OBJECTIVE: Developmental dysplasia of the hip (DDH) is recognized as a frequent cause of secondary osteoarthritis (OA). The purpose in this study was to compare structural and biomechanical properties of subchondral trabecular bone and its relationship with cartilage damage between patients with DDH and patients with primary hip OA.

METHODS: Forty-three femoral head specimens obtained from patients who underwent total hip arthroplasty [DDH, n = 17; primary OA, n = 16; and normal control (NC), n = 10] were scanned by microcomputed tomography and analyzed by individual trabecula segmentation to obtain the microstructural types of subchondral trabecular bone. The biomechanical properties were analyzed by micro-finite element analysis, and cartilage damage was evaluated by histology. The linear regression analysis was used to indicate the association between microstructures, biomechanical property, and articular cartilage.

RESULTS: The DDH group showed the lowest total bone volume fractions (BV/TV) and plate BV/TV in the three groups (p < 0.05). There were also different discrepancies between the three groups in plate/rod trabecular number, plate/rod trabecular thickness, trabecular plate surface area/rod length, and junction density with different modes (plate–plate, rod–rod, and plate–rod junction density). The micro-finite element analysis, histology, and linear regression revealed that the subchondral trabecular bone in the DDH group had inferior biomechanical properties and cartilage damage of patients with DDH was more serious with different subchondral trabecular bone microstructures.

CONCLUSION: Our findings suggested that changes of subchondral trabecular bone play a critical role in DDH progression and that the improvement on subchondral trabecular bone may be a sensitive and promising way in treatment of DDH.
younger than 50 years have clinical indication for total hip arthroplasty (THA) [3]. Given its large population, China may bear the highest DDH-related burden compared with other countries. DDH has become a public health challenge, which underscores the need for urgent comprehensive therapy.

The radiographic measurements remain the primary basis for the diagnosis of DDH [4]. Patients with early-stage DDH definitively develop secondary OA, with 50% of these patients developing advanced OA at the age of 50 years [5]. Compared with primary hip OA, DDH often causes more severe symptoms and imaging manifestations (i.e., early hip instability and limited mobility, abnormal joint loading and stress distribution, absorption of the femoral head, and hip dislocation causing formation of a false joint) [6,7]. Some studies have illustrated changes in the subchondral trabecular bone microstructure in primary OA [8,9]. Although the characteristics of DDH are quite similar to those of primary OA, discrepancies in the subchondral trabecular bone microstructure between these conditions are largely unknown. Knowledge of these discrepancies may help optimise the surgery plan of DDH, and subchondral trabecular bone may be a target for the treatment of early-stage DDH.

Recently, imaging technologies, i.e., microcomputed tomography (micro-CT), have revealed key evidence indicating the imperative role of the subchondral microstructure in the pathogenesis of primary OA [8,10]. Meanwhile, micro-finite element analysis (μFEA) was used to quantitatively analyze the biomechanical properties and their relationship with the subchondral trabecular bone microstructure in patients with OA [11]. However, there are several limitations to the current standard morphological analysis. The individual trabecula segmentation (ITS) analysis maintaining essentially all the plate and rod microarchitecture, number, shape, volume of trabecular plates and rods, and connectivity between trabecular plates–plates, plates–rods, and rods–rods, can evaluate the detailed trabecular bone network [12]. Specifically, the morphological parameters might be insensitive to subtle variations in statistical means of the imaged bone trabecular network, compared with single measurement values of the individual trabecula [12]. In this study, our aim was to indicate the structural and mechanical property of subchondral trabecular bone and articular cartilage in patients with DDH. We explored the relationship among the subchondral trabecular bone microstructure, biomechanical properties, and cartilage damage. We hypothesized that the changes of the subchondral trabecular bone microstructure might be correlated with biomechanical properties and cartilage damage in patients with DDH.

### Materials and methods

#### Specimen preparation

In this study, femoral head specimens were obtained from 17 patients with DDH (DDH group, 45.8–55.2 years) and 16 patients with primary hip OA (OA group, 50.1–68.4 years). All patients who underwent THA were diagnosed as having DDH or primary hip OA based on radiographic and clinical diagnostic criteria by an experienced surgeon in orthopaedic surgery at Shanghai Ninth People’s Hospital [5,13]. Furthermore, 10 healthy femoral heads from patients with the fractured femoral neck who underwent THA in the same hospital constituted the normal control group (NC group, 48.4–61.3 years). This study was approved by the Institutional Review Board of Shanghai Ninth People’s Hospital (IRB reference number: 2018-179-T137). All patients provided informed written consent before their participation in this study. All patients had a detailed history and physical examination, as well as biochemical marker detection tests, to exclude those with systemic diseases, such as haematological disorders, endocrine diseases, thyrotoxicosis, metabolic diseases of the liver and kidney, or malignancies that may have affected systemic bone metabolism. The femoral head specimens were refrigerated for follow-up studies.

#### Micro-CT scanning and ITS-based morphological analysis

All bone specimens were scanned using the high-resolution micro-CT system (Micro-CT 80; Scanco Medical AG, Switzerland) with 36-μm isotropic voxel size [14]. There are particular patterns of the trabecula, which correspond to the compressive and tensile region in the stressed central position of the femoral head [15]. The principal region that is perpendicular to articular surface is well suited for studying bony changes because it can respond to the transferred loading forces from the diseased articular surface [16,17]. Moreover, it has been shown that the cylinder axis of the principal load-bearing region is aligned with the main trabecular direction; thus, measured changes in biomechanical properties, taking into account the testing direction, can be minimized [17].

Based on the aforementioned theory, virtual cylindrical biopsies (Ø, 5.4 mm; L, 5.4 mm) determined by the semiautomatic contouring method as the volume of interest in the principal load-bearing region were extracted from the reconstructed 3D image 2 mm below the surface of subchondral bone (cubic subvolume, Fig. 1). All scanned

![Figure 1](image-url). Micro-CT view of the femoral head specimens. (A) The location of virtual cylindrical biopsies extracted from the image (cubic subvolume of interest). (B–C) The reconstructed 3D image of the DDH and OA group. CT = computed tomography; DDH = developmental dysplasia of the hip; OA = osteoarthritis.
images were segmented by a low-pass filter to remove noise and then to determine the bone phase by the same threshold. The Micro-CT system software (Scanco Medical AG, Image Processing Language version 4.29d; Switzerland) was used to process the microstructural parameters. The total bone volume fraction (BV/TV), bone trabecular number (Tb.N), and bone trabecular thickness (Tb.Th) were obtained automatically. A model based on the type of the structure was used to measure the structure model index (SMI). An ideal segmented plate structural model has an SMI value of 0, whereas a segmented cylindrical rod structural model has an SMI value of 3. Meanwhile, connectivity density was defined to be a topological parameter of total bone trabecular connections per cubic millimetre.

ITS-based morphological analysis was applied for all bone trabecular cubic subvolumes in the DDH, OA, and NC groups. The bone trabecular network was resolved into individual plates and rods using a volumetric decomposition technique. In brief, with the digital topologic analysis, the skeletal network of the bone trabecular network was transformed into a representation skeleton made of surfaces and curves. Then, the plate and rod shapes of the subchondral microstructure were maintained. Each skeletal voxel was uniquely identified using the digital topologic analysis classification. Using the iterative reconstruction measure previously reported, each voxel was classified into an individual type of either the plate or rod [18]. Based on analysis of the separate bone trabecular plate or rod, a series of ITS-based morphological parameters were calculated at the junctions of both the plate and rod trabecula, i.e., plate and rod bone trabecular volume fraction (pBV/TV and rBV/TV), axial bone trabecular volume fraction along the longitudinal axis (aBV/TV), plate and rod tissue fraction (pBV/BV and rBV/BV), a ratio of trabecular plate versus rod (P/R ratio), plate and rod trabecular number density (pTb.N and rTb.N), plate and rod trabecular thickness (pTb.Th and rTb.Th), trabecular plate surface area (pTb.S), trabecular rod length (rTb.L), and plate–plate, plate–rod, and rod–rod junction density (P–P, P–R, and R–R Junc.D). The definition of these ITS-based microstructural parameters and detailed methods of applications were reported in the previous study [18].

**Micro–finite element analysis**

Scanco Medical Finite Element Software 1.06 (Scanco Medical AG, Switzerland) was used for μFEA to simulate the axial compression tests for each trabecular bone cube in the longitudinal directions. Micro-CT images were converted into 3–4 million micro–finite element models to represent hard tissue (36 μm/voxel) with eight-node brick finite elements. Bone material properties at the tissue level were assumed to be homogeneous linear isotropic material with an isotropic linear elastic material with a Young’s modulus (Es) of 15 GPa and a Poisson’s ratio of 0.3 [19,20]. For the bone segment model, uniaxial compression tests were used for μFEA analysis to compute the reaction force under a displacement equal to 1% of bone segment height along the axial direction. Bone stiffness and failure load were the mechanical parameters extracted from the FEA.

**Histological analysis**

After micro-CT scanning, specimens of the DDH, OA, and NC groups corresponding to the volume of interest were processed for cartilage evaluation through the histological analysis. In brief, serial sections (5 μm) were stained using haematoxylin and eosin, Safranin O-Fast Green, and toluidine blue. Cartilage damage was assessed using the Osteoarthritis Research Society International (OARSI) scoring system in the three groups [21]. Five sections from each sample were stained and measured.

**Statistical analysis**

All values are expressed as means ± standard deviations. Independent two-sided Student t tests and one-way analysis of variance were used to test for significant differences in the ITS-based microstructural parameters and biomechanical properties between the DDH, OA, and NC groups. Pearson correlation coefficients were calculated for ITS-based microstructural parameters with each potential covariate. Sets of covariates with p < 0.05 and independent from each other were selected. These included age, height, and Harris Hip Scores. The ITS-based microstructural parameters were compared again after adjusting for the selected covariates by a generalized multiple linear analysis. To indicate the relationship between the microstructure, biomechanical properties, and cartilage damage in the DDH, OA and NC groups, linear regression analyses were performed; a two-tailed p < 0.05 was considered statistically significant. The statistical analysis was performed using the SPSS 19.0 software package (SPSS Inc., Chicago, IL, USA).

**Results**

The study population consisted of 17 patients with DDH and 16 patients with primary hip OA, and 10 patients with the fractured femoral neck constituted the NC group. All patients had no other systemic diseases, such as haematological disorders or endocrine diseases. As shown in Table 1, compared with patients with DDH, those with OA underwent operative treatment at an older age (p < 0.05). However, patients with DDH and OA were not different in weight and body mass index, indicating that obesity was not an interference factor. The Harris Hip Scores showed no joint function and daily activity differences between the DDH and OA groups.

The means and standard deviations of the micro-CT, ITS-based microstructure analysis for the three groups and the percentage differences between the DDH and OA groups are presented in Table 2 and Fig. 2, respectively. The BV/TV of the OA group was 48.9% higher than that of the DDH group (p < 0.05). The values of the SMI showed significantly morphological differences in the bone trabecula between the DDH and OA groups (p < 0.05). The ITS-based microstructural analysis more particularly confirmed this result. The aBV/TV of the OA group was 62.3% higher than that of the DDH group (p < 0.05). The DDH group had an 84.5% lower pBV/TV and a 30.8% lower pBV/BV than the OA group (p < 0.05). There was no difference in rBV/TV between the DDH and OA groups; however, the OA group had an 82.2% higher P/R ratio than the DDH group (p < 0.05). The results of standard micro-CT also showed that Tb.N and Tb.Th were 25.2% and 46.8% lower, respectively, in the DDH group than in the OA group (p < 0.05). Interestingly, the differences may have been caused by the plate-like trabecula because pTb.N, pTb.Th, and pTb.S were 33.1%, 43.7%, and 39.7% higher in the OA group, respectively (p < 0.05), whereas rTb.N, rTb.Th, and rTb.L were not significantly different in the DDH and OA groups (−2.5%, 0.7%, and 0.8%, respectively; p > 0.05). In addition, the connection and

**Table 1**

Basic information of the DDH, OA, and NC groups.

| Characteristics | DDH (mean ± SD) | OA (mean ± SD) | NC (mean ± SD) | p, DDH versus OA |
|-----------------|-----------------|----------------|----------------|-----------------|
| N               | 17              | 16             | 10             |                 |
| Sex (women/men) | 11/6            | 9/7            | 6/4            | 0.03            |
| Age (years)     | 50.3 ± 3.6      | 57.7 ± 4.8     | 55.8 ± 6.4     | 0.03            |
| Height (m)      | 1.61 ± 0.04     | 1.66 ± 0.05    | 1.63 ± 0.09    | 0.04            |
| Weight (kg)     | 58.3 ± 9.2      | 60.4 ± 11.7    | 61.9 ± 7.2     | 0.65            |
| BMI (kg/m²)     | 21.7 ± 1.2      | 22.1 ± 1.5     | 22.9 ± 2.1     | 0.74            |
| Harris Hip Score(HHS) | 62.1 ± 4.8 | 63.2 ± 3.9 | 78.6 ± 5.9 | 0.48 |
| Alcohol use (%) | 23.5%           | 31.2%          | 20%            |                 |
| Current smokers (%) | 17.6%         | 25%            | 37.5%          |                 |
| Other systemic diseases | No | No | No |                 |

BMI = body mass index; DDH = developmental dysplasia of the hip; NC = normal control; OA = osteoarthritis; SD = standard deviation.
Table 2
The subchondral microstructure of the specimens on standard micro-CT scans and ITS-based scans in the DDH, OA, and NC groups.

| Microstructure | DDH (mean ± SD) | OA (mean ± SD) | NC (mean ± SD) | p, DDH versus OA |
|----------------|----------------|----------------|----------------|-----------------|
| **Standard micro-CT** | | | | |
| BV/TV (%) | 23.18 ± 7.14 | 34.48 ± 5.07 | 30.63 ± 0.06* | |
| Conn. Dens. (1/mm³) | 8.63 ± 4.91 | 11.96 ± 10.23* | 6.77 | |
| SMI | 1.71 ± 0.88 | 0.51 ± 0.92 | 0.76 ± 0.63 | 0.03* |
| Tb.N (1/mm) | 2.02 ± 0.51 | 2.53 ± 0.62 | 2.48 ± 0.74* | |
| Tb.Th (mm) | 0.17 ± 0.02 | 0.25 ± 0.03 | 0.23 ± 0.05 | 0.02* |
| R/BV/based micro-CT | | | | |
| aBV/TV (%) | 10.17 ± 3.72 | 16.48 ± 2.13 | 13.17 ± 0.01* | |
| pbV/TV (%) | 13.64 ± 7.16 | 25.13 ± 6.82 | 21.77 ± 0.01* | |
| rBV/TV (%) | 9.54 ± 3.12 | 9.35 ± 4.26 | 8.86 ± 3.46* | |
| rBV/BV (%) | 56.2 ± 22.1 | 73.5 ± 15.2 | 71.1 ± 13.7 | 0.04* |
| rBV/N (%) | 43.8 ± 11.4 | 26.5 ± 8.7 | 28.9 ± 7.9 | 0.03* |
| P/R ratio | 1.69 ± 0.34 | 3.08 ± 0.48 | 2.46 ± 0.53 | 0.001* |
| pTB.N (1/mm) | 1.84 ± 0.33 | 2.45 ± 0.26 | 2.19 ± 0.47 | 0.001* |
| rTB.N (1/mm) | 2.39 ± 0.45 | 2.53 ± 0.64 | 2.21 ± 0.52 | 0.63 |
| pTB.Th (mm) | 0.19 ± 0.03 | 0.27 ± 0.04 | 0.24 ± 0.06 | 0.01* |
| rTB.Th (mm) | 0.13 ± 0.11 | 0.13 ± 0.08 | 0.12 ± 0.06 | 0.79 |
| pTBs. (mm³) | 0.07 ± 0.02 | 0.11 ± 0.02 | 0.10 ± 0.03 | 0.03* |
| rTB.s. (mm³) | 0.45 ± 0.02 | 0.65 ± 0.01 | 0.47 ± 0.04 | 0.88 |
| P-P.Junc.D (1/mm³) | 8.42 ± 3.35 | 13.46 ± 11.28 | 11.28 ± 0.01* | |
| P-R.Junc.D (1/mm³) | 11.97 ± 3.92 | 18.26 ± 15.34 | 15.34 ± 0.008* | |
| R-P.Junc.D (1/mm³) | 11.97 ± 3.92 | 18.26 ± 15.34 | 15.34 ± 0.008* | |

*The p-value remained significant after adjusting for age, height, and Harris Hip Scores.

junction densities (Conn. Dens., P-P Junc.D, P-R Junc.D, and R-R Junc.D) were the microstructural parameter that indicated the interlinking feature of the trabecular network. The Conn. Dens. was 38.4% higher in the OA group. Moreover, the P-P Junc.D and P-R Junc.D were 59.5% and 52.9% higher in the OA group, respectively (p < 0.05). However, the DDH group did show a greater R-R Junc.D than the OA group (−33.8%, p < 0.05). Furthermore, the generalized multiple linear analysis showed that all ITS-based data existed independently; adjustment for covariates (significant data in basic information) did not influence the significant differences in the microstructural variables between the DDH and OA groups (Table 2).

The results of the biomechanical properties determined on mFEA in the DDH, OA, and NC groups are shown in Fig. 3. The subchondral trabecular bone stiffness and failure load of the DDH group were significantly lower than those of the OA and NC groups. In addition, the analysis of linear regression was used to investigate the relationship between the major subchondral trabecular microstructure and biomechanical properties in the DDH, OA, and NC groups (Table 3). The results indicated that BV/TV and pBV/TV might be the key factors for biomechanical properties of subchondral trabecular bone in the three groups (r² > 0.5, Table 3). Moreover, the correlation coefficients for BV/TV and pBV/TV were higher in the OA group than in the DDH and NC groups, suggesting that subchondral trabecular bone in the OA group had greater biomechanical function than that in the DDH and NC groups when given the same amount of the total trabecula or plate trabecula. However, the microstructurses of the rod trabecula were mainly negatively correlated with biomechanical properties in the DDH group, whereas there was no definite correlation in the OA and NC groups, suggesting that the rod trabecula partially influenced the biomechanical properties in patients with DDH, but not in patients with OA or normal people.

The histological results for cartilage evaluation were shown in Fig. 4A. Cartilage damage was obviously severer with cartilage surface destruction and proteoglycan disorder in the DDH group than in the OA group, and there was less cartilage damage in the NC group. The degenerative changes extended into the deep region in the DDH group. The assessment with the OARSI scoring system showed that the patients in the DDH group had worse cartilage condition than those in the OA and NC groups (p < 0.05, Fig. 4B). Meanwhile, the analysis of linear regression proved that BV/TV and pBV/TV were negatively correlated with cartilage damage in the combined data from the DDH, OA, and NC groups (r² > 0.3, Fig. 4C and D), whereas there was no definite correlation between BV/TV and the OARSI score (r² < 0.1, Fig. 4E).

Discussion
In this study, we used the standard micro-CT, ITS morphological analyses, and the mFEA model to investigate the differences in the microstructural and biomechanical characteristics of subchondral bone between the DDH, OA, and NC groups. We found that the changes of the subchondral trabecular bone microstructure may have specific relationship with the biomechanical properties and cartilage damage in patients with DDH or primary OA.

In the trabecular segmentation technique, the subchondral trabecular bone can be resolved into rod-like or plate-like elements because there were microstructural differences between the two types [11,22]. It is increasingly evident that the types of trabecular bone (plate- or rod-like) play the crucial role in determination of trabecular biomechanical properties as rapid changes in these two types occur with disease progression [23,24]. ITS analysis basically contains the plate- and rod-related microstructures (i.e., the number, thickness, and bone volume of trabecular plates and rods), orientation, and connectivity between plate to plate, plate to rod, and rod to rod. By applying ITS-based micro-CT to decompose the subchondral trabecular bone microstructure, we observed differences in trabecular bone types between the DDH, OA, and NC groups that were not obvious in the conventional outcome measures [25]. The DDH group had lower bone volumes fraction, lower numbers of trabecular plates, thinner trabecular plates, and a lower trabecular P/R ratio than the OA and NC groups. However, rod bone volumes and rod trabecular numbers, thicknesses, and lengths were not obviously different between the three groups. These results provide insightful evidence that the plate-like trabecula holds the dominant position in patients with OA or normal people, whereas in patients with DDH, the plate trabecular bone may reduce gradually with disease onset and progression.

Prior studies have proved that trabecular plate number and thickness did not show obvious loss/reduction in a model of spontaneous OA [8, 26]. However, in this study, we found that plate-like trabeculae in DDH were significantly fewer and thinner, and these changes may not affect the rod-like trabecula. DDH is a disabling musculoskeletal disease leading to secondary OA [27], but the reasons for the differing subchondral microstructure types between DDH and OA remain unknown. Some studies have reported that certain factors, i.e., osteoporosis, diabetes, and male vs. female sex, could cause changes in the subchondral trabecular bone microstructure [28-30]. We know that DDH is a congenital abnormality that definitely affects the development of hip joints in newborns [31]. Thus, the aetiology of the subchondral microstructure changes may be related to abnormal joint load and stress distribution, which is different from primary OA. In addition, in-depth examinations of rod–plate dynamics also revealed that patients with DDH had less connectivity between their plate–plate and plate–rod trabeculae, indicating a less connected and more widely separated trabecular network. We believe that the lower trabecular P/R ratio in DDH is the key reason.
for deterioration of the microstructure and increased trabecular spacing and network inhomogeneity. Our results showed that the decreased trabecular P/R ratio in DDH is primarily due to a reduced number of plate-like trabeculae. Notably, patients with DDH have greater junction densities in rod–rod trabeculae than patients with OA, which suggests that the rod-like trabecula occupies an important position in the subchondral trabecular bone microstructure of the patient with DDH, which influences the interconnected bone trabecular network.

In the μFEA analysis, we further discovered that the biomechanical properties of subchondral trabecular bone (bone stiffness and failure load) were significantly lower in the DDH group than the in OA and NC groups, indicating that biomechanical function of subchondral trabecular bone in patients with DDH was severely compromised. Indeed, the subchondral trabecular bone microstructure is also an important determinant for biomechanics [32]. The microstructural types (plate- or rod-like trabecular bone) are independent and an essential factor that significantly correlates with the biomechanical properties [18]. In addition, the bone mass of the plate-like trabecula is more crucial to trabecular biomechanics than that of the rod-like trabecula. However, the existence of the latter is necessary for maintaining the integrity of the trabecular network and optimising the balance of the physiological environment [11,33,34]. In this study, linear regression showed that the changes of the plate trabecula may be positively correlated with the biomechanical properties in the DDH, OA, and NC groups, whereas the influence of the

Figure 2. Percentage difference in the subchondral microstructures by ITS-based micro-CT analysis (OA group/DDH group). *Significant difference between the OA and DDH groups (p < 0.05). a = axial; BV = bone volume; Conn. Dens. = connectivity density; CT = computed tomography; DDH = developmental dysplasia of the hip; ITS = individual trabecula segmentation; OA = osteoarthritis; P = plate; P–P Junc.D = plate-to-plate junction density; P–R Junc.D = plate-to-rod junction density; P/R ratio = plate-to-rod trabecular ratio; r = rod; R–R Junc.D = rod-to-rod junction density; SMI = structure model index; Tb.L = trabecular bone length; Tb.N = trabecular bone number; Tb.S = trabecular bone surface area; Tb.Th = trabecular bone thickness; TV = total volume.

Figure 3. The biomechanical properties in the DDH, OA, and NC groups that are obtained by micro-finite element analysis (μFEA). *Significant difference compared with the DDH group (p < 0.05); **significant difference compared with the OA group. DDH = developmental dysplasia of the hip; NC = normal control; OA = osteoarthritis.
rod trabecula on biomechanical properties was relatively weak. According to the correlation coefficient (the value of $r^2$) in the linear regression analyses, we deemed that the plate-like trabecula held a more essential role in patients with OA or normal people than in those with DDH. These results confirmed worsened biomechanical properties of subchondral trabecular bone in patients with DDH due to the different microstructures, although the clinical manifestations of the two diseases may be similar. Interestingly, in the DDH group, the rod-like trabecula was negatively correlated with biomechanical properties, whereas there was no definite correlation between rod-like trabeculae and biomechanical properties in the OA and NC groups. We speculated that as the extremely key function of the plate-like trabecula in stabilising the trabecular structure and withstanding joint load, the function of relatively fewer rod-like trabeculae on subchondral trabecular bone is not obvious. However, the rod trabecula occupies a certain percentage of bone in patients with DDH, which not only regulates the integrity of the trabecular bone network but also notably impairs biomechanical properties. Studies have proved that the subchondral trabecular bone changes may be accompanied with the entire primary OA development [8,35]. We considered that the synergistic effect of a decrease in the plate trabecula and an increase in the rod trabecula commonly leads to disease progression in patients with DDH.

We also demonstrated that the subchondral trabecular bone microstructural changes were associated with aggravated cartilage damage. Except for the rod-like trabecula, the total trabecula and plate-like trabecula were correlated with more severe cartilage destruction, which explained that secondary OA caused by DDH could exhibit more serious

| Microstructure | DDH | OA | NC |
|----------------|-----|----|----|
| Stiffness | Failure load | Stiffness | Failure load | Stiffness | Failure load |
| BV/TV | 0.54** | 0.51** | 0.88** | 0.85** | 0.63** | 0.68** |
| Tb.N | 0.23 | 0.21 | 0.46* | 0.42* | 0.37* | 0.29 |
| Tb.Th | 0.37* | 0.31* | 0.71** | 0.68** | 0.55** | 0.61** |
| pBV/TV | 0.73** | 0.66** | 0.81** | 0.82** | 0.69** | 0.71** |
| pTb.N | 0.48* | 0.43* | 0.69** | 0.65** | 0.38* | 0.41* |
| pTb.Th | 0.33* | 0.25 | 0.57** | 0.41* | 0.46* | 0.39* |
| pTb.S | 0.38* | 0.28 | 0.59** | 0.46* | 0.51** | 0.58** |
| rBV/TV | −0.39* | −0.41* | 0.19 | 0.24 | −0.13 | 0.08 |
| rTb.N | −0.36* | −0.32* | 0.21 | 0.23 | 0.14 | 0.17 |
| rTb.Th | −0.32* | −0.24 | 0.11 | 0.16 | −0.07 | 0.15 |
| rTb.L | −0.22 | −0.18 | 0.14 | 0.19 | 0.11 | −0.12 |

*The p-value < 0.05 indicated the significant correlation between the subchondral microstructure and biomechanical properties.

**The p-value < 0.001 indicated the significant correlation between the sub-chondral microstructure and biomechanical properties.

$BV =$ bone volume; Conn. Dens. = connectivity density; DDH = developmental dysplasia of the hip; NC = normal control; OA = osteoarthritis; $p =$ plate; P-P Junc.D = plate-to-plate junction density; P-R Junc.D = plate-to-rod junction density; $r =$ rod; R-R Junc.D = rod-to-rod junction density; $Tb.L =$ trabecular bone length; $Tb.N =$ trabecular bone number; $Tb.S =$ trabecular bone surface area; $Tb.Th =$ trabecular bone thickness; $TV =$ total volume.

Table 3
Correlation coefficient of linear regression between the subchondral microstructure and biomechanical properties in the DDH, OA, and NC groups.

Figure 4. The articular cartilage evaluation and its association with the subchondral bone microstructure. (A) Histological analysis of cartilage damage in the DDH, OA, and NC groups. (B) The analytical results with the Osteoarthritis Research Society International (OARSI) scoring system. (C–E) Relationships between subchondral bone microstructure and cartilage damage. Data were combined from the DDH, OA, and NC groups. *Significant difference compared with the DDH group (p < 0.05); #significant difference compared with the OA group. AC = articular cartilage; BV = bone volume; DDH = developmental dysplasia of the hip; H&E = haematoxylin and eosin; NC = normal control; OA = osteoarthritis; SB = subchondral bone; TV = total volume.
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