CLINICAL STUDY

Preterm birth does not affect bone mineral density in young adults

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Abstract

Objective: Previous studies showed conflicting data on the effect of prematurity on bone mineral density (BMD) in infants and children. Only a few studies investigated the long-term effects of prematurity on BMD in early adulthood. The objective of our study was to assess the long-term effects of preterm birth on BMD of the total body (BMDTB), lumbar spine (BMDLS) and bone mineral apparent density of the LS (BMADLS).

Design: Cross-sectional study.

Methods: It consists of two hundred and seventy-six healthy subjects without serious postnatal complications, aged 18–24 years. The contribution of gestational age to the variance in BMD in young adulthood and the differences in BMD between 151 subjects born preterm (median gestational age 32.2 weeks (interquartile range (IQR) 30.3–34.0)) and 125 subjects born at term (median gestational age 40.0 weeks (IQR 39.0–40.0)) were investigated. BMD was determined by dual-energy X-ray absorptiometry.

Results: There were no significant linear correlations between gestational age and BMDTB ($r=0.063$, $P=0.30$), BMDLS ($r=0.062$, $P=0.31$) and BMADLS ($r=0.069$, $P=0.26$). Also after adjustment for possible confounders, gestational age was no significant contributor to the variance in BMDTB ($P=0.27$), BMDLS ($P=0.91$) and BMADLS ($P=0.87$). No significant differences were found between preterm and term subjects with regard to BMDTB, BMDLS and BMADLS.

Conclusion: In our cohort of 276 young adults, aged 18–24 years, gestational age was not a significant determinant in the variance of BMD. Preterm birth without serious postnatal complications is not associated with a lower BMD in young adulthood.

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Introduction

Decreased mineralization of osteoid tissue during the early postnatal period is a known complication of very low birth weight and/or prematurely born infants (1–3). It comprises a variety of disturbances ranging from mild undermineralization to frank radiological rickets with fractures (4). Preterm infants are at an increased risk of low bone mineral density (BMD) as bone mineralization, along with calcium and phosphorus accretion, mainly occurs during the third trimester of pregnancy (5).

Previous studies showed conflicting data on the effect of preterm birth on BMD in infants and children. Prematurity, irrespective of birth weight, was found to be associated with lower BMD in infancy and early childhood (1, 6–9). In contrast, other studies in young children could not show differences in BMD due to prematurity (2, 10).

Owing to advances in neonatal care, survival of preterm and very low birth weight infants has significantly improved and an increasing number of these children reach adulthood. In 2007, the preterm birth rate in the United States was 12.7%, which corresponds with ~550,000 preterm births per year (11). Moreover, this percentage is about 20% higher than the preterm birth rate in 1990 (11). It is, therefore, of increasing importance to assess the effect of prematurity on BMD in adulthood. Previous studies investigated the long-term effects of low birth weight and growth on BMD (12–20), but very few have studied the specific contribution of the duration of gestation on later BMD (14, 21).

BMD in later life depends largely on the peak bone mass achieved in early adulthood and the subsequent bone loss (22). A high peak bone mass provides a larger reserve later in life (22, 23). Osteoporosis is an important and increasing cause of morbidity and mortality in the developed countries. It is characterized by low bone mass, deterioration of bone tissue and disruption of bone architecture, compromised bone strength and an increase in the risk of fracture (24). According to the
WHO diagnostic classification, osteoporosis is defined by BMD at the hip or spine that is $\leq 2.5$ SDS below the young normal mean reference population (24). Since early prevention of osteoporosis is likely to be more successful than treatment of an already established disorder, it is essential to identify potential risk factors.

We hypothesized that prematurity is associated with lower BMD in early adulthood. Therefore, the aim of our study was to assess the long-term effects of gestational age and particularly preterm birth on BMD in a large group of young adults.

Subjects and methods

Subjects

This study investigated a cohort of 276 healthy subjects, aged 18–24 years. Subjects born preterm (gestational age < 36 weeks, $n = 151$) had been admitted to the neonatal intensive care unit of the Erasmus University Medical Centre shortly after birth. In total, 37.7% of all preterm subjects were born small for gestational age. Term controls of similar age (gestational age $\geq 36$ weeks, $n = 125$) were randomly asked to participate from different educational institutes.

All subjects fulfilled the same inclusion criteria: i) age 18–24 years, ii) adult pubertal stage, iii) Caucasian, iv) born singleton, v) a neonatal period without signs of severe asphyxia (defined as an Apgar score below 3 after 5 min) or long-term complications of respiratory ventilation, such as bronchopulmonary dysplasia, vi) maximum duration of respiratory ventilation and/or oxygen supply of 2 weeks in the neonatal period. Subjects with a serious neonatal complication (e.g. necrotizing enterocolitis, degree 3 or more intraventricular haemorrhage, spastic hemilegia or quadriplegia), an endocrine or metabolic disorder, chromosomal defects, syndromes or dysmorphic symptoms suggestive for a yet unknown syndrome were excluded. Subjects with a condition known to interfere with growth, including GH deficiency, severe chronic illness, emotional deprivation, GH treatment, glucocorticosteroid treatment and radiotherapy were also excluded. Birth data were taken from hospital records, and records from community health services and general practitioners.

The Medical Research Ethics Committee of Erasmus University Medical Centre, Rotterdam, The Netherlands, approved this study. Written informed consent was obtained from all the participants.

Methods

Anthropometry Adult height was measured in the upright position to the nearest 0.1 cm using a Harpenden stadiometer (Holtain Ltd, Crymmyth, UK). Weight was measured to the nearest 0.1 kg on a digital scale (Servo Balance KA-20-150S). All anthropometric measurements were performed twice, and the mean value was used for analysis.

Bone mineral density Bone mass of the total body (TB), lumbar spine (LS), lean body mass (LBM) and fat mass (FM) were measured by dual-energy X-ray absorptiometry (DXA, type Lunar-Prodigy, GE Healthcare, Chalfont St Giles, UK). All scans were made on the same machine, and quality assurance was performed daily. The coefficient of variation (CV) was 0.5% for total body BMD and 1.0% for spine BMD (25–27). The CV for lean mass and FM has been reported to be 0.7 and 1.2% respectively (25, 26).

To adjust for differences in bone size, we calculated bone mineral apparent density of the LS (BMAD LS) (g/cm$^2$) with the model $\text{BMAD}_{LS} = \text{BMD}_{LS} \times (4/(\pi \times \text{width}))$ (28). Width was the mean width of the second to fourth lumbar vertebral body. This model was validated by in vivo volumetric data obtained from magnetic resonance imaging of the lumbar vertebrae.

Questionnaire All subjects completed a structured questionnaire, which included questions on socioeconomic status of the participants and their parents, cigarette smoking, alcohol consumption and usage of oral contraceptives. Socioeconomic status was determined by using educational level of the participant, which was assessed by the highest grade of school completed or currently participating in, and categorized into i) high (higher general secondary education or higher), ii) median (junior general secondary education – secondary vocational education) and iii) low education (preparatory middle-level vocational education or lower).

Statistical analysis

Clinical characteristics are expressed as median (interquartile range). Birth weight and length were adjusted for gestational age and gender (29), and baseline data in young adulthood were adjusted for age and gender (30). BMD data are expressed as mean (S.D.). SDSs for BMD TB, BMD LS and BMAD LS were calculated using reference data of Boot et al. (31).

Independent samples $t$-tests were used to evaluate differences in clinical characteristics between preterm and term subjects. To test for linear relationships between gestational age and BMD TB, BMD LS and BMAD LS, Pearson’s correlation coefficient was used. The long-term effect of gestational age on BMD TB, BMD LS and BMAD LS in young adulthood was analysed using multiple regression analyses, corrected for possible confounders (gender, age, birth weight SDS, birth length SDS, adult height SDS, LBM and FM). The interaction term birth length SDS×adult height SDS was added to the multiple regression model to ensure that the effect of these variables was modelled correctly.
To determine differences between preterm and term subjects with regard to BMD, multiple regression analysis was used with correction for age and gender and subsequently also with correction for gender, age, birth weight SDS, birth length SDS, adult height SDS, LBM and FM.

Results were regarded statistically significant at $P<0.05$. Statistics were performed using the computer Statistical Package for Social Science (SPSS version 16.0; SPSS, Inc., Chicago, IL, USA).

**Results**

**Clinical characteristics**

Table 1 shows the baseline characteristics of the study group. The total group of 276 subjects had a median age of 20.9 years. Besides the obvious difference in gestational age, preterm subjects had a lower birth length SDS than term subjects.

**BMD and gestational age**

There were no significant linear correlations between gestational age and BMD TB ($r=0.063$, $P=0.30$), BMD$_{LS}$ ($r=0.062$, $P=0.31$) and BMAD$_{LS}$ ($r=0.069$, $P=0.26$). Multiple regression analyses showed that gestational age was not a significant contributor to the variance in BMD$_{TB}$ ($P=0.27$, $R^2=0.370$), BMD$_{LS}$ ($P=0.91$, $R^2=0.165$) and BMAD$_{LS}$ ($P=0.87$, $R^2=0.262$), after adjustment for age, gender, birth weight SDS, birth length SDS, adult height SDS, LBM and FM as additional independent variables.

Furthermore, SDSs of BMD of the LS were not significantly correlated with gestational age (BMD$_{LS}$ SDS: $r=0.069$, $P=0.254$ and BMAD$_{LS}$ SDS: $r=0.031$, $P=0.605$), whereas BMD$_{TB}$ SDS was significantly correlated with gestational age ($r=0.137$, $P=0.023$). However, after adjustments for possible confounders, this correlation disappeared (BMD$_{TB}$ SDS ($P=0.28$, $R^2=0.290$), BMD$_{LS}$ SDS ($P=0.941$, $R^2=0.175$) and BMAD$_{LS}$ SDS ($P=0.815$, $R^2=0.064$)).

Further analysis showed that adult weight, specified as LBM and TB FM, was an important determinant of BMD in young adulthood, regardless of size at birth and gestational age. LBM was significantly associated with BMD$_{TB}$ ($\beta=0.007$, $P<0.001$) and BMD$_{LS}$ ($\beta=0.008$, $P<0.001$). FM was also associated with BMD$_{TB}$ ($\beta=0.002$, $P<0.001$) and BMD$_{LS}$ ($\beta=0.002$, $P=0.08$).

In addition, we evaluated the possible confounding effects of socioeconomic status, smoking, alcohol and the usage of oral contraceptives on these associations. The socioeconomic status was low in 8.8%, median in 22.6% and high in 68.8% of all subjects, and 22.8% of all subjects smoked. Of all subjects reporting alcohol consumption (80.8%), 28.6% had an alcohol consumption ≥5 units/week. None of these factors had a significant influence on the BMD (TB and LS) outcomes.

**BMD in preterm versus term subjects**

No significant differences were found between preterm and term subjects with regard to BMD$_{TB}$, BMD$_{LS}$ and BMAD$_{LS}$ after correction for age and gender (Table 2). Even after additional adjustment for significant independent variables like birth length SDS, adult height SDS, birth weight SDS, LBM and FM, BMD outcomes did not significantly differ between preterm and term subjects.

**Discussion**

Our study in 276 young adults showed that gestational age was not a significant contributor to the variance in BMD (BMD$_{TB}$ and BMD$_{LS}$) and BMAD (BMAD$_{LS}$) in young adulthood. In addition, our subgroup analysis did not show differences between subjects born preterm or term. Thus, in contrast to our hypothesis, preterm birth was not associated with lower BMD in early adulthood.

Peak bone mass is an important determinant of BMD in later life, but the exact age at which peak bone mass is reached is not well established. Recent data show that peak bone mass is probably attained by the mid-twenties (32). We are one of the first to investigate BMD in subjects born preterm at a mean age of 21 years, thus around the age of reaching peak bone mass. Our results are, therefore, important for the evaluation of the long-term consequences of preterm birth.

Our subgroup analysis showed no differences in BMD between subjects born preterm or term. Dalziel et al.

**Table 1** Clinical characteristics. All values are given as median (interquartile range).

<table>
<thead>
<tr>
<th></th>
<th>Total group</th>
<th>Preterm</th>
<th>Term</th>
</tr>
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<tbody>
<tr>
<td>$n$</td>
<td>276</td>
<td>151</td>
<td>125</td>
</tr>
<tr>
<td>Gender (M/F)</td>
<td>119/157</td>
<td>76/75</td>
<td>43/82</td>
</tr>
<tr>
<td>Gestational age (weeks)</td>
<td>34.9 (32.0 to 40.0)</td>
<td>32.2 (30.3 to 34.0)*</td>
<td>40.0 (39.0 to 40.0)</td>
</tr>
<tr>
<td>Birth length SDS</td>
<td>-0.6 (-1.7 to 0.3)</td>
<td>-1.1 (-2.8 to 0.3)*</td>
<td>-0.4 (-1.3 to 0.2)</td>
</tr>
<tr>
<td>Birth weight SDS</td>
<td>-0.3 (-1.6 to 0.6)</td>
<td>-0.3 (-2.1 to 0.8)</td>
<td>-0.3 (-1.2 to 0.4)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>20.9 (19.6 to 22.3)</td>
<td>21.0 (19.7 to 22.3)</td>
<td>20.6 (19.4 to 22.4)</td>
</tr>
<tr>
<td>Adult height SDS</td>
<td>-0.3 (-1.2 to 0.3)</td>
<td>-0.3 (-1.1 to 0.2)</td>
<td>-0.4 (-1.5 to 0.5)</td>
</tr>
<tr>
<td>Adult weight SDS</td>
<td>-0.3 (-1.0 to 0.3)</td>
<td>-0.2 (-0.9 to 0.4)</td>
<td>-0.4 (-1.0 to 0.3)</td>
</tr>
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</table>

* $P<0.001$ between preterm and term.
investigated a cohort of 174 adults (mean age 31 years) whose mothers had participated in a randomized trial of antenatal betamethasone treatment. Although the main aim of the study was to determine the effect of maternal betamethasone usage on later BMD, they also showed that prematurity had no effect on peak bone mass (21). Hamed et al. showed that, in a cohort of 230 women (aged 20–23 years), BMD was not influenced by prematurity (14), although in that study no adjustments for possible confounders (e.g. LBM, FM or weight) were made and the cohort consisted of female subjects only.

Weiler et al. (33) showed that preterm born adolescents had a lower bone mineral content (BMC) than those born at term. However, after correction for adult height and weight, these differences disappeared, indicating that the effect of gestational age on BMC was largely influenced by adult body size.

Previous studies in infancy and early childhood showed differences in BMD between preterm and term subjects (1, 6, 7, 9), but we could not confirm such results in our young adults. One explanation might be that there was a catch-up in bone mineralization during infancy and childhood (34), but to prove this hypothesis, prospective longitudinal research in preterm and term subjects is mandatory.

A relative high percentage of our premature study population consisted of subjects born small for gestational age (SGA; 37.7%), compared to the normal population in which the prevalence of SGA is only 2.3%. In clinical practice, the majority of children born SGA are born preterm. Furthermore, our study comprises a relatively healthy population. Subjects with, for example, a complicated neonatal period were excluded from our analysis, and it might be that such subjects have a higher risk of lower BMD.

BMD was assessed for TB and LS. As there are differences in cell biology between these two sites (35, 36), gestational age and/or prematurity could have different effects on BMD_{TB} and BMD_{LS}. Despite these differences, gestational age was no significant determinant of both BMD_{TB} and BMD_{LS}. Furthermore, it is important to realize that bone density is just one aspect of bone quality, as bones are complex three-dimensional structures (37).

Analysis of BMD showed that prematurity and also birth size had no long-term consequences on BMD. Instead, adult weight, LBM and TB FM were important determinants of BMD in young adulthood. So it seems important to ensure that appropriate growth and nutrition are maintained throughout childhood and adolescence. Although our data are reassuring, there is no guarantee that BMD will remain normal when subjects become older. Longer term follow-up is, therefore, still warranted.

DXA was used to assess BMD, as it is the most commonly used technique for BMD assessment because of low radiation exposure, great precision and accuracy, and short scanning time (6–10 min). Additionally, DXA performs whole body rather than slice measurements (CT) (31, 38). A shortcoming of DXA is that it measures bone in two dimensions providing only an estimation of bone density. BMD is obtained by dividing BMC (g) by the projected bone image (area in cm²). BMD is, therefore, dependent on bone size, and this might lead to erroneous interpretations of BMD values. A widely used and validated model to correct for bone size is BMAD_{LS} as one of the parameters reflecting bone mineralization.

Conclusions

In conclusion, in a cohort of 276 young adults, gestational age was not a significant determinant in the variance of BMD (TB and LS). BMD was not adversely affected in young adults born moderately preterm. Thus, in contrast to our hypothesis, preterm birth without postnatal complications is not associated with a lower BMD in young adulthood. These findings are of major importance as an increasing number of preterm born subjects reach adulthood.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.
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