Aim: To assess the frequency of bone changes in resected osteonecrotic femoral head (ONFH) specimens at multidetector computed tomography (MDCT) and compare their frequencies between ONFH with limited (<1.5 mm) or advanced (≥1.5 mm) collapse.

Method: Fourteen ONFH were imaged using MDCT (n = 14) and microcomputed tomography (µCT; n = 8). Preoperative staging was performed using radiographs and MRI. Coronal reformats of MDCT images of the specimens were analyzed using the grid overlay method. There were 2,933 grid boxes containing cortical bone and 10,596 containing trabecular bone. Two MSK radiologists assessed in every grid box the presence of interface-related sclerosis, cortical bone interruption, trabecular bone interruption, and trabecular bone resorption. The frequency of grid boxes with bone changes at MDCT was calculated and compared between ONFH with limited (<1.5 mm) or advanced (≥1.5 mm) collapse.

Results: For both readers R1 and R2, there were 1111/10596 (10.5%) and 1362/10596 (12.9%) grid boxes with interface-related bone sclerosis, 557/2933 (19%) and 413/2933 (14.1%) with cortical bone interruption, 796/10596 (7.5%) and 665/10596 (6.3%) with trabecular bone interruption, and 331/10596 (3.1%) and 595/10596 (5.6%) with trabecular bone resorption. The frequency of grid boxes with cortical interruption and trabecular bone resorption was significantly higher in ONFH with advanced than in ONFH with limited collapse. There was no significant difference in frequency of grid boxes with trabecular interruption and interface-related bone sclerosis between ONFH with advanced or limited collapse.

Conclusion: Cortical interruption and trabecular resorption, but not trabecular interruption, were more frequent in osteonecrotic femoral heads with advanced than with limited collapse.

ABSTRACT

Collapse-Related Bone Changes in Osteonecrotic Femoral Heads at Multidetector CT: Comparison between Femoral Heads with Limited and Advanced Collapse

CORRESPONDING AUTHOR:
Charbel Mourad
Hôpital Libanais Geitaoui-CHU, LB
charbel.j.mourad@gmail.com

KEYWORDS:
osteonecrosis of the femoral head; hip; multidetector computed tomography (MDCT); epiphyseal collapse; cortical interruption; trabecular interruption; bone resorption

TO CITE THIS ARTICLE:
Mourad C, Acid S, Michoux N, Awad A, Vande Berg B. Collapse-Related Bone Changes in Osteonecrotic Femoral Heads at Multidetector CT: Comparison between Femoral Heads with Limited and Advanced Collapse. Journal of the Belgian Society of Radiology. 2022; 106(1): 55, 1–7. DOI: https://doi.org/10.5334/jbsr.2735
INTRODUCTION

The occurrence of collapse in osteonecrotic femoral heads (ONFH) is a pivotal step towards osteoarthritis and subsequent total hip replacement. The detection of collapse relies on conventional radiography and MRI [1]. More recently, multidetector computed tomography (MDCT) has been shown to be superior to MRI and radiographs in the detection of subchondral fractures [2–4]. There is a need to better understand bone changes associated with progressive collapse. The aim of this study was to determine the frequency and topology of bone changes in a quantitative analysis performed on MDCT images of resected ONFH specimens using the grid overlay method, and to compare their frequency between femoral heads with limited or advanced collapse.

MATERIAL AND METHODS
1- PATIENTS’ POPULATION
The Institutional Review Board approved the study, with the informed consent of the patients being waived because the study was performed on femoral head specimens that were resected for total hip replacement. An MSK radiologist with four years of experience collected 14 femoral head (FH) specimens with osteonecrosis (ON) from 13 patients: 8 men (46.9 years ±13.9) and 5 women (68.4 years ±16.05). These specimens were resected during hip replacement. The FH specimens were fixated in formalin. In a preliminary study on eight FH specimens, the sensitivity and specificity of in vitro MDCT were calculated in comparison to µCT (see online supplemental material).

2- STAGING OF ONFH AND MEASUREMENT OF ARTICULAR SURFACE COLLAPSE
Staging of ONFH was performed on radiographs and MRI according the ARCO classification [4]. Collapse of the articular surface was measured using the best fitting concentric circle technique [5]. The median collapse was 1.5 mm [1.1–2.1 mm, 95% CI]. There were 10 stage 3-As, 3 stage 3-Bs, and 1 stage 4 ONFH.

3- DATA ACQUISITION AND GRID OVERLAYING
Fourteen resected FH specimens were scanned on a 40-row MDCT (Siemens Somatom Definition 40, Erlangen, Germany). One radiologist with four years of experience viewed the MDCT images on an image viewer with multiplanar capacity. Coronal reformats were obtained and segmented by overlaying a transparent 12 × 12 grid. The anonymized images with the overlaid grid were uploaded on an in-house developed software that enables image analysis (Figure 1 and E1). Further details are provided in online supplemental material.

4- IMAGE ANALYSIS AND LESION DEFINITION
Images were independently analyzed by an MSK radiologist with 30 years of experience (R1), and an MSK radiologist with four years of experience (R2). The reading process was performed in 4 × 4 mm grid boxes using the grid overlay method on in-house developed software (see online supplemental material).

Bone changes at MDCT were defined as follows: interface-related trabecular sclerosis corresponded to thickened trabeculae with a band-like distribution and a concave orientation towards the articular surface. Cortical bone interruption corresponded to a focal interruption of the subchondral bone plate with or without deformity (Figure 2). Trabecular bone interruption corresponded to an interruption of two or more contiguous trabeculae, having irregular, sharp, or angular margins with or without a gas-filled cleft or a dense linear band corresponding to trabecular crushing (Figures 2–3). Trabecular bone resorption corresponded to a lucent zone devoid of mineralized content, with smooth or rounded margins (Figure 2).

Limited FH collapse was defined by the presence of a depression of the FH contours <1.5 mm, and advanced FH collapse by the presence of a depression ≥1.5 mm.

5- STATISTICAL ANALYSIS
Data output was automatically generated on Excel sheets. Frequency of grid boxes with bone changes was compared between FH with limited and advanced collapse using Chi-squared test. Interobserver agreement was computed using Cohen’s Kappa and interpreted according to the scale proposed by Altman [6]. A $p$-value...
<0.05 was regarded as statistically significant. Statistical analyses were performed by using MedCalc Statistical Software version 19.2.6 (MedCalc Software bv, Ostend, Belgium; https://www.medcalc.org; 2020).

**RESULTS**

1- FREQUENCY OF GRID BOXES WITH BONE CHANGES AT MDCT

There were 2,933 grid boxes containing cortical bone and 10,596 grid boxes containing trabecular bone on the MDCT reformats. For R1, there were 557/2933 (19%) with cortical bone interruption, 796/10,596 (7.5%) with trabecular bone interruption, and 331/10,596 (3.1%) with trabecular bone resorption. Results for R2 are given in Table 1.

2- TOPOLOGY OF GRID BOXES WITH NECROTIC LESION AND COLLAPSE-RELATED BONE CHANGES AT MDCT

For R1, grid boxes with collapse-related bone changes (CRBC) at MDCT were more frequent in the anterior, central, and superior thirds of the FH (Table 2). Grid boxes with trabecular interruption were more frequent in the superficial (448/2,621; 17.1%) than in the deep layer (348/7,975; 4.4%). Grid boxes with trabecular resorption were more frequent in the deep (281/7,975; 3.5%) than in the superficial layer (50/2,621; 1.9%). Detailed results for R1 and R2 are provided in Table 2.

For R1, grid boxes with cortical and trabecular bone changes were more frequent within than outside the necrotic lesion (p < 0.001). Detailed results for R1 and R2 are provided in Table 3.
3- COMPARISON OF FREQUENCY OF GRID BOXES WITH NECROTIC LESIONS AND CRBC AT MDCT BETWEEN LIMITED AND ADVANCE COLLAPSE

For R1, grid boxes with cortical interruption were more frequent in FH with collapse ≥1.5 mm (20.49%) than in FH with collapse <1.5 mm (17.02%) (p = 0.018). There was no statistical difference in the frequency of grid boxes with trabecular interruption between FH with collapse ≥1.5 mm (7.13%) than <1.5 mm (8%) (p = 0.094). Grid boxes with trabecular resorption were more frequent in FH with collapse ≥1.5 mm (3.88%) than <1.5 mm (2.15%) (p < 0.001). Detailed results for R1 and R2 are provided in Table 4.

DISCUSSION

In the current study, we assessed bone changes in ONFH specimens using MDCT and found that grid boxes with necrotic lesion and CRBC predominated in the anterior, central, and superior parts of ONFH specimens. We also found that grid boxes with cortical interruption and trabecular bone resorption were more frequent in femoral heads with advanced collapse than in those with limited collapse, with no difference in frequency of bone interruption according to the degree of collapse. These observations suggest that progression of collapse could be more associated with the development of cortical bone fracture and bone resorption than with trabecular bone interruption.

First, the observation that trabecular bone interruption predominated in the superficial layer of the necrotic lesion could be explained by the concentration of mechanical stress on the surface or by its association with cortical interruption [7–11]. The predominance of trabecular bone resorption in the deep regions of the necrotic lesions is associated with the repair process that include increased bone remodeling in vascularized regions surrounding and invading the lesion [11, 12].

Second, the fact that cortical bone interruption was more frequent in ONFH with advanced than with limited collapse is associated with marked deformity of the subchondral bone plate that results from collapse. The observation that bone resorption was more frequent with more advanced collapse suggests that resorption could develop during progressive failure of the FH and is not associated with early fracture [9, 13].

Third, the lack of difference in frequency of trabecular interruption according to the degree of collapse could contradict the general idea that links trabecular fracture with collapse. Trabecular fracture formation could differ from collapse, similar to what is observed in rocks [14]. In ONFH, stress distribution is altered by progressive collapse and development of trabecular fracture can be dissociated from that of cortical fracture [15]. The hypothesis that progression of collapse is associated with bone resorption and cortical fracture but not with trabecular bone fracture deserves further analysis.

Our study has many limitations. In addition to a limited number of specimens, histological confirmation of the findings was not performed. Since µCT can accurately detect cortical and trabecular microfractures in comparison with histology [16, 17], we obtained µCT examinations in eight specimens and demonstrated that MDCT had a high specificity to detect cortical fractures, trabecular fracture, resorption, and sclerosis in comparison with µCT (see supplemental material).

In conclusion, cortical and trabecular bone interruption and trabecular bone resorption are found in collapsed ONFH and their distribution parallels that of the necrotic lesions. Grid boxes with cortical bone interruption and trabecular bone resorption but not with trabecular bone interruption are more frequent in ONFH with advanced than with limited collapse.

Table 1 Frequency of grid boxes with cortical and trabecular bone changes on coronal MDCT reformats of 14 femoral head specimens. R1 and R2 represent two readers.
Table 2 Frequency of grid boxes with CRBC and with necrotic lesion in different thirds regions of the FH in the coronal plane on MDCT reformats.

The denominators represent the total number of grid boxes in every region. (p') Anterior versus Middle; (p") Anterior versus Posterior; (p') Middle versus posterior. (p") = Medial versus Central; (p") = Medial versus Lateral; (p") = Central versus Lateral. (p") = Superior versus Middle; (p") = Superior versus Inferior; (p") = Middle versus Inferior.

| Lesion* | ANT  | MID  | POST | P      | MED  | CENT  | LAT  | P      | SUP  | INF  | P      | RADIAL DIRECTION | P      |
|---------|------|------|------|--------|------|-------|------|--------|------|------|--------|-----------------|--------|
| R1      | NECROTIC | LESEN* | 782/2159 | (36.2%) | 1345/4354 | (30.9%) | 623/2419 | (25.8%) | P < 0.001 | 681/2427 | (28.1%) | 1582/4345 | (36.4%) | 487/2160 | (2.5%) | P < 0.001 | 1713/2273 | (75.4%) | 1030/4632 | (22.2%) | 7/2027 | (0.3%) | P < 0.001 | 823/2232 | (36.9%) | 1972/6700 | (28.8%) |
|         | TROBECULAR | SCerosis | 3242/2562 | (12.6%) | 509/5134 | (9.9%) | 278/2900 | (9.6%) | P < 0.001 | 273/2867 | (9.5%) | 587/5125 | (11.45%) | 251/2604 | (9.6%) | P < 0.001 | 422/2671 | (15.8%) | 650/5447 | (11.9%) | 39/2478 | (1.6%) | P < 0.001 | 203/2621 | (7.7%) | 908/7975 | (11.4%) |
|         | Cortical | interruption | 213/830 | (25.7%) | 236/1199 | (19.7%) | 108/904 | (11.9%) | P < 0.001 | 131/1357 | (6.7%) | 252/846 | (29.8%) | 174/730 | (23.8%) | P < 0.001 | 486/1388 | (35%) | 71/941 | (7.5%) | 0/604 | (0%) | P < 0.001 | NA | NA |
|         | TROBECULAR | interruption | 2432/2562 | (9.5%) | 404/5134 | (7.9%) | 149/2900 | (5.1%) | P < 0.001 | 160/2867 | (5.6%) | 448/5125 | (8.7%) | 188/2604 | (7.2%) | P < 0.001 | 693/2671 | (25.9%) | 99/5447 | (18%) | 4/2478 | (0.2%) | P < 0.001 | 448/2621 | (17.1%) | 348/7975 | (4.4%) |
|         | TROBECULAR | resorption | 162/2562 | (6.3%) | 125/5134 | (2.4%) | 44/2900 | (1.5%) | P < 0.001 | 59/2867 | (2.1%) | 204/5125 | (4%) | 68/2674 | (2.6%) | P < 0.001 | 171/2671 | (6.4%) | 143/5447 | (2.6%) | 17/2478 | (0.7%) | P < 0.001 | 502/2621 | (1.9%) | 281/7975 | (3.5%) |
| R2      | NECROTIC | LESEN* | 884/2159 | (40.9%) | 1323/4354 | (30.4%) | 607/2419 | (25.1%) | P < 0.001 | 685/2427 | (28.2%) | 1585/4345 | (36.5%) | 544/2160 | (2.52%) | P < 0.001 | 1766/2273 | (77.7%) | 1036/4632 | (22.4%) | 12/2027 | (0.6%) | P < 0.001 | 897/2232 | (40.2%) | 1917/6700 | (28.6%) |
|         | TROBECULAR | SCerosis | 421/2562 | (16.4%) | 616/5134 | (12%) | 325/2900 | (11.2%) | P < 0.001 | 332/2867 | (11.6%) | 662/5125 | (12.9%) | 368/2604 | (14.1%) | P < 0.001 | 590/2671 | (22.1%) | 732/5447 | (13.4%) | 40/2478 | (1.6%) | P < 0.001 | 317/2621 | (12.1%) | 1045/7975 | (13.1%) |
|         | Cortical | interruption | 168/830 | (20.2%) | 179/1199 | (14.9%) | 66/904 | (7.3%) | P < 0.001 | 112/1357 | (8.3%) | 186/846 | (22%) | 115/730 | (5.75%) | P < 0.001 | 350/1388 | (25.2%) | 60/941 | (6.4%) | 3/604 | (0.5%) | P < 0.001 | NA | NA |
|         | TROBECULAR | interruption | 228/2562 | (8.9%) | 306/5134 | (6%) | 131/2900 | (4.5%) | P < 0.001 | 108/2867 | (3.8%) | 432/5125 | (8.4%) | 125/2604 | (4.8%) | P < 0.001 | 582/2671 | (21.8%) | 83/5447 | (1.5%) | 0/2478 | (0%) | P < 0.001 | 389/2621 | (14.8%) | 276/7975 | (3.5%) |
|         | TROBECULAR | resorption | 243/2562 | (9.5%) | 248/5134 | (4.8%) | 104/2900 | (3.6%) | P < 0.001 | 116/2867 | (4.1%) | 334/5125 | (6.5%) | 145/2604 | (5.6%) | P < 0.001 | 353/2671 | (13.2%) | 238/5447 | (4.4%) | 4/2478 | (0.2%) | P < 0.001 | 110/2621 | (4.2%) | 485/7975 | (6.1%) |

*Frequency of grid boxes containing the necrotic lesion was calculated in 12 instead of 14 ONFH because the interface was only partially visible.
ABBREVIATIONS

ARCO: Association Research Circulation Osseous
CRBC: collapse-related bone changes
FH: femoral head
MDCT: multidetector computed tomography
ON: osteonecrosis
ONFH: osteonecrotic femoral heads
µCT: Microcomputed tomography

COMPETING INTERESTS

The authors have no competing interests to declare.

AUTHOR AFFILIATIONS

Charbel Mourad orcid.org/0000-0002-6600-5678
Hôpital Libanais Geitaoui- CHU, LB
Souad Acid orcid.org/0000-0002-4128-3233
Cliniques Universitaires Saint Luc, BE
Nicolas Michoux orcid.org/0000-0003-0104-0744
Cliniques Universitaires Saint Luc, BE

Anthony Awad orcid.org/0000-0003-4388-665X
Hôpital Libanais Geitaoui- CHU, LB
Bruno Vande Berg orcid.org/0000-0002-5303-5966
Cliniques Universitaires Saint Luc, BE

REFERENCES

1. Murphey MD, Roberts CC, Bencardino JT, et al. ACR appropriateness criteria osteonecrosis of the hip. J Am Coll Radiol. 2016; 13(2): 147–55. DOI: https://doi.org/10.1016/j.jacr.2015.10.033
2. Stevens K, Tao C, Lee SU, et al. Subchondral fractures in osteonecrosis of the femoral head: Comparison of radiography, CT, and MR imaging. AJR Am J Roentgenol. 2003; 180(2): 363–8. DOI: https://doi.org/10.2214/ajr.180.2.1800363
3. Yeh LR, Chen CK, Huang YL, Pan HB, Yang CF. Diagnostic performance of MR imaging in the assessment of subchondral fractures in avascular necrosis of the femoral head. Skeletal Radiol. 2009; 38(6): 559–64. DOI: https://doi.org/10.1007/s00256-009-0659-0
4. Yoon BH, Mont MA, Koo KH, et al. The 2019 revised version of association research circulation osseous staging system for osteonecrosis of the femoral head. J Knee Surg. 2018; 31(1): 1–7. DOI: https://doi.org/10.1055/a-0711-3274

Table 3 Comparison of frequency of grid boxes with CRBC on coronal MDCT reformats within or outside the osteonecrotic lesions*.

* Frequency of grid boxes containing the necrotic lesion was calculated in 12 instead of 14 ONFH because the interface was only partially visible.

Table 4 Comparison of frequency of grid boxes with necrotic lesions and CRBC at MDCT between ONFH with limited and advanced collapse.
of osteonecrosis of the femoral head. J Arthroplasty. 2020; 35(4): 933–40. DOI: https://doi.org/10.1016/j.arth.2019.11.029
5. Mose K. Methods of measuring in Legg-Calve-Perthes disease with special regard to the prognosis. Clin Orthop Relat Res. 1980; 150: 103–9. DOI: https://doi.org/10.1097/00003086-198007000-00019
6. Altman D. Practical statistics for medical research. London: Chapman and Hall; 1991. DOI: https://doi.org/10.1201/9780429258589
7. Baba S, Motomura G, Ikemura S, et al. Quantitative evaluation of bone-resorptive lesion volume in osteonecrosis of the femoral head using micro-computed tomography. Joint Bone Spine. 2020; 87(1): 75–80. DOI: https://doi.org/10.1016/j.jbspin.2019.09.004
8. Hamada H, Takao M, Sakai T, Sugano N. Subchondral fracture begins from the bone resorption area in osteonecrosis of the femoral head: A micro-computerised tomography study. Int Orthop. 2018; 42(7): 1479–84. DOI: https://doi.org/10.1007/s00264-018-3879-x
9. Gao F, Han J, He Z, Li Z. Radiological analysis of cystic lesion in osteonecrosis of the femoral head. Int Orthop. 2018; 42(7): 1615–21. DOI: https://doi.org/10.1007/s00264-018-3958-z
10. Kenzora JE, Glimcher MJ. Pathogenesis of idiopathic osteonecrosis: The ubiquitous crescent sign. Orthop Clin North Am. 1985; 16(4): 681–96. DOI: https://doi.org/10.1016/S0030-5898(20)30435-1
11. Glimcher MJ, Kenzora JE. The biology of osteonecrosis of the human femoral head and its clinical implications: II. The pathological changes in the femoral head as an organ and in the hip joint. Clin Orthop Relat Res. 1979; 139: 283–312. DOI: https://doi.org/10.1097/00003086-197903000-00040
12. Glimcher MJ, Kenzora JE. Nicolas Andry award. The biology of osteonecrosis of the human femoral head and its clinical implications: 1. Tissue biology. Clin Orthop Relat Res. 1979; 138(138): 284–309.
13. Shi S, Luo P, Sun L, et al. Prediction of the progression of femoral head collapse in ARCO stage 2–3A osteonecrosis based on the initial bone resorption lesion. Br J Radiol. 2021; 94(1117): 20200981. DOI: https://doi.org/10.1259/bjr.20200981
14. Anders MH, Laubach SE, Scholz CH. Microfractures: A review. Journal of Structural Geology. 2014; 69: 377–94. DOI: https://doi.org/10.1016/j.jsg.2014.05.011
15. Utsunomiya T, Motomura G, Ikemura S, et al. Effects of sclerotic changes on stress concentration in early-stage osteonecrosis: A patient-specific, 3D finite element analysis. J Orthop Res. 2018; 36(12): 3169–77. DOI: https://doi.org/10.1002/jor.24124
16. Mourad C, Galant C, Wacheful E, Kirchgesner T, Michoux N, Vande Berg B. Topology of microfractures in osteonecrotic femoral heads at muCT and histology. Bone. 2020; 141: 115623. DOI: https://doi.org/10.1016/j.bone.2020.115623
17. Mourad C, Laperrée K, Halut M, Galant C, Van Cauter M, Vande Berg BC. Fused micro-computed tomography (muCT) and histological images of bone specimens. Diagn Interv Imaging. 2018; 99(7–8): 501–5. DOI: https://doi.org/10.1016/j.dii.2018.01.011

TO CITE THIS ARTICLE:
Mourad C, Acid S, Michoux N, Awad A, Vande Berg B. Collapse-Related Bone Changes in Osteonecrotic Femoral Heads at Multidetector CT: Comparison between Femoral Heads with Limited and Advanced Collapse. Journal of the Belgian Society of Radiology. 2022; 106(1): 55, 1–7. DOI: https://doi.org/10.5334/jbsr.2735

Submitted: 18 December 2021 Accepted: 25 May 2022 Published: 09 June 2022

COPYRIGHT:
© 2022 The Author(s). This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International License (CC-BY 4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited. See http://creativecommons.org/licenses/by/4.0/. Journal of the Belgian Society of Radiology is a peer-reviewed open access journal published by Ubiquity Press.