Maternal vitamin D status affects bone growth in early childhood—a prospective cohort study

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Methods Eighty-seven children were followed from birth to 14 months. Background data were collected with a questionnaire and a 3-day food record. At 14 months bone variables were measured with peripheral computed tomography (pQCT) from the left tibia. Serum 25-OHD and bone turnover markers were determined. Findings were compared with maternal vitamin D status during pregnancy.

Results The children were divided into two groups based on vitamin D status during pregnancy. Despite discrepant S-25-OHD at baseline (median 36.3 vs. 52.5 nmol/l, \( p < 0.001 \)), the values at 14 months were similar (63 vs. 66 nmol/l, \( p = 0.58 \)) in Low D and High D. Serum 25-OHD increased more in Low D (\( p < 0.001 \)) despite similar total intake of vitamin D (mean 12.3 \( \mu \)g/day). In Low D, tibial bone mineral content (BMC) was lower at birth but BMC gain was greater (multivariate analysis of variance [MANOVA] \( p = 0.032 \)) resulting in similar BMC at 14 months in the two groups. In High D, tibial total bone cross-sectional area was higher at baseline; the difference persisted at 14 months (MANOVA; \( p = 0.068 \)). Bone mineral density (BMD) and \( \Delta \)BMD were similar in the two groups.

Conclusions Postnatal vitamin D supplementation improved vitamin D status but only partly eliminated the differences in bone variables induced by maternal vitamin D status during the fetal period. Further attention should be paid to improving vitamin D status during pregnancy.

Keywords Bone growth · Development · Maternal vitamin D status · Pediatric · Vitamin D
fracture risk later in life [1–3]. During the fetal period, it is likely that metabolic and endocrine systems are programmed to allow the fetus to adapt to the in utero environment [4]. Vitamin D is a seco sterol that modifies various biological functions in the body [5], and researchers have identified 37 target organs for vitamin D [5]. Low maternal vitamin D status or inadequate dietary vitamin D intake during pregnancy predisposes children to asthma and allergic rhinitis [6], diabetes [7], acute lower respiratory infection [8], and impaired bone mass accrual. This is evidenced by smaller bone cross-sectional area (CSA) and bone mineral content (BMC) at birth [9, 10] and at 9 years of age [11].

Programming of skeletal growth may occur through growth hormone—IGF-I axis [4, 12], whereas bone quality may be determined by factors related to differentiation of mesenchymal stem cells [13, 14]. The intrauterine environment strongly affects growth rate in infancy, but may also influence growth in puberty [15]. The extent to which changes in nutrient supply between intrauterine and postnatal periods affect growth and development, per se, has not been well established [4]. The most critical views predict that intrauterine nutritional deficits have permanent consequences and that a newborn’s metabolism may not adapt to improved nutritional status; the nutrients may not be utilized efficiently and the risk for disease may be maintained despite improved nutritional status [16]. However, postnatal catch-up occurs in linear growth if the fetal deprivation and its timing and magnitude have not been too critical [17].

Previously the authors of the current study have reported that during the pregnancy, 69% of the women and 37% of the newborns at birth were vitamin D deficient (defined in women as S-25-OHD <50 nmol/l [18, 19] and in the newborn as <37.5 nmol/l [20]). The newborn bone variables were measured with peripheral quantitative computed tomography (pQCT) during the hospital stay. Based on these results, it was concluded that maternal vitamin D status affects bone mineral accrual and influences bone size during the intrauterine period [10].

The present prospective study had two objectives. The first was to follow whether the vitamin D-associated skeletal effects observed at birth persist during the first year of life, and the second was to evaluate the impact of postnatal vitamin D status on bone growth and development in early infancy.

Subjects and methods

Families (N=124) were initially recruited and assessed between October and December 2007 during their labor visit to the birth hospital. They were invited for a follow-up visit approximately 14 months later, between February and March 2009. The recruitment of families has been described in detail elsewhere [10]. Only primiparous mothers who were healthy, non-smoking, aged between 20 and 40 years, of Caucasian origin, and had an uneventful, singleton, full-term pregnancy (37–42 weeks) were included. The study protocol was approved by the Ethics Committee of Helsinki University Hospital. All mothers gave their written informed consent in accordance with the Declaration of Helsinki. Maternal vitamin D status was assessed in communal prenatal clinics during the first trimesters as part of normal follow-up. A second, fasting blood sample from the mother was collected 2 days postpartum during the hospital stay between late October and mid-December 2007. At birth, cord blood was obtained from the umbilical vein after cord clamping in 81 subjects. Background data was collected through an extensive questionnaire. Records on pregnancy follow-up and the birth report were obtained, including birth weight, length and head circumference measured by midwives, and duration of the pregnancy. Birth lengths and weights were transformed into Z-scores using Finnish sex-specific normative data for fetal growth [21]. One newborn and her mother were excluded from the initial analysis due to intrauterine growth retardation.

Eighty-seven (70%) of the original cohort of 124 families agreed to participate in the follow-up visit. Mothers in families agreed on follow-up tended to be younger (p<0.1), they were more educated (p=0.09) and had smaller family (p=0.08) than non-participants, but there were no differences in any pregnancy outcomes. Before the 14-month visit, the families received an extensive questionnaire concerning the child’s health and medical history, sunshine exposure, and use of vitamin supplements. The questionnaire included a 3-day food record. During the study visit, one of the researchers interviewed the family about the child’s development, including motor and language skills. Of those who agreed to participate in the follow-up visit, all but three returned the questionnaire. Anthropometric measurements were obtained for each subject. Height was measured at standing position with a wall-mounted height measuring scale and rounded to the nearest 0.1 cm. Weight was measured while sitting on a scale in light clothing and rounded to the nearest 0.1 kg. Heights were transformed into Z-scores and weights were expressed as height-adjusted weights according to Finnish sex-specific normative data for infants [21]. The newborn bone variables were measured with peripheral computed tomography (pQCT) from the left tibia. A blood sample was obtained for laboratory analyses from all but one child. Local anaesthetic patches (EMLA R; AstraZeneca AB, Södertälje, Sweden) were used to reduce the discomfort of venipuncture.

Dietary intakes were calculated from 3-day food records with Diet32 software (Aivo Oy Finland, Turku, Finland).
The nutrient contents of the foods was based on the Finnish National Food Composition Database, Fineli, version 2001, maintained by the National Public Health Institute of Finland, Nutrition Unit. The total intake of vitamin D included intake from diet and from supplements.

Laboratory measurements

Serum 25-OHD was measured with an OCTEIA immunoenzymometric assay (IDS, Bolton, UK). The intra-assay coefficient of variation (CV) was less than 3.9% and interassay variation (4.5%). Reproducibility was ensured by adhering to the Vitamin D External Quality Assessment Scheme (DEQAS). EIA results were compared with HPLC results in order to determine the reliability of EIA in measuring 25-OHD2 concentration. The results were consistent ($r=0.751$, $p<0.001$, $R^2=0.495$); therefore, the EIA results were used throughout the study. Vitamin D status in children was defined as deficient when S-25-OHD was below 37.5 nmol/l, insufficient when it was between 37.6 and 50 nmol/l, and sufficient when it was above 50 nmol/l, according to the published pediatric reference values [20]. In adults, a concentration of at least 80 nmol/l is considered optimal for multiple health outcomes [22].

Serum bone-specific alkaline phosphatase (S-BALP) was assayed with an OCTEIA Octase BAP immunoenzymometric assay (IDS) in order to characterize bone formation. Samples were diluted 1:5 to meet the standard curve. Intra- and interassay CVs were 6.1% and 6.7%, respectively. The bone resorption marker, serum active isoform 5b of the tartrate-resistant acid phosphatase (S-TRACP), was determined with a bone TRAP assay (SBA Sciences, Turku, Finland). Intra- and interassay CVs were 1.2% and 3.0%, respectively.

pQCT bone measurement

Peripheral bone variables were determined by pQCT from the left tibia. One 2.5-mm slice (voxel size, 0.4 mm) at the 20% site of distal tibia, was measured with a XCT-2000 scanner (Stratec, Pforzheim, Germany) as described previously [10]. Data was analyzed using version 5.50 of the manufacturer’s software package, in which the bone contour was analyzed with a single threshold of 180 mg/cm3 for the detection of total bone mineral density (BMD), BMC, and CSA. The long-term CVs for the phantom BMD and CSA were 1.9% and 1.1%, 2.7% and 0.79%, and 0.50% and 0.78% in the total, cortical, and trabecular bone, respectively. Short-term precision (CV%) was determined with duplicate measurements of five subjects. CVs for the total bone BMD and CSA were 6.0% and 6.5%, respectively. On this basis, the calculated least significant changes for total bone BMD and CSA were 16.7% and 18.1%, respectively. Out of all the pQCT measurements at 14 months, 67 (78%) were successful.

Statistical methods

Statistical analyses were performed with SPSS version 16.0 for Windows (SPSS Inc., Chicago, IL). The two groups were compared using an independent samples t-test. Repeated-measures ANOVA was applied to follow 25-OHD, BMC, CSA, BMD, BALP and TRACP between baseline and the 14-month visit. These time-points were compared using contrasts. Determinants for bone analysis were identified with Pearson correlations. Where necessary, variables were transformed using logarithms in order to satisfy statistical assumptions of normality. Differences between groups in BMC, CSA and BMD at 14 months, as well as in ΔBMC, ΔCSA and ΔBMD (change from birth to 14 months), were tested with multivariate analysis utilizing the same confounding factors. Results are presented as mean (SD) unless otherwise indicated. Results were considered significant when $p<0.05$; $p$ values between 0.05 and 0.10 were considered trends.

Results

A total of 87 children (57% boys) were followed up for 14 months. Their mean (SD) values for age, weight, height-adjusted weight, height, and height Z-score were 14.8 (0.5) months, 10.8 (1.3) kg, 0.68 (7.6)%, 78.6 (3.2) cm, and 0.11 (1.1), respectively. For data analysis, the participants were divided into two groups based on maternal vitamin D status during pregnancy. The median maternal S-25-OHD value, 42.6 nmol/l, was used as the cutoff to define two equal-sized groups of children with below-median (=Low D; mean S-25-OHD 35.7 [5.0] nmol/l) and above-median (=High D; mean S-25-OHD 54.9 [9.1] nmol/l) maternal S-25-OHD concentration. Table 1 presents the background characteristics of these two groups at baseline and at the 14-month follow-up. The duration of exclusive was similar in groups (see Table 1). Eighteen children (21.7%) were still breastfed at the time of the follow-up visit. Dietary intakes of energy, protein, vitamin D and calcium did not differ between the groups and all children had normal development. Only the age when the children started to walk with support differed between the groups; all other developmental milestones were similar.

Despite lower vitamin D concentration during pregnancy and at birth in Low D than in High D (means 35.7 vs. 54.2 nmol/l, and in the cord 40.5 and 59.3 nmol/l, independent samples t-test; $p<0.001$, respectively), the 25-OHD concentrations in the two groups at 14 months were similar (63 vs. 66 nmol/l, $p=0.58$). Serum 25-OHD increased from mean pregnancy value and cord more in
Low D than it did in High D (28 vs. 10 nmol/l and 23.6 vs.
6 nmol/l, respectively; independent samples t-test; p<
0.001), although the total intake of vitamin D was similar,
at an average of 12.3 (3.0) μg/day (Table 2). The total
intake of vitamin D correlated positively with 25-OHD
concentration in the whole cohort (r=0.301, p=0.005) and
in High D (r=0.505, p<0.001), but not in Low D (r=0.219,
p=0.168) (Fig. 1). Vitamin D status according to several
reference values [20, 22] did not differ between the groups.
Of the total cohort, 21%, 62% and 17% had S-25-OHD
below 50 nmol/l, between 50 and 79.9 nmol/l or at least
80 nmol/l, respectively (Table 3). Higher dietary intake of
vitamin D and use of D3 supplements were related to
improved vitamin D status.

Tibia BMC, CSA and BMD

Bone measurements were successful in 68 of subjects (78%) at
the 14-month visit. For the longitudinal bone analysis, complete
baseline and follow-up data were available for 29 subjects in
Low D and 26 subjects in High D. Determinants of bone
variables were gender, birth weight Z-score, walking age,
duration of exclusive breastfeeding and S-25-OHD at 14 months. At the 14-month visit, boys had a higher BMC, ΔBMC and BMD than girls (independent samples t-test; p=0.002, p=0.002 and p=0.02, respectively). Birth weight Z-score correlated strongly with BMC and CSA at 14 months (r=0.507, p<0.001 and r=0.368, p=0.004). Similarly, walking age was inversely associated with BMC, CSA and S-25-OHD at 14 months (r=−0.545, p<0.001, r=−0.433, p<0.001, and r=−0.194, p=0.083, respectively). The duration of exclusive breastfeeding correlated negatively with BMC, ΔBMC, CSA and ΔCSA (r varying from −0.377 to −0.428, p=0.002). S-25-
OHD at the 14-month visit was only modestly related to BMD
and ΔBMD (r=−0.230, p=0.08 and r=−0.142, p=0.250), but
was included in the model as well.

The development of BMC from baseline to 14 months
differed between the groups (repeated-measures multivariate
analysis of variance [MANOVA]; p=0.023) (Fig. 2a) due to
greater baseline BMC in High D. However, the total BMC

Table 1 Background characteristics and changes in them from baseline value given as mean (SD)

|                        | Low D | High D | Independent samples t-test |
|------------------------|-------|--------|---------------------------|
| N                      | 44    | 43     | 0.336                      |
| Age, months            | 14.9 (0.5) | 14.8 (0.5) | 0.842                     |
| Males, %               | 58    | 55     | 0.842a                     |

**Anthropometric and growth variables**
- Weight, kg
- Relative weight
- ΔWeight, kg
- Weight velocity, g/month
- Height, cm
- Height Z-score
- ΔHeight, cm
- Height velocity, cm/month

**History of breast feeding and dietary intakes**
- Duration of exclusive breastfeeding, months
- Currently breastfed, N (%)
- Energy intake, kcal/day
- Fat intake, g/day
- Protein intake, g/day
- Calcium intake, mg/day
- Total intake of vitamin D, μg/day

**Motor and language skills**
- Age when learnt to crawl, months
- Age when learnt to stand, months
- Age when learnt to walk with support, months
- Age when learnt to walk without support, months
- Number of words in use

* Pearson chi square
gain ($\Delta \text{BMC} = \text{BMC}_{14\text{ month}} - \text{BMC}_{\text{baseline}}$) during the first year was 0.062 (SEM=0.029) g/cm greater in Low D (MANOVA; $p=0.032$); consequently, no difference was observed in BMC between the groups at the 14-month visit. TB CSA from baseline to the 14-month visit was significantly higher in High D than in Low D (repeated MANOVA; $p=0.004$) (Fig. 2b) due to the higher baseline CSA in High D. $\Delta \text{CSA}$ did not differ between the groups. Thus, a trend to higher CSA at 14 months by 14.6 (SEM=7.8) mm$^2$ (MANOVA; $p=0.068$) remained in High D. There was no difference between the groups in BMD during the 14 months (Fig. 2c) or in $\Delta \text{BMD}$. The observed decrease in BMD is a consequence of a greater increment in CSA compared with the gain in BMC (69% vs. 18%).

Bone turnover markers

BALP, a surrogate of bone formation, increased dramatically from baseline (repeated measures MANOVA; $p<0.001$) (Fig. 3a), while TRACP concentration remained at the same level during the 14 months (Fig. 3b). At the 14-month visit, TRACP, BALP, or their ratio did not differ between the groups. There was no correlation between BALP and TRACP, but $\Delta \text{TRACP}$ correlated positively with $\Delta 25\text{-OHD} (=25\text{-OHD}_{14\text{ month}} - 25\text{-OHD}_{\text{pregnancy\ mean}})$ ($r=0.345$, $p=0.012$). Correspondingly, $\Delta \text{BALP}$ correlated inversely with $\Delta 25\text{-OHD}$ ($r=-0.213$, $p=0.034$). The correlations were similar in both groups.

Discussion

This prospective study made three key findings. Firstly, distal tibia CSA remained larger at 14 months in infants with higher maternal vitamin D status during pregnancy than in infants with lower maternal vitamin D status. Secondly, the increment in tibial BMC from birth to 14 months was higher in those with inferior maternal vitamin D status during pregnancy. This resulted in similar BMC and BMD at 14 months in both study groups. Finally, 20% of the children had S-25-OHD below 50 nmol/l at 14 months of age, although their median total intake of vitamin D was 12.2 (3.0) $\mu$g, which meets the Nordic recommendation for this age group [23].

Other interesting findings related to bone growth in this prospective cohort were that boys had higher BMC, and BMC increased more during the 14 months and resulted in higher volumetric BMD in distal tibia than in girls. Children in high vitamin D group learnt to walk with support later than children in low vitamin D group.

### Table 2

|                      | Low D | High D | Independent samples t-test |
|----------------------|-------|--------|----------------------------|
| $N$                  | 46    | 40     |                            |
| Mean of first trimester and postpartum maternal 25-OHD, nmol/l | 35.7 (5.0) | 54.2 (9.1) | <0.001 |
| Cord 25-OHD, nmol/l  | 40.3 (7.2) | 59.5 (12.2) | <0.001 |
| At 14-month S-25-OHD, nmol/l | 63.0 (20.7) | 65.6 (21.2) | 0.575 |
| $S$-25-OHD$^a$/total 25-OHD$^a$ | 0.50 (0.28) | 0.50 (0.24) | 0.878 |
| $\Delta$S-25-OHD$^b$, nmol/l | 27.5 (22.2) | 10.2 (19.4) | 0.001 |
| $\Delta$S-25-OHD$^c$, nmol/l | 23.0 (23.2) | 6.0 (22.1) | 0.002 |
| S-TRACP, U/l         | 11.2 (4.0) | 10.0 (4.1) | 0.199 |
| $\Delta$S-TRACP, U/l | $-0.28$ (4.3) | $-0.47$ (4.7) | 0.876 |
| S-BALP, $\mu$g/l    | 124 (38) | 122 (38) | 0.847 |
| $\Delta$S-BALP, $\mu$g/l | 69.2 (37.4) | 62.4 (42.8) | 0.527 |

$^a$ Based on HPLC

$^b$ An increment of S-25-OHD from mean maternal to 14-month visit

$^c$ An increment of S-25-OHD from cord to 14-month visit; $N=30$, $N=31$

![Fig. 1](https://example.com/figure1.png)

**Fig. 1** Total intake of vitamin D correlated positively with serum 25-OHD at 14 months (r=0.505, p<0.001), but not in Low D (r=0.219, p=0.168). Squares indicate High D and circles indicate Low D.
although other developmental milestones were similar. We consider this as a random finding because it is unlikely that higher maternal vitamin D status would contribute to this and several studies have witnessed that vitamin D deficiency is related to delayed age of walking [24, 25]. In this study, walking age without support was inversely related to tibia BMC and CSA, suggesting that earlier walking enhances bone development. Similarly jumping is shown increase the outer diameter of the tibia in a randomized controlled trial of 3- to 5-year-old children [26]. Walking is one of the first weight-bearing exercises modifying the strength of the tibia, but it is unsure if the association between walking age and bone health will preserve in the future. Surprisingly, longer exclusive breastfeeding was linked to lower bone development, which might be a sum of prolonged growth rate [27] and possible lower intakes of nutrients. However, infant 25-OHD at 14-month visit was only modestly related to bone growth; thus, it was included as a confounder in the model.

Maternal vitamin D status regulates skeletal growth and development during fetal life [9, 10, 28]. The present study proves that these effects partly persist in early childhood, as has been suggested in a longer prospective study [11]. Tibia CSA remained somewhat larger in infants whose mothers had better vitamin D status during pregnancy. Besides genetic background bone size is affected by various hormones and it has been shown that growth hormone-IGF-1 axis is responsible for bone size [29] and periosteal expansion [30, 31]. Leptin may favor stem cell differentiation towards osteoblasts rather than adipocytes [32] in infancy. Furthermore, vitamin D stimulates osteoblastogenesis in human mesenchymal stem cells and production of IGF-1 in osteoblasts [14]. In infants with rickets vitamin D supplementation increases serum IGF-1 and accelerates linear growth [33]. In this study, we did not measure IGF-1 and other growth hormone parameters, but height and weight velocities did not differ between the groups. Although all infants received vitamin D supplementation, the difference between the groups in tibia CSA was maintained until 14 months of age. The difference at birth was 16% and 11% at 14 months. This shows that the fetal bone growth tracks during the first year [34], which emphasizes the meaning of maternal nutrition for bone trajectory. Bone size is a major determinant of bone strength [35] and therefore the observed differences in CSA may have significant clinical implications in fracture resistance.

Unlike CSA, BMC or BMD did not differ between the study groups at the 14-month visit. This is explained by the steep increment of BMC in Low D group. In fact, the BMC accrual was about three times higher in Low D than in High D (28.7% vs. 8.4%), and due to this catch-up in Low D there was no difference between the groups in distal tibia BMC at the 14-month visit. The greater increase in BMC in Low

Table 3 Distribution of vitamin D status in 1-year-old children and mean intake in each category

| S-25-OHD (nmol/l) | N (%) | Total intake of vitamin D (μg/day)* | User of D2 (%)* |
|-------------------|-------|-----------------------------------|----------------|
| <37.5             | 2 (2.3) | 12.2 (4.0)                   | 100            |
| 37.6–50           | 16 (18.6) | 10.6 (3.1)                   | 100            |
| 50.1–79.9         | 53 (61.6) | 12.4 (2.8)                   | 88.5           |
| ≥80               | 15 (17.4) | 13.1 (2.5)                   | 50             |
| Total (N)         | 86              | 84                           | 84             |

*Total intake differed between categories of 25-OHD (ANOVA; p=0.03)

*Distribution of D2 users differed between categories (chi square; p=0.001)

![Fig. 2](image-url)

Fig. 2 BMC, CSA and BMD in study groups from baseline to 14 months. Increase in BMC from baseline to 14 months differed between the groups (repeated-measures MANOVA; p=0.023) (a) due to higher baseline BMC in High D. No difference was observed in BMC between the groups at the 14-month visit. TB CSA from baseline to 14 months was significantly higher in High D than in Low D (repeated-measures MANOVA; p=0.004) (b) due to the higher baseline CSA in High D. At 14 months, CSA remained 14.6 (SEM=7.8) mm² (MANOVA; p=0.068) higher in High D. There was no difference between the groups in BMD during the 14 months (c) or in ΔBMD. Low D and High D groups are represented by circles and squares, respectively. Error bars represent SEMs.
D group is likely to be due to increased calcium accrual, as reverted vitamin D status enhances calcium absorption. Some studies have witnessed that insufficient vitamin D status during pregnancy is related to lower bone mineral status in the newborn [9, 10, 28]. Initially, 70% of the mothers were vitamin D deficient during the pregnancy as their mean serum 25-OHD for first trimester and 2 days postpartum was less than 50 nmol/l. Improved postnatal vitamin D status results in catch-up in BMC, but not in CSA. Decline in BMD was similar in both study groups and it reflects a redistribution of bone tissue from the endocortical to the periosteal surface. An explanation for declined BMD might be that cortical thickness decreased during the 14 months while the amount of trabecular bone increased [36].

In early infancy, peripheral bones grow by increasing the outer diameter rather than the mineral content [36]. In the whole group tibia CSA increased dramatically (68%), indicating that the periosteal diameter and circumference increased. The growth in periosteal circumference occurred similarly in groups, but High D group started at a higher level and hence stayed higher at 14-month visit.

Vitamin D supplementation is recommended for all infants aged between 2 weeks and 3 years in Nordic countries in order to guarantee a total intake of 10 μg/day. All subjects in the present study received supplementation, compared to a representative study cohort in Finland, in which 85% of 1-year-old infants and 70% of 2-year-old infants were reported to receive vitamin D supplementation [37]. Thus, families in the present study were somewhat selected and possibly more health-orientated than the Finnish population in general. In the present study, 85% of infants had total vitamin D intake that was in line with the Nordic recommendation [23]. Interestingly, the use of D3 supplements was associated with improved vitamin D status to a greater extent than use of D2 supplements, which is in line with findings of Houghton and Vieth [38]. However, the number of D3 users was very low (N=12), which means that further comparison between different forms of vitamin D is not justified. Because of vitamin D supplementation, S-25-OHD concentration increased during the follow-up. Interestingly, the increase was higher in group with inferior S-25-OHD during pregnancy than in group with higher 25-OHD during pregnancy (ΔS-25-OHD 27.5 vs. 10.2 nmol/l). In line with earlier findings [39, 40], a higher response was observed in those with initially lower status. However, neither S-25-OHD nor ΔS-25-OHD was significantly associated with pQCT bone variables at 14 months or their changes during the 14-month follow-up. The study shows that fetal vitamin D status, rather than postnatal vitamin D status, affects bone growth during the first year. On the other hand, S-25-OHD reflects relatively short-term accumulation of dietary vitamin D and solar exposure [41], whereas observing differences in bone variables takes more time. ΔS-25-OHD correlated positively with ΔS-TRACP and inversely with ΔBALP suggesting that vitamin D affects bone turnover [42]. Consequently, S-25-OHD may be a significant determinant of bone turnover in infants, although growth, diet and motor development also play a part.

There was a positive association between total intake of vitamin D and 25-OHD in the entire group and in High D, but not among those infants in Low D whose vitamin D status during pregnancy was worse. At the 14-month visit, 2.3%, 18.4% and 79.3% were defined as vitamin D deficient, insufficient and sufficient, respectively [20]. Given that more than 20% of the infants had S-25-OHD below 50 nmol/L, despite compliance with supplementation, higher intake of vitamin D is recommended in order to obtain all the potential health benefits of vitamin D [43, 44]. Optimally, vitamin D sufficiency should be guaranteed during fetal life as the findings of the present study suggest that the consequences of poor vitamin D status during the fetal period cannot be totally reverted by postnatal vitamin D supplementation.

One of the limitations of the study was that only 70% of the families were willing to attend the 14-month follow-up...
visit. Furthermore, only 78% of the pQCT measurements at 14 months were successful, which resulted in problems with the sample size in data analysis. A sample size of 35 per group would have been required in order to reach sufficient statistical power. Only total bone parameters were measured with pQCT from the 20% site of tibia. This site contains both cortical and trabecular bone, but we did not quantify those separately because the cortical thickness is relatively small compared to voxel size and partial volume effect obscured the results. However, the strength of this study was a prospective study design with antenatal vitamin D status.

It can be concluded that postnatal vitamin D supplementation improved vitamin D status in infants and partly eliminated the differences in bone variables that had resulted from maternal vitamin D status during the fetal period. The difference remained in total bone CSA, while it disappeared in BMC. It seems unlikely, therefore, that improving vitamin D intake merely in infancy would revert the consequences of poor vitamin D status during the fetal period. Based on these observations, additional efforts should be made to improve vitamin D status during pregnancy.

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Conflicts of interest None.

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