Fractional urinary fluoride excretion of 6–7-year-old children attending schools in low-fluoride and naturally fluoridated areas in the UK

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Abstract

F is an important trace element for bones and teeth. The protective effect of F against dental caries is well established. Urine is the prime vehicle for the excretion of F from the body; however, the relationship between F intake and excretion is complex: the derived fractional urinary F excretion (FUFE) aids understanding of this in different age groups. The present study aimed to investigate the relationships between (1) total daily F intake (TDFI) and daily urinary F excretion (DUFE), and (2) TDFI and FUFE in 6–7-year-olds, recruited in low-F and naturally fluoridated (natural-F) areas in north-east England. TDFI from diet and toothbrushing and DUFE were assessed through F analysis of duplicate dietary plate, toothbrushing expectorate and urine samples using a F-ion-selective electrode. FUFE was calculated as the ratio between DUFE and TDFI. Pearson’s correlation and regression analysis were used to investigate the relationship between TDFI and FUFE. A group of thirty-three children completed the study; twenty-one receiving low-F water (0.30 mg F/l) and twelve receiving natural-F water (1.06 mg F/l) at school. The mean TDFI was 0.076 (SD 0.038) and 0.038 (SD 0.027) mg/kg per d for the natural-F and low-F groups, respectively. The mean DUFE was 0.017 (SD 0.007) and 0.012 (SD 0.006) mg/kg per d for the natural-F and low-F groups, respectively. FUFE was lower in the natural-F group (30 %) compared with the low-F group (40 %). Pearson’s correlation coefficient for (1) TDFI and DUFE was +0.22 (P=0.22) and for (2) TDFI and FUFE was −0.63 (P<0.001). In conclusion, there was no correlation between TDFI and DUFE. However, there was a statistically significant negative correlation between FUFE and TDFI.

Key words: Fluoride; Urinary excretion; Dietary intake

F is a trace element which, following absorption from the gastrointestinal tract, is rapidly incorporated into calcified tissues that contain 99 % of body F. Although the influence of F on bone metabolism is less well defined, the protective effect against dental caries is well established. However, several recent studies in industrialised and developing countries have shown an increase in the prevalence of dental fluorosis in populations from communities with and without water fluoridation, which may suggest that the threshold of F exposure for maximising caries prevention while minimising the potential risk of dental fluorosis has been exceeded. Obtaining the best balance between substantial caries reduction and the avoidance of unsightly dental enamel fluorosis is of critical importance to public health planners.

According to recent epidemiological surveys in the UK, 39 % of 5-year-olds and 33 % of 11-year-olds had evidence of dental caries experience involving dentine, while dental caries experience was even higher (48 %) in 14-year-old English children. The relatively high prevalence of dental caries in UK children highlights the need for primary prevention programmes such as fluoridation schemes. Estimations of total daily F intake at an individual and community level are key when recommendations for F use are being considered. Ingestion of F may occur from water, foods, toothpaste and other therapeutic agents. Increasingly, residence in a non-fluoridated community does not automatically assure low F intake, nor does living in a fluoridated community mean adequate or high F intake, since food, drink or even bottled water produced in a fluoridated area may be...
transported to a non-fluoridated area and vice versa\cite{8}. In addition, some dietary factors can increase or reduce the absorption and excretion of F\cite{9}, making body F retention an important yet variable consideration. In the absence of high concentrations of certain cations (e.g. Ca and Al), almost 90% of F ingested with food is absorbed from the gastrointestinal tract and passed rapidly into the blood. The remaining 10% is excreted with the faeces. Urine is the prime vehicle for excretion of F that is absorbed but not taken up by bones. It is estimated that children under 6 years of age excrete approximately 50% of their ingested F through the urine\cite{10}. F in the urine has been suggested as a suitable non-invasive biomarker for F exposure\cite{11} because collection of information on dietary F and that ingested from toothbrushing, at a community level, is time consuming, costly and requires a high level of expertise. Furthermore, varying degrees of gastrointestinal F absorption from different sources of F intake, such as diet and dental care products, might limit the value of estimated F exposure with regard to its systemic effect. Given these limitations, measurement of urinary F excretion has been recommended as an adequate method for monitoring fluoridation schemes\cite{11,12}.

Studies of F intake and urinary excretion have shown a wide variation in urinary F excretion as a proportion of F intake, ranging from 32 to 80% in children\cite{13–21} as summarised in Table 1. There is, therefore, a need for more assessment of the suitability and validity of urinary F excretion for monitoring fluoridation schemes as well as for predicting total F intake. The aims of the present study were therefore to investigate the relationships between (1) total daily F intake (TDFI) and daily urinary F excretion (DUFE), and (2) TDFI and fractional urinary F excretion (FUFE) in children.

**Materials and methods**

The present study was conducted according to the guidelines laid down in the Declaration of Helsinki and all procedures involving human subjects/patients were approved by the County Durham and Tees Valley 2 Research Ethics Committee (ethics no. 09/H0908/9). Written informed consent was obtained from all participants/patients.

**Study population and recruitment**

The study was conducted in areas of the north-east of England where the water supply was not artificially fluoridated. Before commencing the study, associated ‘Research and Development’ approval was also obtained from the relevant Primary Care Trust’s Research Management and Governance Unit. The Director of Children’s Services Directorate and Local Education Authorities were also contacted and informed of the study.

Parents of children were contacted through the schools which agreed to take part in the study. The inclusion criteria were as follows: healthy children aged 6–7 years who were lifelong residents of the area; children not receiving any professionally applied topical F therapy.

Participating children were not randomly selected but were those for whom parental permission had been obtained. In total, forty-four informed written consents were obtained from parents of the children who met the study inclusion criteria. Following the recruitment, each child and his/her parents were visited twice at their home.

In visit 1, the weight of the child, without shoes and jacket, was measured to the nearest 0·1 kg using a portable digital balance (SOEHNLE Slim Design Linea; ADE (GmbH & Co.)). Their height was also measured to the nearest 0·1 cm using a stadiometer (SOEHNLE MZ10020; ADE (GmbH & Co.)). BMI was then calculated as weight (kg) divided by height squared (m\(^2\)).

At visit 1, parents were provided with a bag that contained equipment required for collection of urine and food (duplicate plate) samples and instructions on how to collect these samples. Information on the toothbrushing habits of the child was also collected and a home tap water sample taken for subsequent F analysis.

**Dietary assessment**

Dietary F intake of the children was monitored by the ‘duplicate plate’ method as described by Guha-Chowdhury et al.\cite{22}. The parents were asked to maintain the usual dietary habits of their children and duplicate portions of all food and drink items as precisely as possible by observing and replicating

<table>
<thead>
<tr>
<th>Country</th>
<th>Age (years)</th>
<th>F exposure</th>
<th>TDFI (mg/d)</th>
<th>DUFE (mg/d)</th>
<th>FUFE (%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sweden\cite{20}</td>
<td>0·19–0·54</td>
<td>Breast-fed</td>
<td>0·011</td>
<td>0·30</td>
<td>359</td>
</tr>
<tr>
<td></td>
<td>0·15–0·31</td>
<td>Formula-fed</td>
<td>0·927</td>
<td>0·39</td>
<td>39</td>
</tr>
<tr>
<td>UK\cite{19}</td>
<td>1–3</td>
<td>Water (0·81 mg F/l)</td>
<td>0·71</td>
<td>0·33</td>
<td>48</td>
</tr>
<tr>
<td>USA\cite{13}</td>
<td>3–4</td>
<td>Water (optimal)</td>
<td>0·33</td>
<td>0·28 (including faeces)</td>
<td>Not reported</td>
</tr>
<tr>
<td>USA\cite{21}</td>
<td>0·19–0·89</td>
<td>Formula-fed</td>
<td>0·190</td>
<td>0·144</td>
<td>78</td>
</tr>
<tr>
<td>Chile\cite{15}</td>
<td>3–5</td>
<td>Water (0·5–0·6 mg F/l)</td>
<td>1·02</td>
<td>0·358</td>
<td>35·5</td>
</tr>
<tr>
<td>Germany\cite{16}</td>
<td>3–6</td>
<td>Salt F tablets (0·25–1 mg/d)</td>
<td>0·931</td>
<td>0·476</td>
<td>51·5</td>
</tr>
<tr>
<td>Iran\cite{14}</td>
<td>4–7</td>
<td>Water (0·30–0·39 mg F/l)</td>
<td>0·428</td>
<td>0·339</td>
<td>80</td>
</tr>
<tr>
<td>Colombia\cite{17}</td>
<td>4–5</td>
<td>Table salt (180–220 mg F/kg)</td>
<td>1·55</td>
<td>0·414</td>
<td>33</td>
</tr>
<tr>
<td>UK\cite{18}</td>
<td>6–7</td>
<td>Water</td>
<td>0·08 mg F/l</td>
<td>0·736</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0·47 mg F/l</td>
<td>0·883</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0·82 mg F/l</td>
<td>1·043</td>
<td>32</td>
</tr>
</tbody>
</table>

* DUFE as a percentage of TDFI.
the actual consumed amounts by the children over 24 h. They were asked to remove parts of food items not normally eaten such as bones, fruit skin, cores, etc., before placing them in the container provided. They were also asked to collect drinks separately in the plastic bottles provided.

Each parent was also supplied with a 1 d food diary and accompanying instructions, for the recording of food and drink consumed on the day the duplicate plate was collected. This was done so that the researchers could cross-reference the information in the diary with the duplicate plate analysis for validation purposes. For those children who were at school during the sample collection period, researchers who were on-site during the school day obtained a duplicate of the child’s school dinner and noted the items consumed. Any snacks and drinks including free school fruit and food consumed at breakfast clubs/breaks were also included in the duplicate plate.

**Assessment of ingested toothpaste**

F intake from ingested toothpaste during toothbrushing was estimated using the method described by Maguire et al.\(^1\)\(^8\). In brief, expectorated toothpaste/saliva samples were obtained during a tooth brushing session, which took place either at the child’s school or at their home. Children provided their own toothpaste and each child was provided with a new toothbrush (Aquafresh Big Teeth for 6–7-year-olds or Aquafresh Milk Teeth for 3–4-year-olds). Toothbrushes were weighed before and after the child or parent dispensed toothpaste. Any toothbrushing expectorate was collected in a small plastic sample collection pot together with the water used to rinse the toothbrush.

**24 h urine collection**

Collection of the 24 h urine sample started on the same day as the duplicate plate collection (day 1). Parents were advised to record the time of the child’s first voided urine sample. All subsequent urine, up to and including the first passing of urine on the following day (day 2), was collected for the 24 h collection. During school hours, urine was collected by the child, supported by trained study researchers. Each child’s voided urine sample was passed to the researchers for storage until the full 24 h sample had been collected.

At visit 2, which was conducted on day 2, the day following duplicate plate collections and after the final collection of urine, all samples were collected from the family home. At the same time, the researchers went through the food diary with the parent and child and checked it against the items in the duplicate plate collection.

**Sample preparation and analysis**

Collected samples were then taken to the F laboratory for processing. Urine collected at home and school (where applicable) was mixed together to produce a pooled sample and the volume recorded. Expectorated saliva/toothpaste samples were vortexed for 30 s. Collected samples of home and school drinks were also mixed together and the volume recorded. Food collected at home and school was mixed, weighed and then homogenised using an industrial blender (Thermomix TM31; Vorwek). Finally, three aliquots each of urine, expectorated saliva, homogenised foods, water and drinks were taken and stored at \(-20^\circ\text{C}\) for further analysis.

Urine, water and drink samples were analysed, in triplicate, for F by a direct F analysis method using a F-ion-selective electrode (Model 9609; Orion Research) coupled to a potentiometer (Model 720A), after sample buffering with total ionic strength adjustment buffer (III)\(^2\)\(^9\). Food and expectorated saliva/toothpaste samples were analysed, in triplicate, for F concentration after overnight hexamethyldisiloxane-facilitated diffusion at room temperature using the F-ion-selective electrode and meter\(^2\)\(^4\). Of these samples, 10\% were re-analysed for F concentration, giving a mean reproducibility of 99.6\%.

The creatinine concentration of each urine sample was measured by the Jaffe method\(^2\)\(^5\) using an autoanalyser (ADIVA 1650; Siemens Medical Solutions Diagnostics).

**Data preparation and analysis**

F intake from toothpaste ingestion during toothbrushing was estimated by subtracting the F content of expectorated saliva/toothpaste from the amount of F initially loaded onto the brush during toothbrushing. F ingestion per brushing was then multiplied by the frequency of brushing (information at visit 1), to calculate the daily F intake from toothpaste for each child. Daily dietary F intake was estimated from the weight of each duplicate plate sample and the F concentrations of their aliquots. Since none of the children used any F supplements, total daily F intake (TDFI, in mg/d) was calculated by adding F intake from diet and F ingested from toothpaste.

Completeness of the 24 h urine samples was checked against two criteria: (1) the lower limits of 5 and 9 ml/h for urinary flow rate in \(<6-\text{and} \geq6\)-year-olds\(^3\)\(^1\)\(^1\), respectively, and (2) a lower limit of 11.3 mg/kg body weight (BW) per d for creatinine excretion\(^2\)\(^6\). Any sample that did not meet either of these criteria was excluded from further analysis.

DUFE (mg/d) was estimated from the 24 h urine volume and F concentration of the urine sample. TDFI and DUFE were also calculated based on body weight (mg/kg BW per d). FUFE (%) was then calculated from the following equation:

\[
\text{FUFE} \, (\%) = \left( \frac{\text{DUFE}}{\text{TDFI}} \right) \times 100.
\]

The data were analysed descriptively using SPSS version 17.0. The percentage of TDFI from diet and FUFE were calculated for each child, individually, before calculating the sample mean and standard deviation. The correlations between TDFI and DUFE and FUFE were examined by regression analysis and Pearson’s correlation.

**Results**

Of the forty-four recruited, thirty-four children completed all aspects of the study. Data from one child were excluded...
because the urine sample was incomplete. Therefore, the final sample was thirty-three children.

F analysis of school water supply showed a mean F concentration of 0·30 (SD 0·12) μg/ml for twenty-one children (low-F group) and 1·06 (SD 0·11) μg/ml for twelve children (natural-F group). The mean F concentration of home water supply for the low-F group was 0·20 (SD 0·10) and 0·49 (SD 0·32) μg/ml for the natural-F group.

The mean age of the low-F and natural-F groups was 6·8 (SD 0·6) and 6·6 (SD 0·3) years, respectively (Table 2). Although the average body weight of the low-F group was heavier (25·4 kg) than that of the natural-F group (22·8 kg), the BMI (kg/m²) was similar: 16·1 and 15·8 kg/m², respectively.

None of the children in the present study took any F tablets or supplements. Diet and toothpaste ingestion were therefore the only sources of F intake for these children. The mean total daily F intake was 1·707 (SD 0·799) mg/d for the natural-F group and 0·393 (SD 0·209) mg/d for the low-F group. On a mg/kg BW basis, this represented 0·076 (SD 0·007) mg/kg BW per d for the natural-F group and 0·012 (SD 0·006) mg/kg BW per d for the low-F group.

Approximately 71% of children used a toothpaste labelled with fluoride as a proportion of TDFI (%) 44 27 41 27

Table 2. Fluoride concentration of water supply, age, height, weight and BMI of children by fluoride area and sex (Mean values and standard deviations)

<table>
<thead>
<tr>
<th></th>
<th>Natural-F* area (n 12)</th>
<th>Low-F† area (n 21)</th>
<th>Girls (n 17)</th>
<th>Boys (n 16)</th>
<th>All children (n 33)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>Mean 6·6 0·3 SD 6·8 0·6</td>
<td>Mean 6·8 0·5 SD 6·8 0·7</td>
<td>Mean 6·8 0·7 SD 6·8 0·7</td>
<td>Mean 6·8 0·7 SD 6·8 0·7</td>
<td>Mean 6·8 0·6 SD 6·8 0·6</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>Mean 119·8 6·3 SD 125·9 12·2</td>
<td>Mean 125·5 13·4 SD 121·9 6·8</td>
<td>Mean 125·5 13·4 SD 121·9 6·8</td>
<td>Mean 125·5 13·4 SD 121·9 6·8</td>
<td>Mean 125·5 13·4 SD 121·9 6·8</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>Mean 22·8 3·2 SD 25·4 4·1</td>
<td>Mean 24·1 2·8 SD 24·8 4·9</td>
<td>Mean 24·1 2·8 SD 24·8 4·9</td>
<td>Mean 24·1 2·8 SD 24·8 4·9</td>
<td>Mean 24·1 2·8 SD 24·8 4·9</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>Mean 15·8 0·3 SD 16·1 2·7</td>
<td>Mean 15·5 0·7 SD 16·8 2·6</td>
<td>Mean 15·5 0·7 SD 16·8 2·6</td>
<td>Mean 15·5 0·7 SD 16·8 2·6</td>
<td>Mean 15·5 0·7 SD 16·8 2·6</td>
</tr>
</tbody>
</table>

Discussion

The knowledge base regarding the usefulness of urinary F excretion as a tool in epidemiological surveillance for prediction of total F intake in children is inadequate. The present study demonstrated that urinary F might not be a reliable estimator for F intake in children aged 6–7 years as suggested previously.

In the present study, no child used F supplements or F tablets, and therefore diet and dentifrice ingestion were the main sources of total daily F intake for all children. In populations using F toothpaste, diet has been reported as contributing to almost 80% of ingested F\(^{-27}\). However, in the present study, toothpaste was the major component of TDFI.

Table 3. Total daily fluoride intake (TDFI) from diet and toothpaste ingestion, daily urinary fluoride excretion (DUFE) and fractional urinary fluoride excretion (FUFE, %) for all participants (Mean values and standard deviations)

<table>
<thead>
<tr>
<th></th>
<th>Natural-F* area (n 12)</th>
<th>Low-F† area (n 21)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daily F intake</td>
<td>Mean 0·578 0·298 SD 0·341 0·254</td>
<td>Mean 0·229 0·166 SD 0·187 0·236</td>
</tr>
<tr>
<td>Food (mg/d)</td>
<td>Mean 0·349 0·263 SD 0·154 0·137</td>
<td>Mean 0·050 0·024 SD 0·024 0·024</td>
</tr>
<tr>
<td>Drink (mg/d)</td>
<td>Mean 1·130 0·820 SD 0·606 0·562</td>
<td>Mean 0·050 0·024 SD 0·024 0·024</td>
</tr>
<tr>
<td>From toothpaste ingestion mg/d</td>
<td>0·076 0·038 SD 0·038 0·027</td>
<td>0·017 0·007 SD 0·017 0·006</td>
</tr>
<tr>
<td>mg/kg BW per d</td>
<td>0·034 0·028 SD 0·024 0·024</td>
<td>0·007 0·004 SD 0·007 0·003</td>
</tr>
<tr>
<td>TDFI† mg/d</td>
<td>Mean 0·045 0·061 SD 0·027</td>
<td>Mean 0·045 0·061 SD 0·027</td>
</tr>
<tr>
<td>mg/kg BW per d</td>
<td>Mean 0·038 0·027 SD 0·038 0·027</td>
<td>Mean 0·038 0·027 SD 0·038 0·027</td>
</tr>
<tr>
<td>Diet as a proportion of TDFI (%) Diet as a proportion of TDFI (%)</td>
<td>41 27 SD 44 27</td>
<td>41 27 SD 44 27</td>
</tr>
<tr>
<td>Urinary F excretion</td>
<td>Volume of urine (ml/d) Mean 547 304 SD 607 314</td>
<td>Mean 547 304 SD 607 314</td>
</tr>
<tr>
<td>DUFE mg/d</td>
<td>Mean 0·393 0·209 SD 0·297 0·131</td>
<td>Mean 0·017 0·007 SD 0·017 0·006</td>
</tr>
<tr>
<td>mg/kg BW per d</td>
<td>Mean 30 21 SD 40 22</td>
<td>Mean 30 21 SD 40 22</td>
</tr>
</tbody>
</table>

* 1·06 μg F/ml.
† 0·30 μg F/ml.
‡ None of the children used F supplements or F tablets.
in both natural-F and low-F areas. Generally, toothpaste can make the largest percentage contribution to TDFI in children younger than 6 years, as they are not in full control of their swallowing reflex and therefore might swallow significant amounts of toothpaste unintentionally. At this age, the crowns of permanent teeth are still undergoing calcification and are therefore susceptible to the uptake of F into enamel apatite, and as a result, excess F intake can result in dental fluorosis. The literature shows a wide variation in the contribution of F toothpaste to TDFI ranging from 22% for 6-year-olds in Iowa to 69% for 4–5-year-olds in Puerto Rico. The differences in the contribution of toothpaste to TDFI in different studies could be explained by the differences in children ages, the F concentrations of the toothpastes used and the diet consumed, as well as the data collection methods and techniques used to measure F intake from these sources.

In the present study, mean daily F intake from drinks was substantially higher in the natural-F area compared with that in the low-F area, which confirms that the impact of F concentration of home water supply on total F intake may be decreasing due to the trend towards consumption of foods and drinks made outside the home.

The mean TDFI of children in the natural-F area (0.076 mg/kg BW per d) was slightly higher than the suggested optimum range of 0.05–0.07 mg/kg BW per d for optimal dental health benefit, whereas for children living in the low-F area, the TDFI (0.038 mg/kg BW per d) was below the optimum range. Therefore, in low-F communities, children might benefit from a community-based preventive programme such as milk fluoridation or supervised toothbrushing at schools.

The mean DUFE when expressed on a body-weight basis for the two groups of children was fairly similar, despite the considerable difference in TDFI between the groups (Table 3). The mean FUFE of children in the natural-F group (30%) was lower than the corresponding value for children in the low-F area (40%). The estimated FUFE varies widely in the literature from 32% for 6–7-year-olds to 359% for breast-fed infants (Table 1). There are several possible explanations for the wide range of reported FUFE. Almost 50% of ingested F is absorbed from the stomach; however, several substances influence the degree of absorption. High dietary levels of fat may increase the absorption of ingested F since the fat reduces the rate of gastric emptying. In addition, foods containing appreciable amounts of divalent or trivalent cations (e.g., Ca, Mg, Fe) may reduce the degree of absorption due to the formation of insoluble complexes. The kidneys are the major route for the removal of F from the body and urinary pH can influence the renal clearance of F. When the tubular fluid is acidic, more ionic F is converted to hydrogen fluoride which is diffusible across the renal tubular epithelium. Differences in the composition of diet and the altitude of residence can significantly influence urinary pH, and consequently F excretion. Age, kidney maturation and body size (existing skeletal mass) are also important variables in F retention.

In a recent study, the relationship between urinary F excretion and TDFI was examined using previously published data on F intake and excretion in children and adults. This study showed a positive linear relationship between urinary F excretion and F intake with a slope of +0.05 and intercept of 0.03 in children, suggesting that urinary F excretion can be used to estimate daily F intake in children younger than 7 years. However, in the present study, daily urinary F excretion did not correlate with TDFI, and there was a lower slope of +0.05 and a higher intercept of 0.27. This result implies that for 6–7-year-old children living in an industrialised country, TDFI cannot be adequately predicted from urinary excretion of F, in contrast to the results of the former study. However, there are two main differences between these two studies; the present study was based on the data from only thirty-three children with a narrow age range (6–7-year-olds), whereas the former study included pooled data from 212 children with a wide age range from 0–19 to 7 years. The stage of bone maturation can influence the rate of uptake of F into bones and teeth. Since the rate of uptake is greater into newly formed bones, F retention would be greater during periods of rapid growth and development. The differences in urinary F excretion between different age groups of young children may be also attributed to the differences in their diet as well as dietary habits. For example, the absorption of F from ingested water is almost 100%, however, when F is taken with milk, the degree of absorption...
might be reduced by up to 50% due to the formation of CaF$_2$ which has a low aqueous solubility\(^{13,33}\).

The negative correlation between FUFE and TDFI observed in the present study implies a higher F retention with increasing F intake. However, Fig. 2 suggests that FUFE remains almost constant above a TDFI of approximately 1.6 mg/d with the estimated FUFE reaching a limiting constant value independent of the magnitude of TDFI.

In conclusion, there was a statistically significant negative correlation between FUFE and TDFI, but no correlation between TDFI and DUFE, in 6–7-year-olds. Therefore, DUFE might not provide the basis for a reliable estimate of total F intake for 6–7-year-old children. However, this relationship should be investigated further in different age groups, separately, with larger sample sizes, in order to establish any conclusion on the use of DUFE as a reliable estimate of TDFI in children.

Acknowledgements

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References