Bone microarchitecture and volumetric bone density impairment in young male adults with childhood-onset growth hormone deficiency

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Abstract

Context: Adult growth hormone deficiency (AGHD) is characterized by low bone density and increased risk of fracture. Bone microarchitecture is insufficiently evaluated in patients with childhood-onset AGHD (CO AGHD).

Objective: To assess volumetric bone density (vBMD) and bone microarchitecture in CO AGHD in early adulthood after cessation of recombinant growth hormone (rhGH) treatment.

Design and subjects: Case–control study in a major academic medical center in Beijing, including 20 young male adults with CO AGHD and 30 age- and weight-matched non-athletic healthy men. High-resolution peripheral quantitative computerized tomography (HR-pQCT) of distal radius and tibia was performed.

Outcomes: The main outcomes were vBMD and morphometry parameters from HR-pQCT.

Results: Compared with healthy controls, CO AGHD group had significantly decreased insulin-like growth factor 1 (IGF-1) level and IGF-1 SDS (P < 0.001). β-CTX and alkaline phosphatase levels in CO AGHD group were significantly increased (P < 0.001). CO AGHD group had significantly decreased total vBMD, cortical vBMD, trabecular vBMD, cortical area, cortical thickness as well as trabecular thickness and trabecular bone volume fraction of both tibia and radius (P < 0.001). CO AGHD patients had an 8.4 kg decrease in grip strength and a significant decrease in creatinine levels (P = 0.001). At both tibia and radius, by finite element analysis, bone stiffness and failure load of the CO AGHD patients were significantly decreased (P < 0.001). After adjusting for age, BMI and serum levels of testosterone and free thyroxin, serum IGF-1 level was a positive predictor for total vBMD, cortical vBMD, cortical area, trabecular vBMD, bone stiffness and failure load of both tibia and distal radius in all subjects.

Conclusions: Young adult male patients with childhood-onset adult growth hormone deficiency who are no longer receiving growth hormone replacement have prominently impaired volumetric bone density and bone microarchitecture and lower estimated bone strength.

Introduction

Adult growth hormone deficiency (AGHD) is an uncommon but debilitating disorder characterized by low bone mineral density, sarcopenia, increased risk of metabolic syndrome and decreased quality of life (1). Growth hormone promotes linear growth and healthy development in childhood and plays crucial
roles in normal body function in adulthood, including maintenance of bone strength, muscle volume and other metabolic parameters. Since recombinant human growth hormone (rhGH) became available in 1985, it was reported to be effective in improving metabolic parameters, bone mineral density and quality of life in AGHD patients (2).

Dual-energy X-ray absorptiometry (DXA) at the hip and lumbar spine is widely used to evaluate skeletal situation in AGHD patients before and after rhGH replacement therapy (3, 4, 5). However, DXA only evaluates 2D areal bone mineral density (aBMD) and does not provide details of microarchitecture of cortical and cancellous bones. Besides, there might be false increases of aBMD due to bone hyperplasia, spinal degeneration and aortic calcification in lumbar spine DXA. Thus, DXA is lacking sensitivity and specificity in predicting fracture risks (6). High-resolution peripheral quantitative CT (HR-pQCT) allows noninvasive assessment of bone microstructure and volumetric bone mineral density (vBMD) at peripheral sites at high resolution (82 μm isotropic voxel size) and with relatively low radiation exposures (3–5 μSv) (7). Recently, bone microarchitecture was assessed by HR-pQCT in 16 male patients with unreplaced adulthood-onset AGHD (AO AGHD) (8). In that report, the mean age of AO AGHD patients was 47.4 years and the age of diagnosis was 37.1 years. No bone microarchitecture deficiency was reported in this series of patients.

As we know, peak bone mass is the maximum amount of bone during individual’s life. It typically occurs in the early 20s in females and late 20s in males. Growth hormone and gonadosteroids play synergistic roles in peak bone mass. Most adolescents with multiple pituitary hormone deficiency (MPHD) will stop rhGH treatment after completion of linear growth but will sustain replacement therapy of other hormones (9). At present, no data are available for vBMD and bone microarchitecture in AGHD patients due to childhood-onset MPHD after cessation of rhGH treatment in early adulthood.

In this cross-sectional study, we assessed the vBMD, bone microarchitecture and estimated bone strength of young male adults with CO AGHD after cessation of rhGH replacement therapy. Female subjects were not enrolled to avoid potential gender-dimorphic variations in bone microarchitecture. The relationship between insulin-like growth factor 1 (IGF-1) levels and changes of vBMD, microarchitecture and estimated bone strength was also evaluated.

**Subjects and methods**

**Subjects**

A total of 20 consecutive male patients with CO AGHD were enrolled in this cross-sectional study from April 2017 to May 2018 in our GHD clinic at Peking Union Medical College Hospital (10). The diagnosis of AGHD was according to the criteria of American Endocrine Society Clinical Practice Guideline (11). Six patients underwent insulin tolerance test and peak value of GH was less than 3 ng/mL in all these patients. The diagnosis was confirmed in the other 14 patients since IGF-1 levels were below the age-adjusted normal range and there were deficiencies in three or more pituitary axes at the same time. All patients had stopped rhGH replacement therapy since attainment of final height. A total of 30 non-athletic healthy male were recruited from general population as controls. Approval from the Institutional Review Board of Peking Union Medical College Hospital was obtained for this study. Written informed consent was obtained from all participants. All data were de-identified before analysis.

**Anthropometrics**

Height, weight, waist circumference and hip circumference of the subjects were measured by standard protocols in the early morning with light clothes. BMI was calculated as weight (kg) divided by height (m) squared.

**Bioimpedance assessments and body composition**

Bioelectric impedance assessments (BIAs) by body composition analyzer (Tanita TBF-410, Japan) were performed on each of the participants according to manufacturer’s instructions. Measurements were taken in the morning after overnight fasting. Fat-free mass (FFM), fat-free mass percentage (FFM%), fat mass (FM), FM percentage (FM%) and total body water (TBW) were obtained according to the body composition model. FFM derived from BIA has been validated previously against DXA (12). The FFM index and FM index were calculated with formula by Van Itallie et al., namely FFMI=FFM/height² (kg/m²) and FMI=fat mas/height² (kg/m²) (13).

**Muscle strength**

Grip strength was measured using a hand dynamometer (Jamar Plus+, Sammons Preston, Rolyon, Bolingbrook, IL, USA) on the non-dominant hand. All participants...
performed three trials of a maximum effort squeeze contracting for 5 s and the maximum values were recorded.

**High-resolution peripheral computed tomography (HR-pQCT) and finite element analysis**

High-resolution images of the non-dominant distal radius and distal tibia were obtained by HR-pQCT scan (XtremeCT II scanner, ScancoMedical, Brüttisellen, Switzerland). vBMD and bone microarchitecture at the distal tibia and radius were investigated using a previously described protocol (14). All participants were scanned with the standard human in vivo scanning protocol (60 kVp, 1000 μA, 100 ms integration time). In short, when a scout scan was finished, reference lines were placed at the distal end plates for both the radius and tibia. Each scan comprised 110 slices, corresponding to a 10.2 mm scan area, with a nominal isotropic resolution of 82 μm. Each scan carried out at the standard location 9.0 mm (radius) and 22.0 mm (tibia) proximal to the reference line. All scans were finished by a specified technologist and analyzed according to the standard manufacturer’s method (15). Scans were graded for motion artifacts as described previously (16).

Bone microarchitecture parameters, including total area (Tt.Ar), trabecular area (Tb.Ar), cortical area (Ct.Ar), cortical perimeter (Ct.Pm), total volume bone mineral density (Tr.vBMD), trabecular vBMD (Tb.vBMD), cortical vBMD (Ct.vBMD), trabecular separation (Tb.Sp), inhomogeneity of network (Tb.1/N.SD), cortical thickness (Ct.Th) and cortical porosity (Ct.Po), were obtained by the standard morphologic analysis using semi-automated software (15). The finite element analysis protocol has been previously described (8, 17). Using dedicated software, HR-pQCT images of the radius and tibia were converted into finite element models to estimate bone stiffness (N/mm). Failure load (N) was calculated using the Pistoia criterion (18).

**Biochemical measurements**

The blood samples were obtained on the day of bone assessment. Liver function, kidney function, HbA1c, serum calcium, serum phosphate, alkaline phosphatase (ALP), 25-hydroxyvitamin D (25 OHD), parathyroid hormone (PTH), β-C-terminal telopeptide region of collagen type I (β-CTX) and hormonal evaluation were all tested in the department of clinical laboratory of Peking Union Medical College Hospital by standard methods. The serum IGF-1 level was measured with a fully automated two-site, solid-phase, chemiluminescent enzyme immunometric assay (Immulite 2000, Siemens Healthcare Diagnostics).

**Statistical analysis**

Data are presented as the mean ± standard deviation (s.d.). The independent-samples t test was used for data analysis between the two groups. Skewed data were in-transformed before t test. The Mann–Whitney U test was used if the data were not normally distributed. Multivariate, forward stepwise linear regression analyses were used to identify predictors of bone microarchitecture and estimated bone strength. All the statistical computations were run using SPSS software, version 22.0 for Windows (SPSS Inc.), and P < 0.05 was considered to be statistically significant.

**Results**

**Demographic, clinical and biochemical characteristics**

General characteristics of CO AGHD patients and healthy male controls are shown in Table 1. A total of 20 male patients with CO AGHD were enrolled in this cross-sectional study. Fifteen patients had pituitary stalk interruption and four patients had pituitary hypoplasia based on MRI. Eleven of them (11/19) were born with breech or foot presentation. Another one patient underwent surgery for craniopharyngioma at the age of 4 years. All patients had MPHD. Fifteen patients sustained levothyroxine replacement therapy. Eight patients sustained glucocorticoid replacement. Testosterone replacement was started at 18 years old and after completion of linear growth in all patients. All patients accepted rhGH replacement therapy since childhood. Duration of rhGH replacement treatment was 11.2 ± 3.5 years. The average time course since the cessation of rhGH replacement was 6.6 ± 3.3 years. No fracture was reported from all our patients.

There was no difference in chronologic age between both groups (28.2 ± 5.4 vs 30.6 ± 4.9 years, P = 0.104). Compared with the control group, CO AGHD group had a 7.4 cm lag in final height (167.1 ± 6.8 vs 174.5 ± 5.3 cm, P < 0.001). CO AGHD group had increased BMI (25.6 ± 3.7 vs 23.2 ± 2.8, P = 0.014), an 11.0 cm increase in waist circumference (95.1 ± 10.3 vs 84.1 ± 6.7 cm, P < 0.001) and increased waist-hip ratio (0.96 ± 0.08 vs 0.87 ± 0.05,
AGHD group also had a 4.2% increase in fat mass percentage (27.0±5.9 vs 22.8±4.9%, P=0.009) and increased FMI (7.0±2.5 vs 5.4±1.7, P=0.010). There was no difference in waist circumference (23.2±0.104 vs 23.7±0.96, P=0.086) between the two groups. Besides, AGHD patients had an 8.4kg decrease in grip strength (37.3±4.0 vs 45.7±8.2kg, P<0.001) and a significantly decreased creatinine level (68.1±9.1 vs 72.3±6.7, P<0.001). Before glucocorticoids replacement, CO AGHD group also had a 4.2% increase in fat mass percentage (27.0±5.9 vs 22.8±4.9%, P=0.009) and increased FMI (7.0±2.5 vs 5.4±1.7, P=0.010). No difference was found in FM, FFM or TBW between two groups. Besides, AGHD patients had an 8.4kg decrease in grip strength (37.3±4.0 vs 45.7±8.2kg, P<0.001) and a significantly decreased creatinine level (68.1±9.1 vs 72.3±6.7, P<0.001).

**Bone microarchitecture of the distal tibia**

Data obtained by HR-pQCT of the distal tibia was shown in Fig.1 and listed in Table 3. Compared with healthy controls, CO AGHD group had significantly decreased total vBMD (209.61±45.73 vs 326.98±66.87mg HA/cm^3, P<0.001), as well as cortical vBMD (850.53±53.21 vs 929.51±37.08mg HA/cm^3, P<0.001) and trabecular vBMD (136.38±32.76 vs 188.86±42.20mg HA/cm^3, P<0.001) in distal tibia.

CO AGHD group also had significant decreased cortical area (95.92±20.22 vs 150.09±28.17mm^2, P<0.001), cortical thickness (0.910±0.204 vs 1.591±0.365mm, P<0.001), intra-cortical porosity (0.010±0.004 vs 0.023±0.013, P<0.001), trabecular thickness (0.236±0.021 vs 0.267±0.026mm, P<0.001) and trabecular bone volume fraction (0.213±0.046 vs 0.282±0.054, P<0.001) in distal tibia.

At the same time, CO AGHD group had significant increased total area (933.37±152.78 vs 801.49±116.45mm^2, P<0.001), cortical perimeter (118.65±10.39 vs 110.24±8.09mm, P=0.002) and trabecular area (843.65±152.51 vs 657.14±124.23mm^2, P<0.001) in distal tibia.

**Bone microarchitecture of the radius**

Data obtained by HR-pQCT of the radius was shown in Fig.1 and listed in Table 4. Compared with healthy controls, AGHD group had significantly decreased total vBMD (226.38±56.90 vs 355.35±66.49mg HA/cm^3, P<0.001), as well as cortical vBMD (747.16±80.22 vs 919.51±38.68mg HA/cm^3, P<0.001) and trabecular vBMD (140.92±37.60 vs 184.65±39.82mg HA/cm^3, P<0.001) in radius.
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with the control group. Similarly, at the radius (Figure 1, Table 2), bone microarchitecture of CO AGHD patient (A) and healthy control (B). Bone microarchitecture of CO AGHD (A′ and C′) compared with healthy control (B′ and D′). Right panel shows deficient trabecular bone in both radius and distal tibia in a patient with CO AGHD (A′ and C′) compared with healthy control (B′ and D′). Bone stiffness (47.397.1 ± 1514.4 vs 89036.9 ± 17207.6, P < 0.001) and failure load (2639.9 ± 889.4 vs 4839.2 ± 905.3, P < 0.001) of the AGHD group were also significantly decreased in comparison with the healthy controls.

Finite element analysis

At the tibia (Table 3), bone stiffness (165377.8 ± 41389.1 vs 239248.1 ± 43866.4, P < 0.001) and failure load (9151.1 ± 2197.9 vs 12889.1 ± 2191.3, P < 0.001) of the AGHD group were significantly decreased when compared with the control group. Similarly, at the radius (Table 4), bone stiffness (47.397.1 ± 1514.4 vs 89036.9 ± 17207.6, P < 0.001) and failure load (2639.9 ± 889.4 vs 4839.2 ± 905.3, P < 0.001) of the AGHD group were also significantly decreased in comparison with the healthy controls.

Association between IGF-1 levels and parameters of bone microarchitecture and estimated bone strength

On multivariate analysis, after adjusting for age, BMI and serum levels of testosterone and FT4, serum IGF-1 level was a positive predictor for total vBMD (β = 0.4710,
Table 3  HR-pQCT parameters of the distal tibia in CO AGHD group and controls. Data are presented as mean ± s.d.

| Parameter                      | Control (n = 30)       | CO AGHD (n = 20)      | P value  |
|-------------------------------|------------------------|-----------------------|----------|
| Total area (mm²)              | 801.49 ± 116.45        | 933.37 ± 152.78       | 0.001    |
| Total vBMD (mg HA/cm³)        | 326.98 ± 66.87         | 209.61 ± 45.73        | <0.001   |
| Cortical area (mm²)           | 150.09 ± 28.17         | 95.92 ± 20.22         | <0.001   |
| Cortical vBMD (mg HA/cm³)     | 929.51 ± 37.08         | 850.53 ± 53.21        | <0.001   |
| Cortical perimeter (mm)       | 110.24 ± 8.09          | 118.65 ± 10.39        | 0.002    |
| Cortical thickness (mm)       | 1.591 ± 0.365          | 0.910 ± 0.204         | <0.001   |
| Intra-cortical porosity       | 0.023 ± 0.013          | 0.010 ± 0.004         | <0.001   |
| Trabecular area (mm²)         | 657.14 ± 124.23        | 843.65 ± 152.51       | <0.001   |
| Trabecular vBMD (mg HA/cm³)   | 188.86 ± 42.20         | 136.38 ± 32.76        | <0.001   |
| Trabecular thickness (mm)     | 0.267 ± 0.026          | 0.236 ± 0.021         | <0.001   |
| Trabecular number (1/mm)      | 1.378 ± 0.177          | 1.331 ± 0.253         | 0.449    |
| Trabecular separation (mm)    | 0.710 ± 0.095          | 0.754 ± 0.186         | 0.501    |
| Tb.1/N SD (mm)                | 0.297 ± 0.043          | 0.300 ± 0.096         | 0.235    |
| Trabecular bone volume fraction| 0.282 ± 0.054          | 0.213 ± 0.046         | <0.001   |
| Bone stiffness (N/mm)         | 239.248 ± 43.866.4     | 165.377.8 ± 41.389.1  | <0.001   |
| Bone failure load (N)         | 12 889.1 ± 2191.3      | 9151.1 ± 2197.9       | <0.001   |

Tb.1/N SD, St. Dev of 1/Tb.N, inhomogeneity of network; vBMD, volumetric bone mineral density.

Table 4  HR-pQCT parameters of the distal radius in CO AGHD group and controls. Data are presented as mean ± s.d.

| Parameter                      | Control (n = 30)       | CO AGHD (n = 20)      | P value  |
|-------------------------------|------------------------|-----------------------|----------|
| Total area (mm²)              | 324.96 ± 54.26         | 343.54 ± 71.24        | 0.301    |
| Total vBMD (mg HA/cm³)        | 355.35 ± 66.49         | 226.38 ± 56.90        | <0.001   |
| Cortical area (mm²)           | 75.54 ± 10.74          | 47.60 ± 9.82          | <0.001   |
| Cortical vBMD (mg HA/cm³)     | 919.51 ± 38.68         | 747.16 ± 80.22        | <0.001   |
| Cortical perimeter (mm)       | 75.65 ± 6.56           | 77.77 ± 9.87          | 0.406    |
| Cortical thickness (mm)       | 1.186 ± 0.193          | 0.706 ± 0.195         | <0.001   |
| Intra-cortical porosity       | 0.006 ± 0.004          | 0.004 ± 0.002         | 0.126    |
| Trabecular area (mm²)         | 253.48 ± 55.27         | 300.12 ± 72.64        | 0.013    |
| Trabecular vBMD (mg HA/cm³)   | 184.65 ± 39.82         | 140.92 ± 37.60        | <0.001   |
| Trabecular thickness (mm)     | 0.246 ± 0.018          | 0.215 ± 0.018         | <0.001   |
| Trabecular number (1/mm)      | 1.462 ± 0.180          | 1.525 ± 0.236         | 0.294    |
| Trabecular separation (mm)    | 0.633 ± 0.083          | 0.641 ± 0.116         | 0.756    |
| Tb.1/N SD (mm)                | 0.250 ± 0.036          | 0.246 ± 0.056         | 0.810    |
| Trabecular bone volume fraction| 0.273 ± 0.056          | 0.203 ± 0.054         | <0.001   |
| Bone stiffness (N/mm)         | 89 036.9 ± 17 207.6    | 47 397.1 ± 5 174.1    | <0.001   |
| Bone failure load (N)         | 4839.2 ± 905.3         | 2639.9 ± 889.4        | <0.001   |

Tb.1/N SD, St. Dev of 1/Tb.N, inhomogeneity of network; vBMD, volumetric bone mineral density.

Discussion

In this study, we focused on vBMD, bone microarchitecture and estimated bone strength of young male adult with CO AGHD and compared with age-matched controls. Our results showed that (1) vBMD of both distal tibia and trabecular thickness (β=0.0001, P<0.001), trabecular bone volume fraction (β=0.0003, P=0.003), bone stiffness (β=161.25, P<0.001) and failure load (β=8.49, P<0.001). Serum IGF-1 level was also a negative predictor for trabecular area (β=−0.1797, P=0.046) of the radius (Table 6).

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and non-dominant radius were significantly decreased in AGHD patients; (2) CO AGHD patients had significantly decreased cortical area and cortical thickness, as well as trabecular thickness and trabecular bone volume fraction of both tibia and distal radius; (3) CO AGHD patients had lower estimated bone strength; (4) after adjusting for age, BMI and serum levels of testosterone and free T4, serum IGF-1 level was a positive predictor for total vBMD, cortical vBMD, cortical area, trabecular vBMD, bone stiffness and failure load. In our series of CO AGHD patients, the average time course since cessation of rhGH replacement was 6.6±3.3 years and all patients sustained testosterone replacement since 18 years old. We thus conclude that young adult male patients with CO AGHD who are no longer receiving GH replacement have abnormalities in bone microarchitecture and estimated bone strength.

Low bone mass is widely believed as a characteristic feature of AGHD since the GH/IGF axis plays a pivotal role in skeletal growth and strength. GH activates both bone formation and resorption via direct and indirect mechanisms (19). IGF-1 regulates radial bone growth and cortical and trabecular bone properties via their effects on osteoblast, osteocyte and osteoclast function (20). Both bone size and BMD increase gradually throughout childhood and rapidly increase during puberty. Calcium is added to bone most rapidly since about age 9 in girls and age 10 in boys (21). During puberty, interactions of IGF-1 and gonadostereoids and parathyroid hormone contribute to the appropriate accumulation of bone mass. Thus, BMD continues to increase until age reaches the mid-to-late 20s and decline gradually throughout the rest of adult life.

However, there were lots of inconsistencies of data of bone density and bone microarchitecture features in AGHD patients. Data from Stephen et al. described normal trabecular bone mineral density and only a 2% decrease in cortical density in radius was described in 13 CO AGHD patients (22). It was suggested that the apparent low BMD observed with DXA was a reduction in cortical bone volume and not density. In our series of patients, rhGH was stopped at the end of puberty and testosterone replacement was sustained in early adulthood. vBMD of both distal tibia and non-dominant radius were significantly decreased. At the same time, the increased β-CTX and ALP suggested active turnover of bone in AGHD patients. While serum levels of both 25 OHD and PTH were all in normal ranges, which suggested that the low vBMD and impaired bone microarchitecture were mainly caused by low IGF-1 levels. As we know, peak bone mass is reached after final adult height, at a mean of 23.1 years in males and 19.9 years in females (23). There was a concurrence of formative years of peak bone development and cessation of rhGH replacement. After adjusting for age, BMI and serum testosterone levels, serum IGF-1 level was a positive predictor for total vBMD, cortical vBMD, cortical area and trabecular vBMD of both tibia and distal radius. Our work thus shed new insight into the seamless transition of childhood-onset GHD from pediatric to teenage and young adulthood.

Our results also provide new insight into the clinical characteristics of AGHD with different age of onset. AGHD is a group of condition with heterogeneous etiologies. There were striking regional variations in pathogenesis and clinical characteristics among adult-onset AGHD (AO AGHD) in Europe and the United States (24). In China, AGHD is less recognized and treated comparing with Western countries and no consensus of management has been reached so far (25). A proportion of patients with childhood-onset GHD are idiopathic.

### Table 5 Associations between bone microarchitecture of the distal tibia and IGF-1 levels after adjusting for age, BMI and serum levels of testosterone and FT4.

|                        | β     | P value |
|------------------------|-------|---------|
| Total vBMD (mg HA/cm²) | 0.4710 | <0.001  |
| Cortical area (mm²)    | 0.2347 | <0.001  |
| Cortical vBMD (mg HA/cm²) | 0.3087 | <0.001  |
| Cortical thickness (mm)| 0.0029 | <0.001  |
| Intra-cortical porosity| 0.00005| 0.004   |
| Trabecular vBMD (mg HA/cm³)| 0.2030| 0.001   |
| Trabecular thickness (mm)| 0.0001| 0.001   |
| Trabecular bone volume fraction| 0.0003| 0.001   |
| Bone stiffness (N/mm)  | 315.14 | <0.001  |
| Bone failure load (N)  | 15.92  | <0.001  |

vBMD, volumetric bone mineral density.

### Table 6 Associations between bone microarchitecture of the distal radius and IGF-1 levels after adjusting for age, BMI and serum levels of testosterone and FT4.

|                        | β     | P value |
|------------------------|-------|---------|
| Total vBMD (mg HA/cm³) | 0.4781 | <0.001  |
| Cortical area (mm²)    | 0.1096 | <0.001  |
| Cortical vBMD (mg HA/cm³) | 0.6503| <0.001  |
| Cortical thickness (mm)| 0.0019 | <0.001  |
| Trabecular area (mm²)  | −0.1797| 0.046   |
| Trabecular vBMD (mg HA/cm³)| 0.1565| 0.008   |
| Trabecular thickness (mm)| 0.0001| <0.001  |
| Trabecular bone volume fraction| 0.0003| 0.003   |
| Bone stiffness (N/mm)  | 161.25 | <0.001  |
| Bone failure load (N)  | 8.49   | <0.001  |

vBMD, volumetric bone mineral density.
and the growth hormone insufficiency is transient (26). While in patients with MPHDD, GHD is usually permanent (27). A large number of patients with MPHD drop out of rhGH replacement therapy during the transition period. There are a lot of associated factors, including insufficient knowledge of treatment necessity in patients, their family and health providers (8), inconvenience of rhGH administration, as well as social-economic issues. In recent data from 16 AO AGHD patients due to sellar masses or traumatic brain injury (8), bone microarchitecture was not deficient in these patients. Our data thus supported that CO AGHD due to MPHD is a distinct entity in AGHD and should be treated in different way from AO AGHD.

The main limitation of our study is the small number of participants. But there was good homogeneity in the etiology, age of onset, duration of rhGH replacement therapy and time course since cessation of rhGH treatment. Inconsistency of etiology and disease duration is one of the main reasons for the inconsistently reported clinical characteristics. Another limitation is the cross-sectional nature of our study. We are going to follow-up these patients in long term and evaluate the dynamic changes of bone microarchitecture and risk of fracture in the future. At the same time, further studies to evaluate effects of rhGH replacement therapy on vBMD and microstructure of CO AGHD patients due to MPHD are needed to elucidate the cause–effect relationship between IGF-1 levels and bone quality. The third limitation is that no questionnaire about exercise was carried out in this study, since appropriate exercise is capable of increasing bone mass and strength.

In conclusion, our findings indicate that CO AGHD patients who are no longer receiving growth hormone replacement had significantly impaired vBMD and microarchitecture and lower estimated bone strength in radius and distal tibia. IGF-1 level is positive predictor for total vBMD, cortical vBMD, cortical area, trabecular vBMD, bone stiffness and failure load. Seamless transition of management is advocated in childhood-onset AGHD patients due to MPHD.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of this study.

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Author contribution statement
H Y designed the study and wrote the primary manuscript. H Z and H P designed and supervised the study and revised the primary manuscript. K Y took part in the collection of clinical data and analyzed the data. Q Z performed the biochemical measurements. L W contributed to the study management. F G helped to revise the primary manuscript.

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References
1 Molitch ME, Clemmons DR, Malozowski S, Merriam GR, Vance ML & Endocrine Society. Evaluation and treatment of adult growth hormone deficiency: an Endocrine Society clinical practice guideline. Journal of Clinical Endocrinology and Metabolism 2011 96 1587–1609. (https://doi.org/10.1210/jc.2011-0179)
2 Hoffman AR, Kuntze JE, Baptista J, Baum HB, Raumann GP, Biller BM, Clark RV, Cook D, Inzucchi SE, Kleinberg D et al. Growth hormone (GH) replacement therapy in adult-onset gh deficiency: effects on body composition in men and women in a double-blind, randomized, placebo-controlled trial. Journal of Clinical Endocrinology and Metabolism 2004 89 2048–2056. (https://doi.org/10.1210/jc.2003-030346)
3 Barake M, Klibanski A & Tittos NA. Effects of recombinant human growth hormone therapy on bone mineral density in adults with growth hormone deficiency: a meta-analysis. Journal of Clinical Endocrinology and Metabolism 2014 99 852–860. (https://doi.org/10.1210/jc.2013-3921)
4 Gotheerstrom G, Bengtsson BA, Bosaeus I, Johannsson G & Svensson J. Ten-year GH replacement increases bone mineral density in hypopituitary patients with adult onset GH deficiency. European Journal of Endocrinology 2007 156 55–64. (https://doi.org/10.1530/eje.1.02317)
5 Hitz MF, Jensen JE & Eskildsen PC. Bone mineral density in patients with growth hormone deficiency: does a gender difference exist? Clinical Endocrinology 2006 65 783–791. (https://doi.org/10.1111/j.1365-2265.2006.02667.x)
6 Nicks KM, Amin S, Atkinson LJ, Riggs BL, Melton LJ 3rd & Khosla S. Relationship of age to bone microarchitecture independent of areal bone mineral density. Journal of Bone and Mineral Research 2012 27 637–644. (https://doi.org/10.1002/jbmr.1468)
7 Geusens P, Chapurlat R, Schett G, Ghasem-Zadeh A, Seeman E, de Jong J & van den Bergh J. High-resolution in vivo imaging of bone microarchitecture. Nature Reviews Rheumatology 2014 10 304–313. (https://doi.org/10.1038/nrrheum.2014.23)
8 Silva PPB, Amlashi FG, Yu EW, Pulaski-Liebert KJ, Gerweck AV, Fazeli PK, Lawson E, Nachtrigal LB, Biller BMK, Miller KK et al. Bone microarchitecture and estimated bone strength in men with active acromegaly. European Journal of Endocrinology 2017 177 409–420. (https://doi.org/10.1530/EJE-17-0468)
9 Yang H, Zhu H, Yan K & Pan H. Childhood-Onset Adult growth hormone deficiency: clinical, hormonal, and radiological assessment in a single center in China. Hormone Research in Paediatrics 2017 88 155–159. (https://doi.org/10.1159/000478527)
10 Wilson JD. Peking Union Medical College Hospital, a palace of endocrine treasures. Journal of Clinical Endocrinology and Metabolism 1993 76 815–816. (https://doi.org/10.1210/jcem.76.4.8473387)
11 Molitch ME, Clemmons DR, Malozowski S, Merriam GR, Vance ML & Endocrine Society. Evaluation and treatment of adult growth hormone deficiency: an Endocrine Society clinical practice guideline.
Clinical Study

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19 Giustina A, Mazzotti G & Canalis E. Growth hormone, insulin-like growth factors, and the skeleton. Endocrine Reviews 2008 29 535–559. (https://doi.org/10.1210/er.2007-0036)

20 Yakar S, Werner H & Rosen CJ. Insulin-like growth factors: actions on the skeleton. Journal of Molecular Endocrinology 2018 61 T115–T137. (https://doi.org/10.1530/JME-17-0298)

21 Abrams SA. Calcium turnover and nutrition through the life cycle. Proceedings of the Nutrition Society 2001 60 283–289.

22 Murray RD, Adams JE & Shalet SM. A densitometric and morphometric analysis of the skeleton in adults with varying degrees of growth hormone deficiency. Journal of Clinical Endocrinology and Metabolism 2006 91 432–438. (https://doi.org/10.1210/jc.2005-0897)

23 Boot AM, de Riddere MA, van der Sluis IM, van Slobbe I, Krenning EP & Keizer-Schrama SM. Peak bone mineral density, lean body mass and fractures. Bone 2010 46 336–341. (https://doi.org/10.1016/j.bone.2009.10.003)

24 Brabant G, Poll EM, Jonsson P, Polydorou D & Kreitschmann-Andermahr I. Etiology, baseline characteristics, and biochemical diagnosis of GH deficiency in the adult: are there regional variations? European Journal of Endocrinology 2009 161 (Supplement 1) S25–S31. (https://doi.org/10.1530/EJE-09-0273)

25 Yang H, Zhang M, Pan H & Zhu H. Management of adult growth hormone deficiency at Peking Union Medical College Hospital: a survey among physicians. Chinese Medical Sciences Journal 2016 31 168–172. (https://doi.org/10.1016/j.cmsj.2016.05.001)

26 Tauber M, Moulin P, Plenkowski C, Jouret B & Rochiccioli P. Growth hormone (GH) retesting and auxological data in 131 GH-deficient patients after completion of treatment. Journal of Clinical Endocrinology and Metabolism 1997 82 352–356. (https://doi.org/10.1210/jcem.82.2.3726)

27 Berberoglu M, Siklar Z, Darendeliler F, Poyrazoglu S, Darcan S, Isguven P, Birdici A, Ocal G, Bundak R, Yuksel B et al. Evaluation of permanent growth hormone deficiency (GHD) in young adults with childhood onset GHD: a multicenter study. Journal of Clinical Research in Pediatric Endocrinology 2008 1 30–37. (https://doi.org/10.4008/jcrpe.v1i1.7)

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