Early Changes of Articular Cartilage and Subchondral Bone in The DMM Mouse Model of Osteoarthritis

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To examine the early changes of articular cartilage and subchondral bone in the DMM mouse model of osteoarthritis, mice were subjected to DMM or SHAM surgery and sacrificed at 2-, 5- and 10-week post-surgery. Catwalk gait analyses, Micro-Computed Tomography, Toluidine Blue, Picrosirius Red and Tartrate-Resistant Acid Phosphatase (TRAP) staining were used to investigate gait patterns, joint morphology, subchondral bone, cartilage, collagen organization and osteoclasts activity, respectively. Results showed OA progressed over 10-week time-course. Gait disparity occurred only at 10-week post-surgery. Osteophyte formed at 2-week post-surgery. BMDs of DMM showed no statistical differences comparing to SHAM at 2 weeks, but BV/TV is much higher in DMM mice. Increased BMD was clearly found at 5- and 10-week post-surgery in DMM mice. TRAP staining showed increased osteoclast activity at the site of osteophyte formation of DMM joints at 5- and 10-week time points. These results showed that subchondral bone turnover might occurred earlier than 2 weeks in this mouse DMM model. Gait disparity only occurred at later stage of OA in DMM mice. Notably, patella dislocation could occur in some of the DMM mice and cause a different pattern of OA in affected knee.

Osteoarthritis (OA) is a prevalent musculoskeletal disease that causes pain and disability for millions of people. It is characterized by progressive cartilage erosion, subchondral bone turnover and osteophyte formation, amongst other pathological alterations within the joint. Recent findings suggested that subchondral changes might precede cartilage degeneration during OA1. Gene expression of subchondral bone in a rat model was reported dramatically dysregulated before noticeable articular cartilage damage2. The correlation between bone remodeling and OA was elegantly reviewed by Burr and Gallant3. Sanchez et al. found subchondral bone osteoblasts can induce phenotypic changes in human osteoartritic chondrocytes4, downregulate aggrecan but upregulate metalloproteinases expression by chondrocytes5. Not only subchondral bone can affect cartilage in OA, but also cartilage can affect subchondral bone. Stock et al. found UPMA from cartilage can be secreted into the cartilage matrix and may migrate to the cartilage-bone interface, causing subchondral bone changes in the process of OA6. All evidence points to a greater role that subchondral bone plays in OA initiation and progression. However, how articular cartilage and subchondral bone changes during the time-course of OA in the predominant model of post-traumatic OA—the DMM model, has not yet been fully described. Understanding the articular cartilage and subchondral bone changes (especially in the early stage) in this commonly used mouse model will provide more information on the pathophysiology of OA.

The DMM (destabilization of the medial meniscus) model is now the most commonly used surgical model in mice. Its reliability, reproducibility, structural similarity to human OA, its relatively slow progression when compared to other surgical models, and the validation of several pain endpoints make it a good model for early OA exploration. Huang et al. reported the age-dependent changes in the articular cartilage and subchondral bone in

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this DMM model and found age and gender matter in articular cartilage and subchondral bone changes after surgery7. Patricia Das Neves Borges et al. described a comprehensive way to automated segment subchondral bone compartments and assess the bone changes in mice using contralateral knee as controls8. However, subchondral bone changes during the time-course of the disease in the DMM model have not been characterized in detail. This current study aims to examine the early changes of articular cartilage, subchondral bone and their progressions in the DMM model, as well as other parameters of OA including standard histology and gait pattern alterations.

Results

Gait disparity. Gait disparity only occurred at 10 weeks post-surgery in the DMM group compared to SHAM but not at 2- and 5-week time-points after surgery (Fig. 1). DMM mice at 10-week time-point showed lower paw intensity than SHAM mice in the operated left leg. However, the same effect was observed for the contralateral un-operated leg (Fig. 1).

Histological analyses of joint changes after DMM surgery. OA phenotype after DMM surgery was confirmed by histology using toluidine blue (TB) staining. Regional proteoglycan loss and chondrocyte clustering occurred on the medial compartments of the joint at 2 weeks post-surgery, especially the medial tibial plateau (MTP) (Fig. 2A,b, white arrow). Collagen disorganization was also shown in this small region in adjacent sections by picrosirius red (PR) staining (Fig. 2A,d, yellow arrow). Cartilaginous tissue formed on the medial edge of tibia and stained strongly with TB at 2-week time-point in DMM mice (Fig. 2A,b, black arrow), with collagen fibril organization similar to that of articular cartilage (Fig. 2A,d, white arrow). 5 weeks post-surgery, mild to moderate cartilage damage can be found in the surgical joints of DMM mice. Cartilage erosion and fibrillation, as well as large osteophytes were noticed on the MTP. Proteoglycan loss and fibrillation were also present on the medial femoral condyle (MFC) (Fig. 2A,i, white arrow). Cartilage lesion on the MTP was deep into the calcified zone 10 weeks post-surgery(Fig. 2A,r). Endochondral ossification can be observed in the osteophyte with bone marrow formed within the osteophyte (Fig. 2A,r, black arrow).

Osteoarthritis Research Society International (OARSI) histopathology scoring system was applied to evaluate OA severity (Fig. 2B). On the medial compartments of the joint, the tibial plateau developed earlier (starting at 2 weeks) and more severe OA compared to femoral condyle (starting at 5 weeks) (Fig. 2B) at the same time-points. Although not statistically significant and much milder, lateral compartments showed a similar trend (Fig. 2B).

TRAP staining. TRAP positive cells were found in the subchondral bone of DMM mice starting at 5 weeks post-surgery, with most of these cells located at the site of osteophyte formation (Fig. 3). Very little positive staining was found in the subchondral bone of the SHAM group, although the expected staining at growth plate was observed (Fig. 3).
Subchondral bone changes after DMM surgery. 3D reconstruction of joints from micro-CT analyses is shown in Fig. 4. Irregular bone surface and abnormal patellar shape (ostearthrosis formations) was noticed in DMM mice as early as 2 weeks post-surgery (Fig. 4A, hollow arrow) and became more obvious at later time-points (Fig. 4E, hollow arrow). Osteophyte formation of the MTP was noticed at 5 and 10 weeks post-surgery, with
those at 10 weeks being more severe (Fig. 4E,I, white arrow). In the coronal plane in the middle of the knee joint, subchondral bone of the MTP didn’t show difference at 2 weeks in DMM Group compared to SHAM (Fig. 4B), but became denser at later time-points (Fig. 4E, circled). BMDs of all compartments are shown in Fig. 5. As seen in the coronal plane images, BMDs of DMM animals showed no statistical differences comparing to their SHAM controls at 2 weeks, with DMM animals displaying a larger spread of their BMDs (Fig. 5A). Clearer differences were detected between DMM and SHAM at both 5 and 10 weeks post-surgery, with increased BMDs in DMM animals at MTP (5 and 10 weeks) and MFC (10 weeks only) (Fig. 5A,B). While the lateral compartments showed no difference between two groups (Fig. 5C,D). No statistical significance was detected among the BMDs of the contralateral right knee (data not shown). BV/TVs were also measured in this study. Statistically differences were only detected in MTP at 2 weeks post-surgery with DMM group presenting a higher BV/TV than their SHAM controls (data not shown). Due to the scan limitation, trabecular thickness or separation couldn’t be calculated in the free version of MicroView 2.5.0 (Parallax Innovations Inc.).

Subluxation/dislocation of patella after DMM. In a few of our mice (5 of 27, 2 mice of 2-week time-point and 3 of 10-week time-point) that underwent DMM surgery, subluxation/dislocation of the patella was noted as early as 2 weeks post-surgery by micro-CT as well as histological staining (Fig. 6A,C). In these mice, joint condition deteriorated faster as shown by 3D reconstruction and histology. The LTP seemed to have more severe OA than the medial side (Fig. 6E,F), which might be due to abnormal mechanical force on the lateral side of the knee caused by the subluxated/dislocated patella. Osteophyte formation can be observed on the lateral side of the femoral condyle and TRAP staining showed highly increased osteoclast activity within the osteophyte which indicating increased bone remodeling (Fig. 6B,D). Of note, mice with tibia dislocation were not included in this study.

Discussion

Subchondral bone has been considered as one of the most promising targets for OA drug development. Botter et al. reported that in ADAMTS5−/− mice, whose joints were protected from OA showed minor subchondral bone changes after DMM surgery. Carlson and colleagues showed that a thickened subchondral plate was present before apparent cartilage deterioration in a spontaneously developing primate OA model. Similar bone changes that occurred earlier than cartilage damage were noticed in other experimental animal models. In Waung’s work, they investigate a digital X-ray microradiography to quantified subchondral BMC in the medial tibial plateau 4 and 8 weeks after DMM surgery and showed increased BMC at both time-points in DMM mice.

This study confirmed in the DMM mouse model, mild cartilage lesion was shown in some of the mice 2 weeks after surgery, presenting with regional proteoglycan loss, chondrocyte clustering and cartilaginous osteophyte formations at the medial side of the surgical joint, which is consistent with the report by Loeser and Das Neves Borges. OARSI scores were significantly higher in DMM joints compared with SHAM at all time-points. Mild to moderate articular cartilage damage was observed at 5 and 10 weeks after surgery. Osteophyte formation progressed over time, and endochondral ossification within the osteophyte can be seen at 10 weeks post-surgery.

Gait disparity appeared only at 10 weeks post-surgery but not at earlier time-points, which could be due to the progressive development of OA pain behaviors reported recently. These changes were relatively mild and,
surprisingly, affected both hind limbs (not only the operated one). In line with these findings, it has been reported that behavioral changes in rodent OA are usually manifested as reductions in total locomotor activity levels (e.g., active time and distance travelled) other than gait pattern alteration. More detail of gait study in mice could be found in our review.

However, Micro-CT results showed that BMDs of MTP in the DMM-operated animals were more variable than SHAM controls at 2 weeks post-surgery (Fig. 5A), which suggests that the destabilization of medial meniscus possibly leads to early loss of subchondral bone mineralization in the MTP of some mice as early as 2 weeks following DMM surgery. Clearer differences were detected at both 5 and 10 weeks post-surgery, with increased BMD in DMM-operated animals at both the MTP and the MFC (Fig. 5A,B).

It has been reported that a large number of genes were dysregulated as early as 6 hours after DMM surgery by microarray analysis of gene expressed in the whole joint, including known pathogenic OA genes Mmp3, Adamts5, and Ccl2. Furthermore, regulation of selected genes was abrogated after immobilization of the surgical joint, and OA was prevented up to 12 weeks post-surgery by immobilization. This indicated that changes in the mouse joint following DMM were rapid and highly mechanosensitive, despite the fact that surgery itself was considered mild. It appears likely that the subchondral bone responds immediately to the altered mechanical load within the joint, and that this response causes or at least partly contributes to the subsequent pathophysiological events of other structures within the joint. However, almost all genome-wide studies carried out in the DMM model used the whole joint instead of more specific joint tissues due to the small size of the mouse joint. Further investigation
of different joint tissues respectively (e.g., subchondral bone, meniscus) would provide additional insights into which tissue responds first.

Low subchondral bone mineralization was present in some animals at the 2-week time-point in our study. However, no statistical significance was found at this time-point. In the meantime, BV/TV of MTP at 2 weeks post-surgery was significantly higher in DMM mice (data not shown), suggesting that increased bone remodeling occurred at very early stages (earlier than 2 weeks post-surgery) of OA in DMM mouse model. In addition, the presence of chondrocyte/osteophyte formation shown in both our and Loeser’s results as early as 2 weeks post-surgery suggested that changes in the subchondral bone caused by DMM were rapid after the surgery. However, no increased osteoclast activity was found in subchondral bone at the 2-week time-point, which suggests that loss of subchondral bone mineralization due to increased osteoclast function might have occurred earlier.

Mice of 5- and 10-week time-points represent late stages of OA in terms of subchondral bone with bone sclerosis, significantly higher BMD in medial compartments and severe osteophyte formation around the joint compared with the SHAM. Positive TRAP staining can mostly be found in the site of osteophytes at these two later time-points. These data suggest that bone sclerosis occurs in the DMM model similar to human OA and other animal models.

Figure 5. Bone mineral density (BMD) of subchondral bone in the DMM model. At 2 weeks post-surgery, BMDs of the MTP in DMM mice didn't showed statistical significant differences comparing to their SHAM controls. However, DMM mice showed a larger spread of BMD at 2 weeks post-surgery (A). Clearer differences were detected between DMM and SHAM at both 5 and 10 weeks post-surgery with higher BMDs in DMM mice. (A). BMDs of the MFC were higher in DMM mice 10 weeks after surgery (B). There was no significant differences found in the lateral compartments between DMM and SHAM at all time-points (C,D). Error bar stands for the 95% confidence interval.
changes driven by osteoclasts/osteoblasts/osteocytes activities as well as gene and protein expressions in the subchondral bone at the early time-points is needed for a better understanding of the molecular mechanism driving bone changes during the initiation of OA in the DMM model.

Materials and Methods

Animals and OA induction. All mice were housed on a 12-hour light/dark cycle with unrestricted access to standard mouse food and water. C57/B6 wild type mice were randomly designated into 2 groups: DMM and SHAM, and three termination time-points (2, 5 and 10 weeks post-surgery). DMM or sham surgery was performed when mice were 12 weeks of age. All animals were housed and used in accordance with Western University’s Animal Care and Use Guidelines. The protocol was approved by the Animal Care and Veterinary Services of Western University (protocol#: 2015-031). Surgery was performed on the left knee joint of mice as previously described20,21. A single dose of Amoxicillin (Novopharm, Toronto, Ontario, Canada) was injected subcutaneously right after the surgery to prevent joint infection (20 mg/kg). Buprenorphine (Schering-Plough, Herfordshire, UK) was administered twice (0 and 4 hours after surgery) subcutaneously to minimized pain (0.05 mg/kg). Mice were housed in colony cage after the surgery (up to 6 mice in one cage) and running wheels were put in two days after the surgery to encourage exercise. Excluding the mice that had dislocated patella after surgery (discussed in Results and Discussion Sections), the final size of each group was as followed: DMM group (2W: 6, 5W: 8, 10W: 6), SHAM group (2W: 5, 5W: 7, 10W: 6).

Gait analysis. Mice were weighed and transferred to the Behavior Study Suite at Robarts Research Institute at Western University two days prior to gait acquisitions. Gait patterns and relative parameters were captured by the Catwalk® gait analysis system (Noldus Inc.) as described22. Briefly, mice were placed on a glass platform located in a dark room and allowed to move freely. A light beam from a fluorescent lamp illuminates the platform. Where the paw touches the glass surface, an image of the paw print is recorded by a camera underneath. Gait was captured and parameters was obtained from the Catwalk® software for statistical analysis.

Figure 6. Possible subluxation/dislocation of the patella after DMM surgery. Safranin-O staining showed the out-grown osteophyte stained strongly for GAG (A). TRAP staining demonstrated increased number and activity of osteoclasts (B, D). μCT showed out-grown osteophyte formation (C, arrow) and deteriorated joint condition (C). Toluidine Blue staining showed cartilage degeneration on medial (E) and lateral (F) compartments at 10 weeks post-Surgery in DMM mice with patella dislocation.
Limb harvest and µCT. Following gait acquisition, mice were sacrificed by CO₂ asphyxiation and both hind limbs were dissected immediately and fixed in 4% paraformaldehyde for 24 hours after the removal of skin and extra tissue.

Mouse hind limbs were then scanned using an eXplore RS in vivo x-ray µCT scanner (GE Healthcare Biosciences, London, ON, Canada). Calibrating standards for air, water, and a cortical bone mimic, SB3 (Gammax RMI), were included with the specimens. The scanning protocol consisted of 900 sy-xray projection images, spaced at an angular increment of 0.4 degrees over a single 6.5-hour gantry rotation at a peak energy of 80 kVp, 450 mA tube current, and 4500 ms per exposure. 5 frames (exposures) per view angle were averaged to reduce noise. Using a filtered-backprojection algorithm, the data sets were reconstructed into 3D volumes with an isotropic voxel tube current, and 4500 ms per exposure. 5 frames (exposures) per view angle were averaged to reduce noise. Using a filtered-backprojection algorithm, the data sets were reconstructed into 3D volumes with an isotropic voxel

Histology. Both hind limbs were then put in 5% ethylenediaminetetraacetic acid (EDTA) in phosphate buffered saline (pH 7.0) for decalcification for 10–12 days after the µCT scan. Decalcified limbs were further processed and embedded in paraffin in frontal orientation, with serial sections taken at a thickness of 4 µm and dried in a 37°C oven for 24 hours, then stored in room temperature. Histology was generally performed as described²³,²⁴. All sections for staining were deparaffinized in two changes of xylene, rehydrated in ethanol with decreasing concentration ending in water, and stained as follows. General histology of cartilage and bone was assessed in toluidine blue (TB, 0.04% toluidine blue in 0.2 M acetate buffer, pH 3.75–4.25, 5 min), Safranin-O and fast-green (0.02% fast green for 30 minutes, 1% acetic acid for 10 seconds, and 1.5% Safranin O for 3 minutes). Collagen was stained with picrosirius red (PR, 0.1% Sirius red in saturated picric acid solution, 60 min; 0.5% acetic acid washes) and polarized microscope was used to evaluate the size and organization of collagen fibrils of articular cartilage and subchondral bone. Osteoclast activity was detected by tartrate-resistant acid phosphatase (TRAP, Sigma Canada) staining with minor alterations to the manufacturer's instructions²⁵. All images were taken using a Leica DFC295 digital camera mounted on a Leica DM1000 microscope and Leica Application Suite 3.8.0.

OARSI scoring. OARSI scoring system was used to evaluate OA severity in three sections with different depth after TB staining. Sections were blinded and scored by three different experienced scientists. Average scores were used in statistical analyses²¹.

Statistical analyses. Statistical analysis was performed in Graph Pad Prism 7. Median and 2.5–97.5 percentile were calculated for OARSI score, BMD (error bar means 95% confidence interval). Gait data (Paw intensity/Weight) followed a lognormal distribution and were log-transformed for parametric test. OARSI scores and BMD data were analyzed by One-way ANOVA. Multiple t tests using Holm-Sidak method was then used to analyze the differences between DMM and SHAM at each time-point. In all cases, p < 0.05 was considered significant.

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Acknowledgements
We would like to thank Tracey Hill for assistance in DMM surgery, and the rest of the Beier lab for their scientific discussion and support in this research. Hang Fang was enrolled in a Ph.D. program in the Third Affiliated Hospital of Sun Yat-Sen University, Guangzhou, P.R. China and funded by China Scholarship Council for a two-year period exchange at Western University, London ON, Canada. He is now an attending orthopedic surgeon in The Third Affiliated Hospital of Southern Medical University, Guangzhou, P.R. China. Frank Beier is the recipient of a Canada Research Chair. This work was supported by an operating grant from the Canadian Institutes for Health Research to F.B. (MOP 86574), and from Science and Technology Planning Project of Tianhe District, Guangdong, China (201404KW005), Medical Scientific Research Foundation of Guangdong Province, China (A2015422), National Natural Science Foundation of China (81371990, 81601945) and a scientific research grant for young investigators from Southern Medical University to H.F (PY2014N086). All funding sources (including CIHR, NSFC and et al.) had no role in the design and interpretation of this study.

Author Contributions
H.F. performed most of research, draft this paper and prepared all figures; I.W. performed some of the data analysis and figure preparations; I.W. performed surgery; C.N. and D.H. performed micro-CT scan and image reconstruction, D.C. and F.B. designed the study and edited the paper. All authors approved the current version of the manuscript.

Additional Information
Competing Interests: The authors declare no competing interests.

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