$ (95% CI $1.13$, $1.77$)). This association was attenuated after adjustment (OR: $1.35$ (95% CI $0.96$, $1.90$)). A linear association was also evident with an unadjusted OR of $1.09$ (95% CI $1.00$, $1.17$), but again this was attenuated after adjustment (OR: $1.06$ (95% CI $0.94$, $1.19$)). There were no associations evident between the ‘processed’ pattern at 7 years and harmful alcohol drinking. For the ‘healthy’ dietary pattern, there was a linear adjusted effect at the age of 7 years (OR: $1.10$ (95% CI $1.00$, $1.21$)) but not at 3 years of age. The ‘traditional’ pattern showed a protective linear adjusted effect at the age of 3 years (OR: $0.90$ (95% CI $0.82$, $1.00$)) and this was stronger at the age of 7 years (OR: $0.87$ (95% CI $0.70$, $0.96$)). In addition, being in the highest quintile of for the ‘traditional’ pattern was associated with a reduced risk of harmful alcohol consumption (OR of $0.71$ (95% CI $0.53$, $0.96$) and $0.64$ (95% CI $0.47$, $0.87$) at the age of 3 and 7 years, respectively). The 3-year ‘snack’ pattern was not associated with harmful alcohol consumption. In the imputed analyses presented in Supplemental Table B, similar patterns of association were seen across all dietary patterns, although associations tended to be weaker. This was particularly the case for the ‘traditional’ pattern at the age of 7 years (adjusted OR: $0.94$ (95% CI $0.88$, $1.00$)).

**Dietary patterns and frequent alcohol consumption**

Associations between dietary pattern scores and whether participants consumed more than one alcoholic drink per week at the age of 17 years are presented in Table 3. Being in the highest quintile of the ‘processed’ pattern score at the age of 7 years was protective (OR: $0.70$ (95% CI $0.54$, $0.91$)) compared to being in the lowest quintile; however, this association was lost after adjustment for confounders. Similarly, the unadjusted linear association with the ‘processed’ pattern scores pre-adjustment was no longer evident after adjustment. The 7-year ‘processed’ pattern was not associated with consuming more than one alcoholic drink per week at the age of 17 years. The ‘healthy’ pattern at both 3 and 7 years of age was linearly associated with this outcome after adjustment, although the effects were stronger at 7 years of age compared to 3 years of age (OR: $1.21$ and $1.11$, respectively). There were no associations between the ‘traditional’ dietary pattern at either age or the ‘snacks’ pattern at the age of 3 years and later alcohol consumption at the age of 17 years. All these patterns of association were replicated in the imputed analyses (see online Supplemental Table C). There was evidence of an interaction between gender and the unadjusted 3-year ‘processed’ pattern and the adjusted 7-year ‘traditional’ pattern. Additional stratified analyses were therefore carried out (see online Supplemental Table E). This indicated that boys with the highest ‘processed’ dietary pattern quintile were $0.55$ (95% CI $0.37$, $0.80$) times less likely to consume alcohol, whilst girls in the same quintile had no association with an OR of $0.92$ (95% CI $0.64$, $1.33$). No association was shown for girls or boys separately for the ‘traditional’ pattern.

**Sugar consumption and alcohol consumption**

There was an increased risk of harmful alcohol consumption for those children in the highest quintile of sugar intake at the age of 3 years, but the OR was attenuated after adjustment (Table 4), and there were no associations with sugar intake at the age of 7 years. There were no associations between sugar consumption at either age and having more than one alcoholic drink per week at 17 years of age. In the imputed analysis (see online Supplemental Table E), a similar pattern was seen.
In this study, we report an association between a ‘healthy’ dietary pattern score at both 3 and 7 years of age and consuming more than one alcohol drink per week at the age of 17 years. In addition, the ‘traditional’ dietary pattern at both ages was protective of harmful alcohol intake at the age of 17 years. However, we found no association between sugar intake and either measure of alcohol consumption after adjustment for ethnicity, maternal level of education, parental social class and maternal AUDIT score.

This study adds to a limited and new area of literature. Nutrition in childhood has been shown to impact on a wide range of later health and behavioural outcomes. Given the negative impact that alcohol can have on health, exploration of factors that contribute to increased usage is important. It is reassuring that in this population, a poor diet in childhood (represented by adherence to the ‘processed’ diet) was a risk factor for harmful alcohol intake at the age of 17 years. This is concerning, given the increasing levels of alcohol consumption among young adults in recent years. Further research is needed to understand the mechanisms underlying these associations and to develop effective interventions to promote healthier eating habits among children.
pattern) is not associated with an increased risk of harmful drinking in early adulthood. The fact that adhering to a ‘healthy’ pattern in early childhood is associated with an increased likelihood of consuming more than one alcoholic drink a week may warrant further investigation. However, harmful levels of intake were not associated with this dietary pattern.

We are not aware of any other studies examining the association between dietary patterns in childhood and alcohol consumption in adolescence. However, other

| Exposure                                      | Unadjusted OR | 95 % CI | P value | Adjusted OR | 95 % CI | P value |
|-----------------------------------------------|---------------|---------|---------|-------------|---------|---------|
| ‘Processed’ diet pattern – 3 years of age     | 1393/3520     | 764/1935|         |             |         |         |
| Quintile 1 (baseline – lowest ‘processed’)    | 1             | 1       |         |             |         |         |
| Quintile 2                                   | 1·32          | 1·08, 1·60| 0·006   | 1·26        | 0·98, 1·62| 0·077   |
| Quintile 3                                   | 1·21          | 0·98, 1·48| 0·070   | 1·12        | 0·85, 1·46| 0·431   |
| Quintile 4                                   | 1·10          | 0·89, 1·36| 0·399   | 0·96        | 0·71, 1·30| 0·776   |
| Quintile 5 (highest ‘processed’)             | 1·41          | 1·13, 1·77| 0·003   | 1·35        | 0·96, 1·90| 0·063   |
| Linear effect                                | 1·09          | 1·00, 1·17| 0·023   | 1·06        | 0·94, 1·19| 0·353   |
| ‘Processed’ diet pattern – 7 years of age     | 1323/3311     | 759/1905|         |             |         |         |
| Quintile 1 (baseline – lowest ‘processed’)    | 1             | 1       |         |             |         |         |
| Quintile 2                                   | 1·15          | 0·93, 1·41| 0·193   | 1·17        | 0·90, 1·53| 0·237   |
| Quintile 3                                   | 1·05          | 0·85, 1·30| 0·667   | 0·32        | 0·69, 1·22| 0·541   |
| Quintile 4                                   | 1·31          | 1·06, 1·62| 0·013   | 1·25        | 0·94, 1·67| 0·131   |
| Quintile 5 (highest ‘processed’)             | 1·07          | 0·85, 1·35| 0·564   | 1·09        | 0·79, 1·50| 0·618   |
| Linear effect                                | 1·04          | 0·96, 1·12| 0·336   | 1·03        | 0·92, 1·15| 0·599   |
| ‘Healthy’ diet pattern – 3 years of age       | 1393/3520     | 764/1935|         |             |         |         |
| Quintile 1 (baseline – lowest ‘healthy’)      | 1             | 1       |         |             |         |         |
| Quintile 2                                   | 1·13          | 0·90, 1·41| 0·296   | 1·26        | 0·92, 1·73| 0·151   |
| Quintile 3                                   | 1·07          | 0·86, 1·34| 0·526   | 1·27        | 0·94, 1·73| 0·122   |
| Quintile 4                                   | 1·08          | 0·87, 1·35| 0·475   | 1·13        | 0·83, 1·55| 0·435   |
| Quintile 5 (highest ‘healthy’)                | 1·18          | 0·95, 1·47| 0·129   | 1·32        | 0·97, 1·80| 0·079   |
| Linear effect                                | 1·04          | 0·97, 1·11| 0·253   | 1·08        | 0·98, 1·19| 0·143   |
| ‘Healthy’ diet pattern – 7 years of age       | 1323/3311     | 759/1905|         |             |         |         |
| Quintile 1 (baseline – lowest ‘healthy’)      | 1             | 1       |         |             |         |         |
| Quintile 2                                   | 0·98          | 0·78, 1·23| 0·870   | 0·99        | 0·72, 1·35| 0·930   |
| Quintile 3                                   | 0·74          | 0·59, 0·93| 0·010   | 0·79        | 0·58, 1·08| 0·142   |
| Quintile 4                                   | 0·85          | 0·68, 1·07| 0·165   | 0·79        | 0·58, 1·08| 0·144   |
| Quintile 5 (highest ‘healthy’)                | 1·09          | 0·87, 1·36| 0·452   | 1·14        | 0·83, 1·55| 0·420   |
| Linear effect                                | 1·04          | 0·97, 1·11| 0·291   | 1·10        | 1·00, 1·21| 0·063   |
| ‘Traditional’ diet pattern – 3 years of age    | 1393/3520     | 764/1935|         |             |         |         |
| Quintile 1 (baseline – lowest ‘traditional’)  | 1             | 1       |         |             |         |         |
| Quintile 2                                   | 0·99          | 0·80, 1·22| 0·892   | 0·78        | 0·59, 1·05| 0·100   |
| Quintile 3                                   | 0·95          | 0·77, 1·18| 0·642   | 0·80        | 0·59, 1·07| 0·137   |
| Quintile 4                                   | 0·87          | 0·70, 1·08| 0·205   | 0·73        | 0·54, 0·97| 0·031   |
| Quintile 5 (highest ‘traditional’)            | 0·89          | 0·72, 1·10| 0·290   | 0·71        | 0·53, 0·96| 0·024   |
| Linear effect                                | 0·95          | 0·89, 1·01| 0·161   | 0·90        | 0·82, 1·00| 0·034   |
| ‘Traditional’ diet pattern – 7 years of age    | 1323/3311     | 759/1905|         |             |         |         |
| Quintile 1 (baseline – lowest ‘traditional’)  | 1             | 1       |         |             |         |         |
| Quintile 2                                   | 0·87          | 0·70, 1·09| 0·223   | 0·78        | 0·58, 1·05| 0·105   |
| Quintile 3                                   | 0·96          | 0·77, 1·20| 0·717   | 0·83        | 0·62, 1·12| 0·220   |
| Quintile 4                                   | 0·92          | 0·74, 1·15| 0·448   | 0·84        | 0·63, 1·13| 0·254   |
| Quintile 5 (highest ‘traditional’)            | 0·75          | 0·60, 0·94| 0·011   | 0·64        | 0·47, 0·87| 0·005   |
| Linear effect                                | 0·94          | 0·87, 1·01| 0·075   | 0·87        | 0·70, 0·96| 0·007   |
| ‘Snack’ diet pattern – 3 years of age         | 1393/3520     | 764/1935|         |             |         |         |
| Quintile 1 (baseline – lowest ‘traditional’)  | 1             | 1       |         |             |         |         |
| Quintile 2                                   | 0·78          | 0·62, 0·99| 0·037   | 0·74        | 0·53, 1·03| 0·073   |
| Quintile 3                                   | 0·81          | 0·64, 1·01| 0·065   | 0·79        | 0·58, 1·09| 0·154   |
| Quintile 4                                   | 0·84          | 0·67, 1·05| 0·123   | 0·81        | 0·59, 1·11| 0·190   |
| Quintile 5 (highest ‘traditional’)            | 0·91          | 0·73, 1·14| 0·407   | 0·87        | 0·64, 1·19| 0·390   |
| Linear effect                                | 0·99          | 0·93, 1·07| 0·986   | 0·97        | 0·88, 1·07| 0·572   |

AUDIT, Alcohol Use Disorders Identification Test.

*Data adjusted for ethnicity, maternal level of education, parental social class and maternal AUDIT score.
studies have identified associations between dietary patterns and alcohol consumption in adulthood which helped to inform our hypothesis. For example, a positive association was identified between heavier alcohol consumption during pregnancy and scoring highly on the ‘processed’ dietary pattern among mothers in the ALSPAC cohort\(^34\). The FinDrink project in Finland identified mixed results from a study of the relationship between alcohol consumption and dietary patterns\(^35\), and they identified that moderate alcohol consumption (compared to non-drinkers) was identified with higher fish intake but also higher energy intake from total fats and monosaturated fats. A key conclusion from the FinDrink study was that further study into the relationship between alcohol consumption and dietary habits is needed.

### Table 3

Association between dietary patterns at ages 3 and 7 years, and consumption of more than one drink per week at the age of 17 years; associations with quintiles of dietary pattern score and continuous pattern score

| Exposure | Unadjusted OR | 95% CI | P value | Adjusted OR | 95% CI | P value |
|----------|---------------|--------|---------|-------------|--------|---------|
| 'Processed' diet pattern – 3 years of age | | | | | | |
| Number of cases/total | 860/3371 | 492/1856 |
| Quintile 1 (baseline – lowest 'processed' pattern) | 1 | 1 |
| Quintile 2 | 0.90 | 0.72, 1.12 | 0.340 | 0.83 | 0.62, 1.10 | 0.199 |
| Quintile 3 | 0.97 | 0.78, 1.22 | 0.805 | 1.00 | 0.75, 1.35 | 0.476 |
| Quintile 4 | 0.74 | 0.58, 0.95 | 0.018 | 0.72 | 0.50, 1.02 | 0.061 |
| Quintile 5 (highest 'processed' pattern) | 0.70 | 0.54, 0.91 | 0.009 | 0.90 | 0.61, 1.32 | 0.597 |
| Linear effect | 0.89 | 0.82, 0.97 | 0.009 | 0.94 | 0.82, 1.08 | 0.376 |
| 'Processed' diet pattern – 7 years of age | | | | | | |
| Number of cases/total | 809/3175 | 485/1832 |
| Quintile 1 (baseline – lowest 'processed' pattern) | 1 | 1 |
| Quintile 2 | 0.94 | 0.75, 1.19 | 0.621 | 0.83 | 0.61, 1.20 | 0.211 |
| Quintile 3 | 0.92 | 0.72, 1.17 | 0.508 | 0.86 | 0.62, 1.17 | 0.332 |
| Quintile 4 | 0.99 | 0.78, 1.27 | 0.965 | 1.08 | 0.79, 1.48 | 0.638 |
| Quintile 5 (highest 'processed' pattern) | 0.80 | 0.61, 1.05 | 0.102 | 0.76 | 0.52, 1.09 | 0.147 |
| Linear effect | 0.97 | 0.89, 1.06 | 0.494 | 1.00 | 0.89, 1.14 | 0.960 |
| 'Healthy' diet pattern – 3 years of age | | | | | | |
| Number of cases/total | 860/3371 | 492/1856 |
| Quintile 1 (baseline – lowest 'healthy' pattern) | 1 | 1 |
| Quintile 2 | 0.87 | 0.67, 1.14 | 0.308 | 0.80 | 0.55, 1.15 | 0.221 |
| Quintile 3 | 1.09 | 0.84, 1.41 | 0.520 | 0.83 | 0.59, 1.18 | 0.311 |
| Quintile 4 | 1.21 | 0.94, 1.56 | 0.137 | 1.05 | 0.75, 1.49 | 0.768 |
| Quintile 5 (highest 'healthy' pattern) | 1.37 | 1.07, 1.75 | 0.013 | 1.13 | 0.80, 1.58 | 0.494 |
| Linear effect | 1.15 | 1.06, 1.23 <0.001 | 1.11 | 1.00, 1.24 | 0.056 |
| 'Healthy' diet pattern – 7 years of age | | | | | | |
| Number of cases/total | 809/3175 | 485/1832 |
| Quintile 1 (baseline – lowest 'healthy' pattern) | 1 | 1 |
| Quintile 2 | 1.03 | 0.78, 1.34 | 0.854 | 0.96 | 0.67, 1.37 | 0.813 |
| Quintile 3 | 0.82 | 0.62, 1.08 | 0.156 | 0.66 | 0.45, 0.96 | 0.028 |
| Quintile 4 | 1.13 | 0.87, 1.47 | 0.352 | 0.95 | 0.66, 1.35 | 0.763 |
| Quintile 5 (highest 'healthy' pattern) | 1.52 | 1.18, 1.96 | 0.001 | 1.35 | 0.95, 1.91 | 0.009 |
| Linear effect | 1.19 | 1.10, 1.29 <0.001 | 1.21 | 1.09, 1.35 | 0.001 |
| 'Traditional' diet pattern – 3 years of age | | | | | | |
| Number of cases/total | 860/3371 | 492/1856 |
| Quintile 1 (baseline – lowest 'traditional' pattern) | 1 | 1 |
| Quintile 2 | 1.21 | 0.95, 1.57 | 0.120 | 1.10 | 0.79, 1.52 | 0.576 |
| Quintile 3 | 0.97 | 0.75, 1.25 | 0.811 | 0.95 | 0.68, 1.33 | 0.776 |
| Quintile 4 | 1.05 | 0.82, 1.34 | 0.718 | 0.97 | 0.70, 1.35 | 0.674 |
| Quintile 5 (highest 'traditional' pattern) | 1.02 | 0.80, 1.30 | 0.884 | 0.88 | 0.63, 1.24 | 0.469 |
| Linear effect | 1.00 | 0.92, 1.08 | 0.946 | 0.95 | 0.85, 1.05 | 0.312 |
| 'Traditional' diet pattern – 7 years of age | | | | | | |
| Number of cases/total | 809/3175 | 485/1832 |
| Quintile 1 (baseline – lowest 'traditional' pattern) | 1 | 1 |
| Quintile 2 | 0.89 | 0.69, 1.15 | 0.384 | 0.94 | 0.67, 1.32 | 0.736 |
| Quintile 3 | 1.00 | 0.78, 1.29 | 0.979 | 1.02 | 0.73, 1.43 | 0.899 |
| Quintile 4 | 0.93 | 0.72, 1.20 | 0.598 | 0.97 | 0.70, 1.36 | 0.880 |
| Quintile 5 (highest 'traditional' pattern) | 0.89 | 0.69, 1.15 | 0.375 | 0.89 | 0.63, 1.26 | 0.506 |
| Linear effect | 0.97 | 0.89, 1.06 | 0.484 | 0.98 | 0.87, 1.09 | 0.663 |
| 'Snack' diet pattern – 3 years of age | | | | | | |
| Number of cases/total | 1393/3520 | 764/1935 |
| Quintile 1 (baseline – lowest 'traditional' pattern) | 1 | 1 |
| Quintile 2 | 0.85 | 0.65, 1.10 | 0.205 | 0.77 | 0.53, 1.21 | 0.169 |
| Quintile 3 | 0.83 | 0.64, 1.08 | 0.160 | 0.88 | 0.62, 1.26 | 0.481 |
| Quintile 4 | 0.90 | 0.70, 1.16 | 0.424 | 0.82 | 0.58, 1.17 | 0.275 |
| Quintile 5 (highest 'traditional' pattern) | 1.04 | 0.81, 1.34 | 0.765 | 1.02 | 0.71, 1.45 | 0.928 |
| Linear effect | 1.03 | 0.95, 1.11 | 0.450 | 1.01 | 0.91, 1.14 | 0.804 |
The study by Mehlig et al. also informed our hypothesis, and specifically our interest in examining the relationship between NMES percentage intake and alcohol consumption. The results from the ALSPAC study are notably different to the IDEFICS study and could be due to multiple factors. Firstly, there are likely to be differences between the demographics, eating habits and alcohol use of the populations from different countries\(^{(12-14)}\). Secondly, the ALSPAC data offered the opportunity to study a larger population than the IDEFICS study. Approximately 25.5% of the adolescents were consuming at least one drink per week and 39.9% had an AUDIT \(\geq 8\), which provided a rich data source for identifying if any associations exist. This contrasts with the study undertaken by Mehlig et al.\(^{(10)}\) which had 107 participants regularly consuming alcohol. This is a small sample size when considering that the results may have been affected as the participants were underage.

The use of AUDIT scores also strengthens our study. AUDIT is a widely used, validated tool\(^{(25)}\), and the overall score offers a broad understanding of a person’s risk of alcohol harm. To add sensitivity to the study, the measure of alcohol frequency was included as an additional outcome variable. This additional measure also had more similarity to the outcome measure in the IDEFICS study, providing an opportunity to better consider our paper in the context of previous findings. Finally, it should be highlighted that the IDEFICS study concluded that both sugar and fat intake were associated with later alcohol consumption. High fat intake was shown to be independently associated with alcohol consumption, but high sugar intake was not\(^{(10)}\). Our findings support this lack of independent association with sugar; however, we did not find any evidence to suggest that increased ‘fat’ intake, using our ‘processed’ pattern as a proxy was associated with alcohol consumption.

As well as the strengths outlined above, this paper has a number of limitations. Some children were lost to years\(^{(10)}\), so it is more unlikely that this cohort would be regularly consuming alcohol\(^{(6)}\). Reporting of alcohol intake may have been affected as the participants were underage. The use of AUDIT scores also strengthens our study. AUDIT is a widely used, validated tool\(^{(25)}\), and the overall score offers a broad understanding of a person’s risk of alcohol harm. To add sensitivity to the study, the measure of alcohol frequency was included as an additional outcome variable. This additional measure also had more similarity to the outcome measure in the IDEFICS study, providing an opportunity to better consider our paper in the context of previous findings. Finally, it should be highlighted that the IDEFICS study concluded that both sugar and fat intake were associated with later alcohol consumption. High fat intake was shown to be independently associated with alcohol consumption, but high sugar intake was not\(^{(10)}\). Our findings support this lack of independent association with sugar; however, we did not find any evidence to suggest that increased ‘fat’ intake, using our ‘processed’ pattern as a proxy was associated with alcohol consumption.

As well as the strengths outlined above, this paper has a number of limitations. Some children were lost to
follow-up, reducing the amount of data available at the later time points. However, the results of the imputation indicated that this did not bias our findings. On the whole, effect sizes in the imputed analyses were smaller than those in the complete case analysis. The loss of participants to follow-up reduces the external validity of the study, because the adolescents included were more likely to be from advantaged backgrounds, more likely to be from a White British background, more likely to have highly educated mothers and less likely to have mothers with an AUDIT \( \geq 8 \).

A further potential limitation is the reliability of self-reported alcohol data amongst adolescents\(^{37,38}\). Williams et al.\(^{57}\) have shown that self-report of substance abuse by adolescents only had fair validity and recommended biochemical corroboration be routinely used for this age group, although Winters et al.\(^{38}\) suggest that adolescents reliably self-report alcohol intake. The data collected on alcohol were part of a much wider clinic visit with many other measures being obtained. Whilst there is still a risk that participants may have under-reported their alcohol consumption, it is less likely in such a setting where multiple facets of an individual are being examined.

The external validity of the study may also be affected by the changes in children’s diets over the last 30 years. The dietary intake data used in the study were collected in the 1990s, and between data collection and the present day, childhood obesity has increased and some elements of childhood diet have changed in association with this\(^{39}\). It is likely that the broad patterns found in the PCA would remain, but some loadings and scores that contributed to the patterns may have strengthened, for example, to accommodate an increase in consumption of sugary drinks\(^{40}\).

Conclusion

In this large prospective cohort study, no association was found between sugar intake in childhood and alcohol consumption in later adolescence, which is in contradiction to previous research on this topic. However, we do report an association between adherence to a ‘healthy’ dietary pattern at both 3 and 7 years of age and consuming more than one drink per week and adherence to a ‘traditional’ dietary pattern intake at the age of 7 years being protective of harmful alcohol intake. We do not know enough from this study alone to know if adherence to a specific dietary pattern in early childhood would affect harm from alcohol. However, the study adds to a field of relatively limited literature and further research is required to elucidate the associations under study. For the population studied, it suggests that changes to diet in early childhood are unlikely to have an impact on harmful alcohol use in adolescence given the lack of consistency across the results.

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Supplementary material

For supplementary material accompanying this paper visit https://doi.org/10.1017/S1368980021004183

References

1. GBD Alcohol Collaborators (2018) Alcohol use and burden for 195 countries and territories, 1990–2016: a systematic analysis for the global burden of disease study 2016. Lancet 392, 1015–1035.
2. Chikritzhs T, Stockwell T, Naimi T et al. (2015) Has the leaning tower of presumed health benefits from ‘moderate’ alcohol use finally collapsed? Addiction 110, 726–727.
3. Bacolod M, Cunha JM, Shen Y-C et al. (2017) The Impact of Alcohol on Mental Health, Physical Fitness, and Job Performance. Cambridge, MA: National Bureau of Economic Research.
4. McQuire C, Tilling K, Hickman M et al. (2019) Forecasting
Childhood dietary patterns and adolescent alcohol intake

the 2021 local burden of population alcohol-related harms using Bayesian structural time-series. *Addiction* **114**, 994–1003.
5. Home Office (2012) A Minimum Unit Price for Alcohol. https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/157763/ia-minimum-unit-pricing.pdf (accessed August 2019).
6. Donghue KPD, Rose H, Boniface SFD et al. (2017) Alcohol consumption, early-onset drinking, and health-related consequences in adolescents presenting at emergency departments in England. *J Adolesc Health* **60**, 438–446.
7. Pitkänen T, Kokko K, Lyyyra A-L et al. (2008) A developmental approach to alcohol drinking behaviour in adulthood: a follow-up study from age 8 to age 42. *Addiction* **103**, 48–68.
8. Degenhardt L, O’Loughlin C, Swift W et al. (2015) The persistence of adolescent binge drinking into adulthood: findings from a 15-year prospective cohort study. *BMJ Open* **3**, e003015.
9. Hingson RW & White AM (2013) *An Easy Guide to Factor Analysis*. Philadelphia, PA: W.B. Saunders.
10. North K & Emmett P (2005) Multivariate analysis of diet in children at 4 and 7 years of age and associations with socio-demographic characteristics. *Eur J Clin Nutr* **59**, 751–760.
11. Northstone K & Emmett P (2000) Multivariate analysis of nutrient intake and gambling: is fat and sugar consumption associated with gambling? *Int J Addict* **35**, 1094–1099.
12. Fortuna J (2010) Sweet preference, sugar addiction and the familial history of alcohol dependence: shared neural pathways and genes. *J Psychoactive Drugs* **42**, 147–151.
13. Avena NM, Rada P & Hoebel BG (2009) Sugar and fat binging have notable differences in addictive-like behavior. *J Nutr* **139**, 623–628.
14. Colantuoni C, Rada P, McCarthy J et al. (2002) Evidence that intermittent, excessive sugar intake causes endogenous opioid dependence. *Obes Res* **10**, 478–488.
15. DiNicolantonio JJ, O’Keefe JH & Wilson WL (2018) Sugar addiction: is it real? A narrative review. *Br J Sports Med* **52**, 910–913.
16. Chamberlain SR, Redden S, Leppink E et al. (2017) Caloric intake and gambling: is fat and sugar consumption ‘impulsive’? *J Gambl Stud* **33**, 783–793.
17. Tapsell LC, Neale EP, Hu FB et al. (2016) Foods, nutrients, and dietary patterns: interconnections and implications for dietary guidelines. *Adv Nutr* **7**, 445–541.
18. Boyd A, Golding J, MacLeod J et al. (2013) Cohort profile: the ‘children of the 90’s’ – the index offspring of the Avon longitudinal study of parents and children. *Int J Epidemiol* **42**, 111–127.
19. Fraser A, Macdonald-Wallis C, Tilling K et al. (2013) Cohort profile: the Avon longitudinal study of parents and children: ALSFAP mothers cohort. *Int J Epidemiol* **42**, 97–110.
20. North K & Emmett P (2000) Multivariate analysis of diet among 5-year-old children and associations with socio-demographic characteristics. The Avon longitudinal study of pregnancy and childhood (ALSFPAC) study team. *Eur J Clin Nutr* **54**, 73–80.
21. Northstone K & Emmett P (2005) Multivariate analysis of diet in children at 4 and 7 years of age and associations with socio-demographic characteristics. *Eur J Clin Nutr* **59**, 751–760.
22. Criibb V, Emmett P & Northstone K (2013) Dietary patterns throughout childhood and associations with nutrient intakes. *Public Health Nutr* **16**, 1801–1809.
23. Gorsuch RL (1974) *Factor Analyses*. Philadelphia, PA: W.B. Saunders.
24. Kline P (1994) *An Easy Guide to Factor Analysis*. London: Routledge.
25. Holland B, Welch AA, Unwin ID et al. (1991) *McCance & Widdowson’s The Composition of Foods*, 5th ed. Cambridge: The Royal Society of Chemistry.
26. Public Health England (2017) Guidance on the 5 Alcohol Use Screening Tests. https://www.gov.uk/government/publications/alcohol-use-screening-tests/guidance-on-the-5-alcohol-use-screening-tests (accessed August 2019).
27. Heron J, Maughan B, Dick DM et al. (2015) Conduct problem trajectories and alcohol use and misuse in mid to late adolescence. *Drug Alcohol Depend* **153**, 100–107.
28. Lewer D, Meier P, Beurd E et al. (2016) Unravelling the alcohol harm paradox: a population-based study of social gradients across very heavy drinking thresholds. *BMC Public Health* **16**, 599.