Vitamin D status and its determinants during autumn in children at northern latitudes: a cross-sectional analysis from the OPUS School Meal Study

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Vitamin D status and its determinants during autumn in children at northern latitudes: a cross-sectional analysis from the optimal well-being, development and health for Danish children through a healthy New Nordic Diet (OPUS) School Meal Study

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Abstract
Sufficient summer/autumn vitamin D status appears important to mitigate winter nadirs at northern latitudes. We conducted a cross-sectional study to evaluate autumn vitamin D status and its determinants in 782 Danish 8–11-year-old children (55°N) using baseline data from the Optimal Well-being, Development and Health for Danish Children through a Healthy New Nordic Diet (OPUS) School Meal Study, a large randomised controlled trial. Blood samples and demographic and behavioural data, including 7-day dietary recordings, objectively measured physical activity, and time spent outdoors during school hours, were collected during September–November. Mean serum 25-hydroxyvitamin D (25(OH)D) was 60.8 (SD 18.7) nmol/l. Serum 25(OH)D levels ≤50 nmol/l were found in 28.4 % of the children and 24 % had concentrations <25 nmol/l. Upon multivariate adjustment, increasing age (per year) (β = 2.9; 95 % CI 5.2, –5.1, –0.7 nmol/l), female sex (β = 3.2; 95 % CI 5.9, –0.7 nmol/l), sampling in October (β = 5.2; 95 % CI 10.1, –0.4 nmol/l) and November (β = 13.3; 95 % CI 17.7, –9.1), and non-white ethnicity (β = 5.7; 95 % CI 11.1, –0.3 nmol/l) were negatively associated with 25(OH)D (all P < 0.05). Likewise, immigrant/descendant background was negatively associated with 25(OH)D, particularly in females (β = 16.3; 95 % CI 21.9, –10.7) (P < 0.001) (Pinteraction = 0.003). Moderate-to-vigorous physical activity (MVPA) (min/d) (β = 0.06; 95 % CI 0.01, 0.12), outdoor walking during school hours (min/week) (β = 0.4; 95 % CI 0.1, 0.6) and intake of vitamin D-containing supplements ≥3/day (β = 8.7; 95 % CI 6.4, 11.0) were positively associated with 25(OH)D (all P < 0.05). The high proportion of children with vitamin D status below the recommended sufficiency level of 50 nmol/l raises concern as levels expectedly drop further during winter months. Frequent intake of vitamin D supplements was strongly associated with status. MVPA and outdoor activity during school hours should be investigated further in interventions to improve autumn vitamin D status in children at northern latitudes.

Key words: Vitamin D: Children: Northern latitudes: Determinants

Determinants of vitamin D status are complex to evaluate but require attention as vitamin D deficiency has been found to occur commonly among, for example, healthy European children and adolescents1, and as vitamin D facilitates intestinal Ca absorption and optimal bone health2–5. Vitamin D plays a particularly crucial role during childhood growth stages, when accumulation of skeletal Ca increases vastly6,7. The vitamin D metabolite 25-hydroxyvitamin D (25(OH)D) has a half-life of 2–3 weeks, and currently serves as a suitable marker of vitamin D status5. Although different cut-off levels are advocated6, the European Society for Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) has recently re-established the recommendation of a 25(OH)D concentration >50 nmol/l (20 ng/ml) to indicate sufficiency7. Severe vitamin D deficiency during childhood, established by ESPGHAN as 25(OH)D concentrations <25 nmol/l7, may lead to painful joints, muscle/
bone disorders and overt rickets (7,8). In addition, vitamin D status
has been inversely associated with extra-skeletal conditions
such as type 1 diabetes, CVD and certain cancers (9,10). Under
conditions of regular skin exposure to sunlight, dietary vitamin D
intake is estimated to contribute about 10% of human vitamin D
requirements (3), and only few foods such as egg yolks and fatty
fish are naturally rich in vitamin D (3,11). The remaining 90% is
estimated to originate from the production that occurs in the
skin upon exposure to UVB waves in sunlight (5). Indeed, sun
exposure of hands, arms and face in non-skin-redening doses
2-3 times/week has been suggested to satisfy the vitamin D
requirement of most people (12). Yet, decreased solar intensity
makes cutaneous production unattainable from November
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skin upon exposure to UVB waves in sunlight (5). Indeed, sun
exposure of hands, arms and face in non-skin-redening doses
2-3 times/week has been suggested to satisfy the vitamin D
requirement of most people (12). Yet, decreased solar intensity
makes cutaneous production unattainable from November
through February at latitudes of 52°N and above (13–15). Children
living in northern latitude countries are therefore thought to be at
risk of vitamin D deficiency, particularly in winter/early spring.
Indeed, sufficient summer/autumn vitamin D status appears
important to mitigate such winter nadirs in these children (16). It is
therefore of great relevance to identify behavioural factors of
summer/autumn vitamin D status in children at these latitudes.

In addition to seasonal UVB fluctuations, the extent of skin
pigmentation has been shown to modify the cutaneous vitamin D
production so that dark-skinned children generally require more
sun exposure compared with fair-skinned peers to produce the
same amount of vitamin D (17–19). Indeed, previous studies on
vitamin D in Scandinavian children have consistently found
vitamin D concentrations to be lower in winter than in summer,
and among immigrant or dark-skinned children, and they have
shown a positive association between vitamin D status and intake
of vitamin D supplements (20–25). Other observed, but less
consistent, correlations with vitamin D status in Scandinavian
children have included dietary vitamin D intake (24), sun habits (16)
and, inversely, BMI (22,24). Yet, previous studies have had relatively
small sample sizes (16,20–25), and most have included only specific
groups such as adolescent girls or immigrant children (16,20–25).

The aim of the present cross-sectional study was to evaluate
autumn vitamin D status and identify demographic and beha-
vioral determinants hereof in a large and highly representative
sample of children in Denmark (55°N). The study was based on
baseline data from the Optimal well-being, development and
health for Danish children through a healthy New Nordic Diet
(OPUS) School Meal Study and included 782 third and fourth
graders from nine schools in Denmark. The study provided
innovative and previously unexamined vitamin D determinants at
this latitude and time of year, including the role of being outdoors
during school hours.

Methods

Study design and subjects

The OPUS School Meal Study was a large cluster-randomised
controlled cross-over trial that aimed to assess the impact of
providing school meals based on the New Nordic Diet on dietary
intake and nutrient status, growth, early disease risk markers,
well-being and absence from school, cognitive function, food
waste and cost, as well as social features in third- and fourth-grade
children. The design of the OPUS School Meal Study has been
described in detail elsewhere (20). Participants were recruited from
May to October 2011. Briefly, initial contact was established with
thirty-nine schools, of which nine were included. Inclusion
criteria for the schools in the OPUS Study were (a) location in the
eastern part of Denmark (Zealand and Lolland-Falster), (b) a total
of four or more classes at third and fourth grade, (c) suitable
kitchen facilities available for food production during school
hours and (d) a high motivation for participation as determined
by the study team. As more than nine schools were eligible for
participation, logistical matters were also taken into consideration.
Moreover, the aim was that ≥50 % of the schools should belong to
municipalities where (a) the income was below the national mean
for families with children and (b) the percentage of households in
which the adults had lower secondary education (≤10 years) was
above the national mean for families with children, as defined
by Statistics Denmark (27). Ultimately, three of the nine (33 %)
included schools fulfilled these criteria. Written information about
the study was sent to the families of all third- and fourth-grade
children at the nine participating schools, and the families were
invited to an information meeting. In order to recruit families of
various socio-economic and ethnic backgrounds, we held several
information meetings at each school, sent out several reminders,
and had translators present at meetings when relevant. Exclusion
criteria for the invited third- and fourth-grade children were
(a) diseases or conditions that might obstruct the measurements
or put the child at risk by eating the intervention school meals, or
(b) concomitant participation in other scientific studies that
involved radiation or blood samples. We obtained written
informed consent for participation from all custody holders of 834
children, corresponding to 82 % of the 1021 invited children. The
enrolment percentage at the nine schools ranged from 60 to 89 %.
The schools were located in eight separate municipalities and the
included children came from rural as well as urban areas.

Ethics statement

This study was conducted according to the guidelines laid down in
the Declaration of Helsinki and all procedures involving human
subjects were approved by the Committees on Biomedical
Research Ethics for the Capital Region of Denmark (H-1-2010-124);
the study was registered in the database www.clinicaltrials.gov
(no. NCT 01577277). Written informed consent was obtained
from custody holders of all subjects.

Socio-demographic characteristics, screen time and puberty stage

Information on demographic and socio-economic characteristics
was collected at baseline by an in-depth 2-h personal interview
with the child and parents. The highest education level attained in
the household was used to categorise parental educational level
into lower secondary education (≤10 years), upper secondary
education, vocational education, short higher education, a
Bachelor’s degree or equivalent, or a Master’s degree or higher
(≥17 years). The birth countries of parents and grandparents were
used to categorise the children as immigrant/descendants or not.
Children were categorised into ‘immigrants/descendants’ if all
grandparents and ≥1 parent were born outside Denmark. Though
not identical, this classification was based on the definitions of immigrants and descendants used by Statistics Denmark.280

Parents estimated the child's daily ‘screen time’ during the previous week – that is, watching television or using the computer and game consoles – and the registered min/weekday was applied for this variable. Finally, puberty status was self-evaluated by child and parents on the basis of breast development in girls and pubic hair in boys (Tanner stages) as validated by Morris & Udry.289 Registrations were dichotomised into a variable indicating entered puberty (≥ Tanner stage 2) (yes/no).

Registration of dietary intake, physical activity and outdoor activity during school hours

Before the clinical measurements, the children, aided by their parents, recorded their daily intake of food and beverages for 7 consecutive days using a Web-based Dietary Assessment Software for Children developed specifically for children aged 8–11 years.280 The software was validated in the OPUS School Meal Pilot Study, although not specifically for vitamin D intake.31,32 Intake data were processed by a General Intake Estimation System, originally developed for the Danish National Survey of Diet and Physical Activity 2003–2008 (Division of Nutrition, The National Food Institute, Technical University of Denmark, Denmark).331 Energy intake (EI) relative to estimated BMR was evaluated following the equations of Henry,34 and energy under-reporters (EI:BMR ≤ 0.5) and over-reporters (EI:BMR ≥ 1.28)331 were excluded from the dietary variables, as were children who registered their diet for <4 d. Dietary intake did not include the intake from supplements.

The intake of dietary supplements was registered separately in the dietary recordings and the daily intake of vitamin D-containing supplements, including multivitamins, was summarised in the current study. Supplement intake was applied without excluding children who recorded for <4 d and without excluding energy over- and under-reporters.

To measure physical activity, the children were asked to wear an accelerometer (ActiGraph GT3X+ Tri-Axis Accelerometer Monitor) in an elastic belt tightly at the right hip for the same seven days and eight nights as the dietary recordings, and to remove it only during water activities – that is, showering or swimming.

Analyses of the physical activity data are described in detail elsewhere.331 Derived variables included time spent sedentarily (0–100 counts per minute (cpm)) and moderate-to-vigorous physical activity (MVPA) (≥2296 cpm). Also, the families recorded the child's amount of bicycling during weekdays.

School settings and policies on outdoor activity differed between the nine schools. From a questionnaire to the principal of each school we obtained information on whether the children were obligated by school policy to spend all recesses outdoors (obligatory outdoor recesses), and approximately how many minutes per week the children spent outside during school hours in walking between classrooms and other school facilities (outdoor walks between classrooms). Outdoor walks between classrooms indicated a necessary non-voluntary outdoor walking activity certain to take place for the child to get between classrooms. This variable depended on the school architecture as some schools were made up of several buildings scattered around an area, whereas other schools were made up of one entire building that therefore did not require any outdoor walking to get between classrooms during school hours.

Anthropometry and ethnicity

Anthropometric measurements were conducted during morning hours at each school, after the child had been fasting since midnight, had emptied the bladder and was wearing only light clothing. Height was measured in cm to the nearest one decimal using a transportable stadiometer (CMS Weighing Equipment Ltd). Height was derived as the mean of three consecutive measurements with the child in standing position, holding his head in the Frankfurter plane. Body weight was measured in kg to the nearest one decimal using a digital scale (BWB 800; Tanita). BMI was calculated as body weight (kg)/height (m)², and sex- and age-adjusted z-scores for BMI were calculated using WHO AnthroPlus software.372 The OPUS School Meal Study also included dual-energy X-ray absorptiometry (DXA) scans of each child conducted at the same occasion as the blood sampling. In relation to the DXA scan the ethnicity of the child was originally recorded by an investigator based on physical presence and statement of the child. Ethnicity was recorded according to the terminology of the Lunar Prodigy Pro™ (GE Medical Systems) DXA scanner with EnCore™ software – that is, Caucasian, Asian, African, Latin or Other. These recordings were applied for ethnicity categorisation in the current study – that is, children recorded as Caucasian were named white ethnicity. These also included children from Turkey, the Middle East, Pakistan and India. Because of small numbers in the four non-Caucasian groups (non-Caucasian n 41), we dichotomised ethnicity recordings into a variable indicating white ethnicity (yes/no).

Blood markers of vitamin D status and fatty fish intake

Blood samples were collected in August/September (n 311), October (n 146) or November (n 325) 2011 at the same time as the anthropometric measures. As August samples were few (n 21), and were collected only on 30 and 31 August, these samples were combined with the September samples. Local anaesthetic patches were offered to the children to reduce discomfort, and blood was sampled by means of venepuncture after an overnight fast from all children at one school before moving on to the next school. Serum was separated by centrifuging collected blood at 2500 g at 4°C for 10 min after 30 min at room temperature. Serum was then stored at −80°C for later vitamin D analysis. Heparinised whole blood was mixed with 0·1 % butylated hydroxytoluene (Sigma-Aldrich) in ethanol (0·1 ml/ml blood) and stored at −80°C for later analysis of fatty acid composition. As a measure of vitamin D status, 25(OH)D concentrations (25-hydroxy-vitamin D₂+D₃ (DTOT25)) were assessed in serum by automatic Chemiluminescence ImmunoAssay technology on DiaSorin Liaison (DiaSorin AB) at the Department of Clinical Biochemistry, Aalborg University Hospital, Aalborg, Denmark, a laboratory partaking in the vitamin D External Quality Assessment Scheme (DEQAS).380 Before analysis, serum was defrosted at room temperature for half an hour and centrifuged at 4000 rpm for 3 min. Concentrations are
presented in nmol/l (1 nmol/l = 0.4006 ng/ml), and the lowest detection limit was 10 nmol/l. Intra-assay CV for 25(OH)D was 7.6%, whereas inter-assay CV for 25(OH)D was 5.4%. As a biomarker of fatty fish intake, EPA + docosahexanoic acid (DHA) status was assessed in whole blood by means of high-throughput GC. Fatty acid methyl esters were prepared from whole-blood samples by direct trans-esterification with convectional heat as previously described39,40. The amount of EPA + DHA is presented as w/w% of the total whole-blood fatty acids. Intra- and inter-assay CV for EPA wt% were 1.2 and 4.5%, whereas intra- and inter-assay CV for DHA wt% were 2.4 and 6.4%, respectively.

### Statistical analyses

The statistical analyses were carried out using Stata (version 12.1; StataCorp LP). Significance was established at \( P < 0.05 \). Between-group differences in population characteristics were evaluated using the two-sample \( t \) test (unequal variances), the Wilcoxon and Mann-Whitney test (for non-normally distributed outcomes) and the \( \chi^2 \) or Fisher’s exact test for categorical outcome variables. Linear mixed effects models were used to identify factors associated with serum 25(OH)D concentrations (the outcome). Model checking was based on visual inspection of residual and normal probability plots. For all mixed models, school, grade and class were included as random effects to account for the dependency structures imposed by the hierarchical design. A priori, we considered month of sampling and intake of vitamin D supplements to be well-established determinants of vitamin D status to be included in all models. Initially, a basic demographic linear mixed model was constructed with outcome and random effects as specified above, and with month of sampling, intake of vitamin D-containing dietary supplements (in proportion to days of dietary recordings – i.e. days with intake of vitamin D-containing supplements/total number of days of dietary recordings) and all demographic variables included as fixed effects: that is, white ethnicity (yes/no), immigrant/descendant background (yes/no), age, sex, entered puberty (i.e. Tanner stage \( \geq 2 \) (yes/no)) and parental education level. Subsequently, interactions between sex and puberty, sex and immigrant/descendant background, sex and ethnicity, and sex and month of sampling were included one by one in this demographic model and tested by likelihood ratio tests. Significant interactions were retained in the model. In the next step, each behavioural variable identified (see below) was assessed one by one through inclusion in the above demographic model. Because of the behavioural complexity of vitamin D sources, this approach was applied to increase the transparency of the contribution of each of the behavioural variables distinct from the effects of the remaining explanatory variables in the model, and to address potential collinearity between variables. The behavioural variables included were as follows: MVPA (min/d), sedentary time (min/d), ‘screen time’ (min/weekday), bicycling (min/weekday), BMI-for-age z-score, outdoor walking between classrooms during school hours (min/week), obligatory outdoor recesses during school hours (yes/no), whole-blood EPA + DHA (wt%) as a biomarker of fatty fish intake, intake of dietary vitamin D (\( \mu \)g/d) and intake of dietary Ca (mg/d). Eventually, to evaluate the strength of the associations found for each single behavioural variable, all significant behavioural variables were simultaneously included in the demographic model.

In addition, to further elaborate on the association between serum 25(OH)D concentration and frequency of vitamin D-containing supplement intake, we conducted a more detailed sub-analysis with a categorical vitamin D supplement variable including only children who recorded their dietary intake for all 7 d (days with intake of vitamin D-containing supplements/7 d). This categorical supplement profile was then evaluated in the demographic model.

### Results

#### Subject characteristics

Of the 834 children enrolled in the OPUS School Meal Study, eleven children withdrew from the study before the clinical measurements at baseline, seven were ill or on holiday on the day of sampling, and thirty-four children failed to deliver a blood sample or had a sample with insufficient blood volume for 25(OH)D analysis. Ultimately, baseline samples for 25(OH)D analysis were successfully collected from 782 children. These \( n = 782 \) children comprised the study population for evaluation of the vitamin D status in the present study and had an age range of 8–11 years. Boys were older, spent more time with MVPA, spent more screen time, and had higher serum 25(OH)D concentrations compared with girls. More girls than boys had entered puberty (Table 1). Overall, 62% of the children were obligated to go outdoors during all recesses of their school day. This applied for more third graders (89%) than fourth graders (60%) \( (P < 0.001) \). On average, children spent a mean of 8 min/week (range 0–30) walking outdoors between classrooms and these numbers were higher for fourth graders (9.5 min/week; range 0–20) than for third graders (6.5 min/week; range 0–30) \( (P < 0.001) \).

A total of sixty-four children (8%) were excluded from analyses with dietary variables because of being under-reporters, fourteen (2%) because of being over-reporters, and twenty-three (3%) for recording their diet for \( < 4 \) d. Of the remaining sample, thirty-nine (6%) children consumed \( \geq 7.5 \mu \)g of dietary vitamin D/d (the official recommended intake at the time of the study\(^{41-43} \)), whereas thirty children (4%) consumed \( \geq 10 \mu \)g/d (the new official recommendation\(^{41-43} \)), 172 (25%) had an intake between 2.5 and 7.5 µg/d, and 470 (69%) consumed \( \leq 2.5 \mu \)g/d. Boys had higher daily intakes of energy, protein (energy %), vitamin D, Ca and milk (Table 2). Yet, because of their higher EI, the higher daily intake of dietary vitamin D in boys compared with girls was not retained when dietary vitamin D intake was calculated as µg/MJ. The same occurred with the intake of milk and dietary Ca.

An overall ninety children (11.5%) were categorised as immigrants-descendants, 44% of them were girls and 91% originated from a single foreign country only. Turkish roots were predominant along with roots in Pakistan and the former Yugoslavian Republic, accounting for 30, 18 and 17% of the immigrant/descendant children, respectively. The majority (80%) of the immigrant/descendant children were white; that is, only eighteen of the forty-one non-white children in the study had an immigrant/descendant background. Valid dietary recordings were...
Table 1. Characteristics of the study population according to sex
(Mean values and standard deviations; medians and 25th–75th percentile tested for sex differences)

<table>
<thead>
<tr>
<th></th>
<th>Girls</th>
<th>Boys</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>Mean ± SD</td>
<td>Mean ± SD</td>
<td></td>
</tr>
<tr>
<td>%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
<td>782</td>
<td>47.7 ± 52.3</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>782</td>
<td>142.3 ± 7.1</td>
<td>0.24</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>781</td>
<td>39.9 ± 34.3</td>
<td>0.06</td>
</tr>
<tr>
<td>25th–75th percentile BMI</td>
<td>781</td>
<td>29.7–38.5</td>
<td>0.08</td>
</tr>
<tr>
<td>25th–75th percentile</td>
<td></td>
<td>29.7–38.5</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Puberty stage (%)</td>
<td>757</td>
<td>53/36/10/0/3</td>
<td>0.50</td>
</tr>
<tr>
<td>White ethnicity (%)</td>
<td>779</td>
<td>95.2 ± 94.4</td>
<td>0.61</td>
</tr>
<tr>
<td>Immigrants/descendants (%)</td>
<td>782</td>
<td>10.7 ± 12.2</td>
<td>0.51</td>
</tr>
<tr>
<td>Parental education (%)</td>
<td>779</td>
<td>6/4/33/10/28/19</td>
<td>0.76</td>
</tr>
<tr>
<td>Sedentary activity (min/d)</td>
<td>756</td>
<td>470 ± 475</td>
<td>0.25</td>
</tr>
<tr>
<td>Moderate-to-vigorous activity (min/d)</td>
<td></td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>37</td>
<td>57</td>
<td></td>
</tr>
<tr>
<td>25th–75th percentile</td>
<td>26–48</td>
<td>39–73</td>
<td>0.24</td>
</tr>
<tr>
<td>Bicycling on weekdays (min/d)</td>
<td>750</td>
<td></td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Median</td>
<td>11</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>25th–75th percentile</td>
<td>4–18</td>
<td>6–19</td>
<td></td>
</tr>
<tr>
<td>Screen time on weekdays (min/d)**</td>
<td></td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>120</td>
<td>130</td>
<td></td>
</tr>
<tr>
<td>25th–75th percentile</td>
<td>75–155</td>
<td>90–185</td>
<td></td>
</tr>
<tr>
<td>Obligatory outdoor recesses during school hours (% of yes)</td>
<td>782</td>
<td>39.8 ± 6.2</td>
<td>0.26</td>
</tr>
<tr>
<td>Median</td>
<td>5</td>
<td>5</td>
<td>0.51</td>
</tr>
<tr>
<td>25th–75th percentile</td>
<td>0–13</td>
<td>0–10</td>
<td></td>
</tr>
<tr>
<td>Month of blood sampling (%)</td>
<td>782</td>
<td>38/18/44</td>
<td>0.47</td>
</tr>
<tr>
<td>Serum 25-hydroxyvitamin D (nmol/l)</td>
<td>782</td>
<td>54.9 ± 62.9</td>
<td>0.001</td>
</tr>
<tr>
<td>Whole-blood EPA + DHA (wt%)</td>
<td>782</td>
<td>3.6 ± 1.0</td>
<td>0.08</td>
</tr>
<tr>
<td>Supplements intake (%)</td>
<td>693</td>
<td>45/8/5/5/7/8/17</td>
<td>0.80</td>
</tr>
</tbody>
</table>

uw, Underweight; nw, normal weight; ow, overweight; ob, obese; A, ≤ lower secondary education; B, upper secondary education; C, vocational education; D, short higher education; E, Bachelor’s degree or equivalent; F, Master’s degree.
* Not including intake from supplements.
† Tanner stages as validated by Morris & Udrey(29).
‡ Determined as 4 grandparents + ≥ 1 parent born outside Denmark, motivated by definitions of immigrants and descendants used by Statistics Denmark(28).
§ Based on the highest parental education level in the household.
|| Based on age- and sex-specific cut-offs defined to pass through BMI of 18.5, 25 and 30 kg/m² at age 18 years, as described by Cole et al.(41,42).
|| Based on the highest parental education level in the household.
| Days per 7 d-registration with intake of a vitamin D-containing supplement, also including multivitamins.

Table 2. Dietary intake of the study population according to sex
(Mean values and standard deviations; medians and 25th–75th percentile tested for sex differences)

<table>
<thead>
<tr>
<th></th>
<th>Girls</th>
<th>Boys</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>Mean ± SD</td>
<td>Mean ± SD</td>
<td></td>
</tr>
<tr>
<td>Subjects (n)</td>
<td>7210</td>
<td>322</td>
<td>359</td>
</tr>
<tr>
<td>Energy (kJ/d)</td>
<td>8290 ± 1406</td>
<td>8290 ± 1406</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Protein (E%)</td>
<td>15 ± 16</td>
<td>2 ± 2</td>
<td>0.016</td>
</tr>
<tr>
<td>Fat (E%)</td>
<td>32 ± 33</td>
<td>4 ± 4</td>
<td>0.32</td>
</tr>
<tr>
<td>Carbohydrates (E%)</td>
<td>53 ± 52</td>
<td>5 ± 5</td>
<td>0.06</td>
</tr>
<tr>
<td>Median</td>
<td>1.7 ± 1.3</td>
<td>2.0 ± 1.5</td>
<td>0.002</td>
</tr>
<tr>
<td>25th–75th percentile</td>
<td>2.0–2.9</td>
<td>1.5–2.9</td>
<td></td>
</tr>
<tr>
<td>Dietary Ca (mg/d)*</td>
<td>843 ± 796</td>
<td>953 ± 796</td>
<td>0.001</td>
</tr>
<tr>
<td>Fish (total) (g/d)</td>
<td>12 ± 12</td>
<td>12 ± 12</td>
<td>0.48</td>
</tr>
<tr>
<td>Fatty fish (g/d)</td>
<td>3 ± 0</td>
<td>0 ± 0</td>
<td>0.24</td>
</tr>
<tr>
<td>Egg (g/d)</td>
<td>14 ± 14</td>
<td>14 ± 14</td>
<td>0.53</td>
</tr>
<tr>
<td>Milk (g/d)</td>
<td>329 ± 264</td>
<td>371 ± 264</td>
<td>0.002</td>
</tr>
</tbody>
</table>

* Not including intake from supplements.
handed in by fifty-eight (64%) of the immigrant/descendant children; this was fewer than the 90% in the non-immigrant group \((P<0.001)\). Immigrant/descendant children recorded a 571 kJ lower EI per day \((P=0.006)\), an approximately 0.6 µg lower intake of dietary vitamin D/d \((P=0.03)\) and an approximately 122 mg lower intake of dietary Ca/d \((P=0.001)\) compared with non-immigrant children. However, the differences in dietary vitamin D and Ca intake disappeared when calculated as µg/MJ and mg/MJ, respectively. No children carried veils or other cultural or religious attires.

**Serum 25-hydroxyvitamin D concentrations**

Mean serum 25(OH)D concentration for the total population \((n = 782)\) was 60.8 (so 18.7) nmol/l (range 15.2–132 nmol/l). Serum 25(OH)D concentrations \(\leq 50\) nmol/l were found in 222 (28.4%) of the children, of whom nineteen (2.4%) had concentrations <25 nmol/l (Fig. 1). As a subgroup, immigrant/descendant girls \((n = 40)\) had the highest deficiency prevalence; 80% had concentrations \(\leq 50\) nmol/l (Fig. 2). The proportion of immigrant/descendant children who were sampled in August/September, October and November, respectively, did not differ from that of non-immigrant/descendant children \((P=0.10)\), nor did the proportion of boys and girls \((P=0.47)\).

**Determinants of 25-hydroxyvitamin D concentrations**

Analysis on determinants of vitamin D status comprised \(n = 724\) children, except for analyses on bicycling \((n = 624)\) and dietary vitamin D and Ca intake \((n = 681)\). No interaction effects were found between sex and puberty, sex and ethnicity, or sex and month of sampling \((P=0.073, P=0.051\) and \(P=0.429\), respectively), whereas an interaction effect between sex and immigrant/descendant background was observed \((P=0.003)\). Consequently, the demographic model included age, proportion of days with intake of vitamin D-containing supplements, sex, ethnicity, immigrant/descendant background, puberty, month of sampling, parental education and the interaction between sex and immigrant/descendant background (Table 3).

In the subsequent analyses MVPA and outdoor walking between classrooms during school hours were found to be behavioural determinants of serum 25(OH)D (Table 3), whereas no associations were observed with obligatory outdoor recesses, BMI-for-age z score, sedentary time, amount of bicycling, dietary vitamin D and Ca intake, and whole-blood EPA+DHA concentrations.

In the demographic model adjusted simultaneously for the two significant behavioural determinants, MVPA and outdoor walking between classrooms, serum 25(OH)D increased 0.06 nmol/l \((95\%\ CI 0.01, 0.12; P=0.022)\) per min d in MVPA and 0.4 nmol/l \((95\%\ CI 0.1, 0.6; P=0.003)\) per min per week of walking outdoors between classrooms (Table 3). In this model, children sampled in November had a −13.3 nmol/l \((95\%\ CI −17.7, −9.1; P<0.001)\) lower serum 25(OH)D compared with those sampled in August/September (Table 3) and a −8.4 nmol/l \((95\%\ CI −12.6, −4.3; P=0.001)\) lower serum 25(OH)D concentration compared with those sampled in October (data not shown). Also, children sampled in October had a −5.2 nmol/l \((95\%\ CI −10.1, −0.4; P=0.035)\) lower serum 25(OH)D compared with those sampled in August/September (Table 3). Increasing age, being a girl, being non-white and being immigrant/descendant were also associated with lower serum 25(OH)D concentrations (Table 3). Children from households with a parental education level of at least a Master’s degree (≥17 years) were found to have a higher serum 25(OH)D level compared with children from households with parental level of lower secondary education (≤10 years) \(6.9\) (95% CI 1.0, 12.8 nmol/l; \(P=0.021\) (Table 3), with vocational education \((5.1; 95\%\ CI 2.1, 8.1\) nmol/l; \(P=0.001)\), with short higher education \((4.3; 95\%\ CI 0.2, 8.4\) nmol/l; \(P=0.039)\) and with Bachelor’s degree or equivalent \((4.5; 95\%\ CI 1.5, 7.6\) nmol/l; \(P=0.004)\) (data not shown). The 25(OH)D status in immigrant/descendant boys was estimated to be 9.4 nmol/l (95% CI 2.2, 16.6; \(P=0.010)\) higher than in immigrant/descendant girls. In comparison, the difference between non-immigrant boys and girls was 3.3 nmol/l (95% CI 0.7, 5.9; \(P=0.012\)).

The categorical vitamin D supplements variable constructed for sub-analysis on the association between frequency of vitamin D-containing supplements intake and serum 25(OH)D included \(n = 695\) children who recorded dietary intake for all 7 d (Table 1). The sub-analysis with this categorical vitamin D supplements variable in the demographic model included \(n = 673\) children. Here, we found higher serum 25(OH)D concentrations in those children who took vitamin D-containing supplements 3, 4, 5, 6 and 7 d/d compared with children who consumed supplements 0 out of 7 d \((P<0.001)\;\text{Fig. 3).}\) Serum 25(OH)D concentration was
found to be constant among children taking supplements for 0–2 d/7 d as well as among children taking supplements for 3–7 d/7 d ($P=0.64$). Consequently, the categorical supplements variable was collapsed into a dichotomous variable of 0–2 d/7 d and 3–7 d/7 d, respectively. When this collapsed variable was included in the demographic model, intake of vitamin D-containing supplements for 3–7 d ($n=282$) was associated with an increase of 9·0 nmol/l (95% CI 6·7, 11·2; $P<0.001$) as compared with intake for 0–2 d/7 d ($n=391$). A comparable estimate was observed, 8·7 nmol/l (95% CI 6·4, 11·0; $P<0.001$), when additionally adjusted for MVPA and outdoor walks.

**Discussion**

Approximately 70% of the school children in this cross-sectional study had serum 25(OH)D levels >50 nmol/l during autumn months. Consequently, concentrations ≤50 nmol/l were observed in almost one-third of the children at the time of year when concentrations are expected to be close to the annual peak. Compared with studies conducted at similar latitudes and time of year, mean 25(OH)D status in the present study (approximately 61 nmol/l) was fairly equal to that observed in 4–6-year-old Swedish children (60 nmol/l, August–September, 63°N) and 11–13-year-old Danish girls (60·3 nmol/l, August–September, 55°4’N). In contrast, mean 25(OH)D status in the present study was lower than in 4–18-year-old British children (71·2 nmol/l, June–November, 50–59°N) and 8-year-old Swedish children (82·8 nmol/l, summer, 57°N). Yet, between-study comparisons are problematic because of different methods of 25(OH)D analysis. Also, differences in, for example, fortification programmes between countries must be taken into account. Indeed, food fortification with vitamin D is neither mandated nor common in Denmark. On the basis of the well-established association between serum 25(OH)D and season at northern latitudes, the mean serum 25(OH)D concentration above sufficiency level in the present study matched the
Table 3. Determinants of serum 25-hydroxyvitamin D (β values and 95% confidence intervals)

<table>
<thead>
<tr>
<th>Demographic model* (n 724)†</th>
<th>Demographic model + MVPA (n 724)†</th>
<th>Demographic model + outdoor walks to classrooms (n 724)†</th>
<th>Demographic model + MVPA + outdoor walks to classrooms (n 724)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>β 95% CI  P</td>
<td>β 95% CI  P</td>
<td>β 95% CI  P</td>
<td>β 95% CI  P</td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-2.6 [-4.8, -0.4] 0.022</td>
<td>-1.8 [-3.8, 0.09] 0.062</td>
<td>-3.2 [-5.4, -0.9] 0.006</td>
<td>-2.9 [-5.1, -0.7] 0.009</td>
</tr>
<tr>
<td>Vitamin D-containing supplements‡</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10.6 [8.0, 13.3] &lt;0.001</td>
<td>10.5 [7.8, 13.1] &lt;0.001</td>
<td>10.5 [7.9, 13.2] &lt;0.001</td>
<td>10.4 [7.8, 13.1] &lt;0.001</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Girls Ref.</td>
<td>4.5 [2.1, 6.9] &lt;0.001</td>
<td>4.5 [2.1, 6.9] &lt;0.001</td>
<td>0.001</td>
</tr>
<tr>
<td>Boys Ref.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White Ref.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-white</td>
<td>-5.9 [-11.3, -0.5] 0.034</td>
<td>-5.7 [-11.2, -0.3] 0.038</td>
<td>-5.7 [-11.1, -0.3] 0.040</td>
</tr>
<tr>
<td>Background</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-immigrant descendant§</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immigrant/descendant</td>
<td>-16.7 [-22.3, -11.1] 0.001</td>
<td>-16.6 [-22.2, -11.0] 0.001</td>
<td>-16.3 [-21.9, -10.7] &lt;0.001</td>
</tr>
<tr>
<td>Immigrant/descendant × sex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Puberty¶</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not entered puberty Ref.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Entered puberty ( Tanner stage ≥2)</td>
<td>-0.2 [-2.6, 2.3] 0.876</td>
<td>-0.2 [-2.6, 2.3] 0.897</td>
<td>-0.1 [-2.6, 2.3] 0.921</td>
</tr>
<tr>
<td>Month of sampling</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>August/September</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>October Ref.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>November</td>
<td>-13.3 [-18.8, -7.8] &lt;0.001</td>
<td>-13.9 [-18.2, -9.6] &lt;0.001</td>
<td>-13.3 [-17.7, -9.1] &lt;0.001</td>
</tr>
<tr>
<td>Parental education**</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower secondary education (≤10 years)</td>
<td>6.0 [-2.0, 14.0] 0.141</td>
<td>6.4 [-1.6, 14.4] 0.115</td>
<td>6.3 [-1.7, 14.3] 0.125</td>
</tr>
<tr>
<td>Upper secondary education</td>
<td></td>
<td></td>
<td>6.7 [-1.3, 14.7] 0.103</td>
</tr>
<tr>
<td>Vocational education</td>
<td>1.6 [-4.1, 7.3] 0.578</td>
<td>1.9 [-3.8, 7.5] 0.514</td>
<td>1.6 [-4.1, 7.2] 0.590</td>
</tr>
<tr>
<td>Short higher education</td>
<td>2.4 [-4.0, 8.8] 0.460</td>
<td>2.6 [-3.8, 9.0] 0.433</td>
<td>2.5 [-4.0, 8.9] 0.453</td>
</tr>
<tr>
<td>Bachelor degree or equivalent</td>
<td>2.2 [-3.5, 7.9] 0.452</td>
<td>2.4 [-3.3, 8.1] 0.403</td>
<td>2.2 [-3.5, 7.9] 0.448</td>
</tr>
<tr>
<td>Master's degree (≥17 years)</td>
<td>6.6 [0.9, 12.7] 0.025</td>
<td>6.9 [1.0, 12.8] 0.022</td>
<td>6.8 [0.9, 12.7] 0.024</td>
</tr>
<tr>
<td>MVPA (min/d)††</td>
<td>0.06 [0.01, 0.1] 0.022</td>
<td>0.06 [0.01, 0.1] 0.022</td>
<td>0.06 [0.01, 0.12] 0.022</td>
</tr>
<tr>
<td>Outdoors walks between classrooms (min/week)‡‡</td>
<td>0.3 [-0.1, 0.6] 0.006</td>
<td>0.4 [-0.1, 0.6] 0.003</td>
<td></td>
</tr>
</tbody>
</table>

MVPA, moderate-to-vigorous physical activity; Ref., referent values.
* By linear mixed effects model with school, grade and class held as random effects and the above left column variables as fixed effects.
† Including the same n 724 children in all models hereof seventy-one children with immigrant/descendant background and thirty children with non-white ethnicity.
‡ Proportion of days with intake of a vitamin D-containing supplements/total days of dietary registration (%).
§ Determined as 4 grandparents + ≥1 parent born outside Denmark, motivated by definitions of immigrants and descendants used by Statistics Denmark(28).
∥ Interaction identified between immigrant/descendant background and sex.
¶ Tanner stages as validated by Morris & Udy(29).
** Based on the highest parental education level in the household.
†† MVPA ≥2296 cpm, measured by ActiGraph.
‡‡ Min/week the child had to spend outdoors on walking between classrooms during school hours.
expectation that serum 25(OH)D peaks in summer/autumn. And the seasonal fluctuation was further reaffirmed in the present study by the lower vitamin D levels measured in November compared with October and August/September, and was consistent with the hypothesis of November being the first month in the ‘vitamin D winter’ at these latitudes. Yet, our results also demonstrated that 25(OH)D insufficiency is not to be perceived solely as a winter phenomenon. Indeed, it is of particular concern that 24% of the children had autumn 25(OH)D concentrations below 25 nmol/l. As 25(OH)D concentrations are expected to decrease further during winter and early spring, these children are believed to be at risk for critically low winter status(16). This result underlines the relevance of an evaluation of potential demographic and behavioural determinants of children’s autumn vitamin D status. Indeed, although diverging cut-off levels for the optimal vitamin D level in children have been proposed and applied(50), the 25 nmol/l threshold is established on the basis of its clinically observed association with rickets and osteomalacia(1).

In the present study we did not find dietary intake to be associated with autumn serum 25(OH)D concentrations. This is consistent with other studies at similar latitudes with summer/autumn data(16,45), and underlines sunlight as the major source at consistent with other studies at similar latitudes with summer/autumn data(16,45). Sub-analysis with the children who registered for all 7 d/week. Adjusted for age, sex, white ethnicity, immigrant/descendant background, month of sampling and parental education, n 673. Statistical significance tested against reference: 0 days/week: * P < 0.05, ** P < 0.01, *** P < 0.001.

## Fig. 3. Association between intake of vitamin D-containing supplements in d/week and serum 25-hydroxyvitamin D (25(OH)D). Sub-analysis with the children who registered for all 7 d/week. Adjusted for age, sex, white ethnicity, immigrant/descendant background, month of sampling and parental education, n 673. Statistical significance tested against reference: 0 days/week: * P < 0.05, ** P < 0.01, *** P < 0.001.

status, also observable at the time of year when sunlight is abundant, but only if intake is frequent enough.

Among the demographic variables, our study pointed to non-white ethnicity, immigrant/descendant background, girl sex, and particularly the combination of girl sex and immigrant/descendant background as determinants of lower vitamin D status. The Danish authorities have identified certain population groups to be at risk for vitamin D deficiency. These include children 0–2 years of age, individuals with dark skin complexion, individuals who dress to cover most of the body during summer, individuals who rarely go outdoors and individuals who avoid sunlight(46). In addition to these officially identified groups, our results imply a need for also focusing on girls, children with an immigrant/descendant background, and older children such as adolescents and teenagers.

Other studies have found similar associations between immigrant background and low vitamin D status(22,25,49), but definitions of immigrant background vary, as does the suggested explanations for the association(50). In the present study, factors such as parental education, vitamin D supplementation, BMI, puberty stage, physical activity, screen time and dietary intake were all excluded as potential explanations of the association, and none of the children wore veils. Children from Turkey, the Middle East, Pakistan and India were categorised as white, and it cannot be ruled out that these children had darker skin tones than the white non-immigrant/descendant children. Yet, this does not explain that there was a difference between immigrant/descendant boys and girls. Interestingly, studies in Caucasian adolescent girls and women at similar latitudes as the present study have previously observed good correlation between usual sun habits and summer vitamin D status(16). We therefore speculate that behavioural factors such as time spent outside, area of skin exposed and overall sun-seeking behaviour, which was unfortunately not measured in the present study, may be culturally influenced, and different in children, particularly in girls, with an immigrant/descendant background(51,52). Unfortunately we obtained dietary recordings from only 64% of the immigrant/descendant children compared with 90% of the non-immigrant children. It is therefore also possible that actual differences in dietary vitamin D intake between immigrant and non-immigrant children are substantially larger than captured in this study.

The decrease in serum 25(OH)D with age in childhood has been observed in other studies(45,46,49), although not consistently(20,22). Our finding of higher status in boys compared with girls was not seen in previous studies conducted at similar northern latitudes(22,24,45). Varying behavioural patterns may likewise be relevant to consider in regard to these observed age and sex differences in serum 25(OH)D. Interestingly, an American study in children and adolescents has, for example, shown that 13-19-year-olds spend the least time outdoors compared with any other age group, and that teenage girls decreased their outdoor activities more dramatically than did teenage boys(53). Similar findings occurred in a UK population comparing 9–10-year-olds and 14–15-year-olds(54). We hypothesise that the association between age and serum 25(OH)D was mediated by such age-determined behavioural changes in outdoor activity in the present study, particularly because the association with age disappeared when MVPA was added to the demographic model.
Indeed, we speculate that MVPA may be positively associated with outdoor activity, but evidence on the nature of this association is needed.

Our results indicated that age was not a contributor to serum 25(OH)D status when MVPA was included in the demographic model, whereas age remained significant when outdoor walks was included. In our sample outdoor walks between classrooms was higher among fourth graders compared with third graders \((P<0.001)\). Hence, correlations and associations between these variables along with potential residual confounding are relevant considerations for interpretation of these outcomes.

To our knowledge the present study is the first to explicitly investigate the association between children’s vitamin D status and outdoor activity during school hours. Interestingly, and consistent with the hypothesis that only a few minutes of daily sun exposure is adequate to cover the vitamin D requirement, we found that even a few weekly minutes of outdoor walking between classrooms during school hours appeared to have a strong impact on the vitamin D status of children in autumn. This is also in accordance with the fact that school hours are placed at the time of day when sunshine is brightest. Contrary to outdoor walking between classrooms, obligatory outdoor recesses did not predict serum 25(OH)D concentrations. This was surprising. However, we speculate that children may spontaneously prefer to go outdoors during recesses at this time of year even if they are not obligated to do so by school policy, particularly as autumn weather is usually good in Denmark and all schools have outdoor facilities for children during recesses. Consequently, this ‘voluntary factor’ is likely to have affected the validity of the outdoor recess variable and might explain why an association was not observed even though it scientifically appears a likely determinant. Consequently, outdoor walks between classrooms is speculated to be a more valid measure of outdoor activity as it holds a clear non-voluntary behavioural necessity in order to get to class, and it proved robustly and highly associated with serum 25(OH)D. Inexpensive and easily applicable, brief intervals of outdoor activity during school hours may be worth investigating further as a way to safely and effectively improve vitamin D status in children, especially in older children.

The DiaSorin LIAISON assay applied to measure 25(OH)D in the current study has been shown to be the superior among automated immunoassays\(^{(55)}\). It has, however, previously been found to report lower 25(OH)D concentrations compared with the liquid chromatography-tandem spectrometry method considered the golden standard\(^{(56)}\). A risk of overestimation of low vitamin D status therefore exists in our study. The 2010 DEQAS survey found DiaSorin LIAISON to have a mean deviation from the all-laboratory-trimmed mean of minus 4 \(\pm 19 \) \(\text{nmol/l}\). As a result 24-4 \% of the children would then be categorised as insufficient and 1-9 \% as deficient compared with the respectively reported 28-4 and 2-4 \%, which is still a substantial part of the children. Also, DiaSorin LIAISON has been suggested to have a mean bias of only 0-5 \(\text{nmol/l}\) compared with the liquid chromatography-tandem spectrometry method\(^{(57)}\) – that is, a lesser underestimation than the one presented here.

The descriptive nature of our cross-sectional study naturally limits the ability to draw conclusions about causality. In view of the explorative nature of our study we have opted for reporting \(P\) values unadjusted for multiple comparisons in combination with cautious interpretation of results, which need to be reinforced through similar trends found in different analyses. The robust findings throughout the adjustments and models in Table 3 do not, however, suggest random findings. The risk of collinearity was addressed by testing each behavioural factor one at a time in the same basic demographic model. This approach also ensured transparency in the analyses. Still, although we were able to adjust for important potential confounders such as sex, age, month of sampling, parental education, supplemental intake, immigrant/descendant background and non-white ethnicity, there is always a risk of residual confounding in a cross-sectional study like the present. Records of direct sun exposure and overall behaviour regarding sun exposure would have been a strength, as factors such as sunny vacations and outdoor activities after school are possible residual confounders of the results.

There are several strengths to the present study. The extent and validity of determinants investigated, such as ≥4-day web-based dietary recordings and validated objective accelerometry measurements of physical activity along with the explorative inclusion of innovative determinants regarding outdoor activity during school hours, makes this study unique. In addition, data were collected at the time of year when behavioural factors are indeed speculated to be of particular importance at this latitude. Importantly, the study population was substantially larger than in previous Scandinavian studies, and a specific goal was representativeness of the study sample. Recruitment was conducted from nine geographically and demographically varying areas. The enrolment rate was high at 82 \%, and, although the proportion of non-white ethnicity in our sample might appear small, it is considered fairly representative of the Danish population as a whole. All parental education levels were present, and the 11-5 \% of children characterised as immigrants/descendants corresponded well with the 10-1 \% of immigrants/descendants reported in the general Danish population\(^{(58)}\). Also, distribution of country of origin of the included immigrant/descendant children was in accordance with that found in the general Danish population\(^{(59)}\). Similarly, the prevalence of dietary supplement users in our study population, 55 \% among girls and 50 \% among boys, appears to mirror well that of the general population, although our variable included vitamin D-containing supplements only, hereunder also multivitamins. Indeed, in national surveys, 46 and 55 \% of 11–14-year-old Danish girls and boys and 60 and 51 \% of adult Danish women and men reported taking dietary supplements\(^{(59,60)}\). All together, this indeed indicates a high degree of representativeness in our study.

In conclusion, almost one-third of the 8–11-year-old Danish children in this large representative study population did not reach the recommended 25(OH)D sufficiency level of 50 \(\text{nmol/l}\) during autumn. This raises concern as status is expected to drop further during winter, and indicates that vitamin D deficiency might not be regarded solely as a winter phenomenon in children at northern latitudes. Older age, female sex, non-white ethnicity and immigrant/descendant background, particularly in females, are all
factors associated with lower vitamin D status and therefore relevant when evaluating the determinants of vitamin D status in children. Frequent intake of vitamin D-containing supplements in autumn was an observable positive determinant of autumn vitamin D status in the children. Interestingly, increased MVPA and brief outdoor walks between classrooms during school hours were also found to be positive behavioural determinants of vitamin D status. In contrast, obligatory outdoor recess was not. Yet, the validity of this variable was potentially affected by children who chose to go outdoors during recesses even when it was not obligatory by school policy. Hence, more valid data on outdoor activity during recesses are needed in future studies, and these potential behavioural initiatives to improve autumn vitamin D status in children should be investigated further in randomised trials.

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References


