Association of dietary energy density in childhood with age and body fatness at the onset of the pubertal growth spurt

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Abstract
The aim of the present study was to examine the association of pre-pubertal dietary energy density (ED) with both age and body fatness at the start of the pubertal growth spurt (age at take-off, ATO). Analyses included 219 DOrtmund Nutritional and Anthropometric Longitudinally Designed Study participants with sufficient height measurements to estimate ATO who provided 3 d weighed dietary records at baseline, i.e. 2 and 3 years before ATO (mean age 6·9 (SD 1·2) years). Mean energy intakes and amounts of foods/drinks consumed at baseline were derived from the records. ED (kJ/g) was calculated based on (1) all foods and drinks (ED_all), (2) foods and energy-containing drinks (ED_energy), (3) foods and milk as a drink, but no other beverages (ED_milk) and (4) foods only, solid or liquid (ED_food). Using multiple regression analyses, the association between the ED variables and ATO was investigated. Furthermore, Z-scores of BMI and fat mass index (FMI) at ATO were considered as outcomes to reflect body fatness at puberty onset. The results showed that ED at baseline was not associated with ATO, regardless of the ED method used. For example, mean ATO in the lowest v. highest tertile of ED_food was 9·3 (95 % CI 9·0, 9·5) v. 9·4 (95 % CI 9·1, 9·7) years, PTrend = 0·8 (adjusted for sex, maternal age, birth weight, dietary protein, dietary fibre, baseline BMI Z-score). Similarly, ED was not independently associated with BMI or FMI Z-score at ATO (PTrend = 0·3–0·9). In conclusion, dietary ED in childhood did not influence timing or body fatness at ATO in this cohort of healthy, free-living children.

Key words: Energy density: Puberty: BMI: Children

Earlier puberty represents a risk factor for hormone-related cancers such as breast cancer and has been related to other adverse outcomes such as mortality (1–3). Higher childhood body fatness may result in earlier onset and/or more rapid progression of puberty (4–6). Since higher adiposity can be a result of excess energy intake (EI), one can also assume a role of EI for pubertal timing. Several studies have addressed this question with respect to age at menarche with conflicting results. Higher EI has been associated with both earlier (7) and delayed (8) menarche, yet others have found no association (9,10).

Recently, interest in dietary energy density (ED), i.e. the ratio of the amount of food to total EI, has emerged. According to experimental studies that manipulated ED under laboratory conditions, ED is a major determinant of EI and potentially affects adiposity (11). It has been postulated that increased body fat resulting from excess EI and higher ED in childhood could influence pubertal timing. EI, irrespective of adiposity, may also be a regulator of childhood growth (velocity) and associated hormones, which have been related to pubertal timing and/or cancer risk (12,13). In addition to these mechanisms, higher ED can be associated with lower diet quality and hence nutrient intake. Higher intakes of thiamin, Fe (14) and fibre (15) have been related to delayed pubertal onset in girls. Children with lower diet quality, indexed by the Nutritional Quality Index, indeed experienced their pubertal growth spurt (age at take-off, ATO) earlier (16).

However, data on ED in children are limited, particularly if they choose their diet spontaneously and consume foods

[Abbreviations: ATO, age at take-off; DONALD, DOrtmund Nutritional and Anthropometric Longitudinally Designed; ED, dietary energy density; ED_all, all foods and drinks; ED_energy, foods and energy-containing drinks; ED_food, foods only, solid or liquid; ED_milk, foods and milk as a drink, but no other beverages; EI, energy intake; FMI, fat mass index.

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ad libitum. Its role on pubertal timing has not previously been addressed. The few observations on ED and body fat have yielded mixed results\(^{(17-19)}\). In part, this may be explained by different methods of calculating ED and including beverages\(^{(19)}\). The aim of the present study was to examine the association between ED, body fat and ATO in the DOrtmund Nutritional and Anthropometric Longitudinally Designed (DONALD) Study participants. ED was calculated using four different approaches.

### Methods

The DONALD Study is an ongoing, open cohort study started in 1985\(^{(20)}\). Information on diet, growth and metabolism between infancy and adulthood is regularly collected: four visits in the first year of life; two in the second; then one annually. The Ethics Committee of the University of Bonn, Germany, approved the study. Examinations are performed with parental consent.

ATO could be estimated from repeated height measurements in 376 term singletons with a birth weight >2500 g\(^{(21)}\). Of these, 219 had anthropometry measured, plausible dietary data 2 and 3 years before ATO, and information on potential confounders (maternal overweight/education, breast-feeding and baseline anthropometry). The time points represent mean age at ‘adiposity rebound’, a critical period in childhood. This sample size of 219 was sufficient to detect mean differences in ATO of 0·42 years or 0·47 BMI/fat mass index (FMI) Z-scores, between ED tertiles (80% power, \(\alpha = 0·05\), two-tailed).

### Anthropometry

Weight, height and skinfolds (triceps, biceps, subscapular and iliac) were measured at each visit as reported previously\(^{(20)}\). From age 2 years onwards, standing height was measured using a digital stadiometer. BMI Z-scores were calculated using the German reference\(^{(22)}\). Percentage of body fat was estimated using the Slaughter formula\(^{(23)}\). Parents weighed all foods and beverages consumed by their children, including water, for three consecutive days with regularly calibrated electronic scales (32·6% weekend days and 67·4% weekdays). The records were checked for completeness and accuracy, and mean EI and nutrient intakes were calculated using the in-house nutrient database\(^{(20)}\). The ratio of reported EI and calculated BMR was used to exclude two potential under-reporters\(^{(27)}\). Habitual dietary intakes were calculated as means 2 and 3 years before ATO.

We calculated ED (kJ/g) using four approaches\(^{(19,28,29)}\): (1) ED_all included all foods and drinks, regardless of energy content; (2) ED_energy included all foods and energy-containing drinks (>21 kJ, or 5 kcal, per 100 g)\(^{(19)}\); (3) ED_milk included all foods and milk as a drink, but no other beverages; (4) ED_food included foods only (solid/liquid).

### Puberty

ATO was estimated using the parametric Preece and Baines formula\(^{(24)}\). As described previously\(^{(21,25)}\), Preece and Baines formula 1 was fitted on various age ranges of height-for-age, beginning with age 2 years, to determine the optimal data range. ATO represented the age at minimal height velocity at onset of the pubertal growth spurt. Best fit was determined by four criteria, including graphical inspection of growth curves and comparison of residual standard deviations\(^{(21,25)}\). Finally, data from age 5 to 13 years (girls) and age 6 to 13 years (boys) were selected. Preece and Baines formula 1 was also used to estimate age at peak height velocity (\(n = 216\)). Menarcheal age was available in 104 girls.

### Diet

Dietary intake was assessed by 3 d weighed dietary records, as described previously\(^{(20)}\). Parents weighed all foods and beverages consumed by their children, including water, for three consecutive days with regularly calibrated electronic scales (32·6% weekend days and 67·4% weekdays). The records were checked for completeness and accuracy, and mean EI and nutrient intakes were calculated using the in-house nutrient database\(^{(20)}\). The ratio of reported EI and calculated BMR was used to exclude two potential under-reporters\(^{(27)}\). Habitual dietary intakes were calculated as means 2 and 3 years before ATO.

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### Other data

At study entry, parents were interviewed, weighed and measured. Parents of children were regularly asked which

### Table 1. Puberty, family and dietary characteristics of the DOrtmund Nutritional and Anthropometric Longitudinally Designed Study sample (Mean values, standard deviations, number of participants and percentages, \(n = 219\))

<table>
<thead>
<tr>
<th>Variables</th>
<th>Mean</th>
<th>SD</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Birth weight &lt; 3000 g</td>
<td></td>
<td></td>
<td>24</td>
<td>11·0</td>
</tr>
<tr>
<td>Age at baseline (years)</td>
<td>6·9</td>
<td>1·2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATO</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
<td>9·4</td>
<td>1·2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Girls ((n = 116))</td>
<td>8·7</td>
<td>0·9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Boys ((n = 103))</td>
<td>10·3</td>
<td>0·9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overweight*</td>
<td>34</td>
<td>15·5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Family and lifestyle</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal age at birth (years)</td>
<td>30·1</td>
<td>3·9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal overweight†</td>
<td></td>
<td></td>
<td>70</td>
<td>32·0</td>
</tr>
<tr>
<td>Maternal schooling &gt; 12 years</td>
<td></td>
<td></td>
<td>115</td>
<td>52·5</td>
</tr>
<tr>
<td>High physical activity at age 5 years†</td>
<td></td>
<td></td>
<td>46</td>
<td>21·0</td>
</tr>
<tr>
<td>Diet at baseline</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total energy (kJ/d)</td>
<td>6386·4</td>
<td>1105·8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total amount (g/d)</td>
<td>1602·9</td>
<td>334·2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carbohydrates (%E)</td>
<td>50·7</td>
<td>4·8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fat (%E)</td>
<td>36·4</td>
<td>4·3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Protein (%E)</td>
<td>12·8</td>
<td>1·6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fibre (g/100 kJ)</td>
<td>4·4</td>
<td>1·0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ED</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ED_all (kJ/g)</td>
<td>4·1</td>
<td>0·6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ED_energy (kJ/g)</td>
<td>5·1</td>
<td>0·7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ED_milk (kJ/g)</td>
<td>6·0</td>
<td>0·9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ED_food (kJ/g)</td>
<td>6·9</td>
<td>1·0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ATO, age at take-off; %E, percentage of total energy intake; ED, dietary energy density; ED_all, all foods and drinks; ED_food, foods and energy-containing drinks (>21 kJ/100 g); ED_milk, foods and milk as a drink, but no other beverages; ED_food, foods only.

* According to the International Obesity Task Force definition\(^{(22)}\).
† BMI \(>25\) kg/m\(^2\), yes/no.
‡ Based on the parental interview (which sports do children perform and how often).
Dietary energy density and puberty onset

Statistics

Normality of continuous variables was assessed by statistical testing and graphical inspection. We calculated adjusted mean outcome levels (ATO or BMI/FMI Z-score at ATO) in tertiles of ED. $P$-trend across tertiles was determined by multiple linear regression, with ATO or BMI/FMI Z-score at ATO as outcomes and ED as the independent variable. There was no significant interaction of ED with sex ($P$>0·1), and stratification did not indicate sex differences. Potential confounders were selected for their relationship with ED and the outcomes, or based on the literature(9,15,21,25,30): sex; birth year; birth weight (>3000 g, yes/no); maternal overweight (BMI $\geq$ 25 kg/m$^2$, yes/no); maternal education ($\geq$ 12 years schooling, yes/no); maternal age at birth; full breastfeeding ($\geq$ 4 months, yes/no); household smoking (yes/no); physical activity (low/moderate/high); protein, fat (percentage of EI, percentage of total energy intake); fibre (g/100kJ); baseline BMI/FMI Z-score. Variables changing the ED estimate or associated with the outcome levels (ATO or BMI/FMI Z-score) were used in the models. Diet quality according to the Nutritional Quality Index (low/moderate/high) formed a potential mediator, since it could lie on the pathway between ED and ATO(16). Analyses were performed using SAS 9·1 (SAS, Inc., Cary, NC, USA).

Results

ATO took place approximately 1·5 years earlier in girls than in boys (Table 1). ED ranged between 4·1 and 7·0 g/kJ, depending on the calculation method.

ED_food at baseline was not associated with ATO, BMI or FMI Z-score and ED as the independent variable. There was no significant interaction of ED with sex ($P$>0·1), and stratification did not indicate sex differences. Potential confounders were selected for their relationship with ED and the outcomes, or based on the literature(9,15,21,25,30): sex; birth year; birth weight (>3000 g, yes/no); maternal overweight (BMI $\geq$ 25 kg/m$^2$, yes/no); maternal education ($\geq$ 12 years schooling, yes/no); maternal age at birth; full breastfeeding ($\geq$ 4 months, yes/no); household smoking (yes/no); physical activity (low/moderate/high); protein, fat (percentage of EI, percentage of total energy intake); fibre (g/100kJ); baseline BMI/FMI Z-score. Variables changing the ED estimate or associated with the outcome ($P$<0·1) were retained in the models. Diet quality according to the Nutritional Quality Index (low/moderate/high) formed a potential mediator, since it could lie on the pathway between ED and ATO(16). Analyses were performed using SAS 9·1 (SAS, Inc., Cary, NC, USA).

Table 2. Adjusted mean age at take-off (ATO, in years), BMI Z-score and fat mass index (FMI) Z-score at ATO in tertiles of pre-pubertal dietary energy density (ED) based on foods only, DOrtmond Nutritional and Anthropometric Longitudinally Designed Study

(Least square mean (lsmean) values and 95% confidence intervals, n 219)

<table>
<thead>
<tr>
<th>Tertile of ED_food* at baseline (kJ/g)</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>lsmean 95% CI</td>
<td>lsmean 95% CI</td>
<td>lsmean 95% CI</td>
</tr>
<tr>
<td>ATO (years)†</td>
<td>9·3 9·0, 9·5</td>
<td>9·2 8·9, 9·4</td>
<td>9·4 9·1, 9·7</td>
</tr>
<tr>
<td>BMI Z-score at ATO‡</td>
<td>0 −0·1, 0·2</td>
<td>0 −0·1, 0·1</td>
<td>0 −0·1, 0·2</td>
</tr>
<tr>
<td>FMI Z-score at ATO§</td>
<td>0 −0·2, 0·1</td>
<td>−0·1 −0·2, 0·1</td>
<td>0·1 −0·1, 0·2</td>
</tr>
</tbody>
</table>

* ED_food corresponds to ED based on food intake only.
† Adjusted for sex, maternal age at birth (years), birth weight (<3000 g, yes/no), protein percentage of total energy intake (%E), fibre (g/100kJ) and baseline BMI Z-score.
‡ Adjusted for sex, maternal overweight (BMI $\geq$ 25 kg/m$^2$, yes/no), birth weight (<3000 g, yes/no), protein (%E), fibre (g/100kJ) and baseline BMI Z-score.
§ Adjusted for sex, maternal overweight (BMI $\geq$ 25 kg/m$^2$, yes/no), birth weight (<3000 g, yes/no), protein (%E), fibre (g/100kJ) and baseline BMI Z-score.

Discussion

The present results suggest that healthy children with higher pre-pubertal ED do not experience their pubertal growth spurt earlier.

One mechanism by which ED could influence ATO would be increased body fatness. In another DONALD subsample (n 215), neither BMI nor FMI were clearly associated with ATO(10) which supports the present results. However, ED was related to later pubertal markers (age at peak height velocity, menarche) either – and these were predicted by pre-pubertal body composition in the previous analysis. Another mechanism could be that higher ED lowers diet quality, related to ATO in 222 DONALD Study participants(16). Nutritional Quality Index scores were inversely related to ED in the present study (data not shown), but considering diet quality had no impact.

Similar to puberty onset, ED was not related to body fatness at ATO, i.e. at a distinct physiological age in a child's development. This is a worthwhile addition since previous studies have relied on chronological age. However, at a given chronological age in childhood differences in body composition may just stem from different developmental stages. In one study(13), ED (excluding beverages) at 6–8 years was related to BMI gain at 13–17 years. However, no association existed for percentage of body fat, BMI or waist circumference Z-score. Johnson et al.(17) found that only higher ED at 7 years, not at 5 years, was associated with a higher risk of a BMI >80th percentile at 9 years.
ED may have been below a threshold above which unfavourable effects (increased adiposity, earlier puberty onset) might operate; it was lower than in other paediatric studies when similar calculations were compared\(^{17–19}\). DONALD Study participants have a higher socio-economic background, a general interest in health and might have healthier dietary behaviour\(^{20}\). Besides the ED range, this could be important since an effect of ED may exist on excess adiposity in particular\(^{17–19}\). However, extremes of anthropometry were not represented in this analysis either. Correspondingly, the risk for overweight at ATO did not differ significantly between ED tertiles (data not shown). The homogeneity of our sample by contrast makes it less susceptible to residual confounding.

An additional limitation lies in the dietary assessment, since 3 d might not be sufficient to derive valid estimates of EI\(^{31}\). We addressed this by considering two records per child, reflecting habitual intake, and excluded potential under-reporters. Lastly, only crude measures of physical activity were available.

In conclusion, a higher habitual ED in childhood was not associated with an earlier puberty onset or higher body fatness at that time point in healthy, free-living children with a low overall ED.

Acknowledgements

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References


