Micro-scale assessment of the postoperative effect of anterior cruciate ligament reconstruction preclinical study using a 7.1T micro-magnetic resonance imaging

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Abstract. High-field micro-magnetic resonance imaging (MRI) scanning may provide additional information for quantitative analysis of graft bone healing processes, thus serving as a promising supplementary method in graft and bone healing evaluation following anterior cruciate ligament reconstruction (ACLR) surgery during preclinical studies. The present study included 12 New Zealand white rabbits that underwent ACLR with polyethylene terephthalate (PET) ligament. At 4, 8, and 16 weeks following surgery, 4 rabbits were euthanized and knee joint samples were harvested for a 7.1T micro-magnetic resonance imaging (MRI) scan. The graft bone tunnel diameter and signal noise ratio (SNR) at the region of interest (ROI) were measured. Hematoxylin-eosin staining was performed at each time point to verify the graft bone healing process in histology. The bone tunnel diameter at the graft tunnel interface decreased over time in both femoral and tibial parts. Notably, the tunnel size was smaller than the diameter of the drilling Kirschner wire that was used to observe the femoral part and proximal site of the tibial part at 16 weeks following surgery. SNR research demonstrated that both the femoral and tibial part PET ligaments selected in the ROI exhibited a marked increase in SNR from the initial 4-week results. The micro-MRI result was consistent with that of histological analysis. Micro-MRI scanning was applied in an animal model that underwent ACL reconstruction surgery with PET ligament, and it was determined that micro-MRI is promising in quantitatively observing graft bone healing processes directly with a focus on graft tunnel distances and SNRs.

Introduction

The anterior cruciate ligament (ACL) is an important stabilizing structure of the knee, and ACL injury is listed as one of the most common injuries in sports medicine; >120,000 ACL reconstruction (ACLR) surgeries are reportedly performed annually in the United States alone (1). Early rehabilitation and return to sports following an operation are considered core objectives following an ACLR. To reach these targets, graft and bone healing is crucial in the postoperative course; therefore, various studies have been conducted under the scope of graft and bone healing processes (2-4).

In clinical studies, the graft and bone healing condition is primarily evaluated indirectly via physical examination for ligament stability (e.g. Lachman test or pivot-shift test (5) and return to sports capability (e.g. single-leg hop test, and isokinetic evaluation (6), subjective functional scoring scales (e.g. International Knee Documentation Committee (IKDC) Knee Ligament Evaluation Form (7) and a personal questionnaire regarding patients' return to sports, activity frequency and condition. Radiographic assessments are typically performed in the form of bilateral weight-bearing X-ray scans based on the IKDC knee arthritis grading system (8). Recent studies have utilized a magnetic resonance imaging (MRI) scan to evaluate graft maturity; this approach may illustrate graft and bone healing conditions at each time point more clearly (9-11). For animal studies, histological examination is thought to be the most suitable method in the evaluation of graft bone healing processes. Specimens are typically harvested, fixed, embedded, sectioned and prepared for multiple staining protocols to evaluate different components, and graft and bone healing conditions are revealed clearly in slides under microscopy. However, a number of limitations are also present with histological examination modalities, as histology studies analyze graft and bone healing in segments, focusing more on the regional healing condition and failing to elaborate on the entire graft bone healing result.

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Micro-MRI has previously been introduced in preclinical studies in orthopedics research (12,13). Due to its superiority in high spatial resolution, high soft tissue contrast and visualizing tendons, ligaments, cartilage, menisci and other non-osseous tissues directly, micro-MRI with a high field coil has been applied in a variety of osteoporosis, osteoarthritis and cartilage lesion animal studies (14,15). Disease progression and cartilage derogation have been identified in animal models, including mouse, rat, rabbit and dog models (14-16). In the study of ACL, however, few studies have been performed with micro-MRI scanning as a possible method for evaluating graft bone healing processes. This may be associated with difficulties in accessing devices or lack of previous experience.

In the present study, high-field micro-MRI was performed on animal models that underwent ACLR surgery at different time points following surgery to assess progressive changes in tunnel diameter and MRI signal noise ratio in graft and bone structures. It was hypothesized that high-field micro-MRI scanning may provide additional information on graft bone healing processes, thus serving as a promising supplementary method in graft and bone healing evaluations following ACLR surgery during preclinical studies.

Materials and methods

Animal ACLR model. The present animal study was reviewed and approved by the Animal Experiment Ethics Committee of the College of Pharmacy, Fudan University (Shanghai, China). A total of 12 skeletally mature male New Zealand white rabbits (age, 1 year; weight, 3.4±0.4 kg) were provided by the Animal Efficacy Evaluation Center, College of Pharmacy, Fudan University (Shanghai, China). Rabbits were housed at 20°C with a 40-60% humidity under a 16/8 h light/dark cycle. Animals also received ad libitum access to food and water. Following 2 weeks of quarantine, ACLR surgeries were performed on the rabbits’ right legs. Briefly, the rabbits were fixed on the surgery table in a supine position following general anesthesia with 3% pentobarbital (30 mg/kg body weight; Shanghai Wokai; Sinopharm Chemical Reagent Co., Ltd. Shanghai, China). A medial parapatellar incision was made, and the native ACL was exposed and transected. Joint laxity was confirmed by a Lachman test following removing the native ACL. A 3.0-mm diameter Kirschner wire was used to drill the bone tunnels from the ACL footprints and remnants towards the femoral and tibial insertion sites of the ACL. The polyethylene terephthalate (PET) ligament fabricated prior to surgery was fed through the tunnel with the help of a PDS II wire (Ethicon, Inc., Cincinnati, OH, USA). The femoral and tibial ends of the ligament were fixed with the periosteum adjacent soft tissue. Following graft transplantation, consecutive cycling loads were performed 20 times to ensure the ligaments were functional. Then, the wounds were sutured in layers (Fig. 1). Postoperatively, rabbits were returned to their cages and allowed free cage activity without immobilization. Intramuscular prophylactic antibiotic injections (800,000 IU penicillin) and wound cleaning were performed immediately following the operation and once daily for 3 consecutive days.

Rabbits were euthanized via administration of 3% pentobarbital (100 mg/kg) in the auricular vein at 4, 8 and 16 weeks following surgery and the femur-graft-tibia complex specimens (n=4 at each time point) were harvested immediately from the rabbits following sacrifice. Soft tissues around the complex were well preserved.

Micro-MRI scan and data analysis. In all rabbit specimens, MRI examinations were performed with a 7.1T Biospec 70/20R MRI magnet (Bruker AXS GmbH, Karlsruhe, Germany). An automatic shimming process was performed at the beginning of each scan to homogenize the magnetic field. Specimens were placed supine on the animal bed with a 38-mm transmit-receive cylindrical birdcage radio frequency coil fixed to it. A two-dimensional T2-TurboRare sequence (repetition time, 3,500 ms; echo time, 26 ms; flip angle, 90°; field of view, 16x16 mm; slice thickness, 0.35 mm; gap between slices, 0 mm; acquisition resolution, 256x256 matrix size) was used in the scanning.

The PET ligament and bone tunnel interface distance were measured in axial position images, and the distance was identified as the largest diameter from the center of the ligament fiber to the edge of the bone tunnel. The tunnel diameters at the proximal, middle and distal sites in the bone tunnel were measured simultaneously, both in the femoral and tibial tunnels (Fig. 2).

The signal noise ratio (SNR) is a measure of signal strength in a certain area relative to background noise. SNR analyses were performed using ImageJ 1.51e (National Institutes of Health, Bethesda, MD, USA). Briefly, the range of interest (ROI) was set based on the oblique sagittal plane in which the PET ligament graft and bone tunnels were mostly linear and parallel to each other in the longitudinal direction. The PET ligament in the femoral bone tunnel was highlighted in a linear region with a width of 10 mm set on the boundary of the
joint surface side parallel to Blumensaat’s line. The PET ligament in the tibia bone tunnel was highlighted in a linear region with a width of 10 mm set on the boundary of the joint surface side parallel to the joint surface at the opening site of the bone tunnel. The signal intensity of the background was measured at 1 cm anterior to the tibia tuberosity. The SNR was calculated as signal intensity in the ROI divided by signal intensity in the background area (Fig. 3) (17). Two sports medicine surgeons who were blinded to the sample information at the measurement moment were requested to provide an analysis, and the mean values were regarded as final results.

**Histological examination.** Bone-graft complexes were harvested from each group at 4, 8 and 16 weeks (n=4 at each time point) following micro-MRI and fixed in 10% formalin for 48 h at 10°C immediately prior to histological examination. Specimens prepared for undecalcified bone slicing were embedded in resin and sectioned with a sliding microtome (SM2500; Leica Microsystems GmbH, Wetzlar, Germany) perpendicular to the longitudinal axis of the femur and tibia tunnel at a thickness of 5 µm. The sections were stained at 20 kC for 6 h with hematoxylin-eosin to evaluate the graft, host bone and the interface between them. Images were visualized via inverted microscopy (IX71SBF2; Olympus Corporation,
Tokyo, Japan) and captured with DP72 Manager (Olympus Corporation).

Statistical analysis. Statistical analysis was performed using SPSS 22.0 (IBM Corp., Armonk, NY, USA). Data are presented as the mean ± standard deviation. One-way analysis of variance with Fisher’s Least Significant Difference was used to analyze the mean diameter between the graft and bone tunnel in the femoral and tibial tunnels, and the mean SNR at the femur and tibia. P<0.05 was considered to indicate a statistically significant difference.

Results

Bone tunnel diameter measurements. The bone tunnel diameter at the graft tunnel interface site was largest at 4 weeks following surgery (femoral part, proximal site=3.55±0.18 mm, middle site=3.33±0.41 mm and distal site=3.20±0.41 mm; Fig. 4; tibial part, proximal site=3.15±0.41 mm, middle site=3.68±0.43 mm, and distal site: 3.80±0.19 mm; Fig. 5). Compared with samples at 4 weeks, the size was decreased in the samples collected at 8 and 16 weeks following surgery. At 8 weeks, the tunnel diameter at the middle site in the femoral part and at the proximal site in the tibial part showed significant shrinkage when compared with the results at the same site at 4 weeks (P<0.05). At 16 weeks following surgery, all 6 measuring sites, except the tibial medial site, in the tibial tunnel reported significantly decreased tunnel diameters in comparison with the 4-week results (P<0.05). It is worthwhile to note that in certain measuring sites (proximal, middle and distal site of the femur part and proximal site of the tibia part; diameters=2.93±0.69, 2.88±0.40, 2.58±0.33 and 2.92±0.35 mm, respectively), the tunnel diameter was smaller than the diameter of the Kirschner wire used in tunnel drilling at 16 weeks following surgery.

SNR analysis. SNR analysis revealed that both the femoral and tibial PET ligaments selected in the ROI exhibited a marked increase in SNR from the initial 4-week results (femoral at 4 weeks, 5.68±0.36; tibial at 4 weeks, 5.08±1.28). However, statistical significance was only detected in the comparison between 16 and 4 weeks in the SNR value in the tibia (P=0.0464; Fig 6).

Evaluation of graft bone interfaces. As presented in Fig. 7, the interface between host bone and the PET ligament graft was
The graft and bone healing processes following surgery heavily affect the therapeutic effects as well as patients' abilities to return to sports, which has always been emphasized in sports medicine practice (1). In the hope of accelerating the ligamentation process, various methods have been applied in preclinical studies (18,19). To analyse the association between those enhancement solutions and graft bone healing processes, daily observation, gross observation, anatomic study, biomechanics test, histology, and medical imaging were included in the present study. New Zealand Rabbits are frequently utilized in animal experimental research. The size and morphological similarity to human knee joint made the animal an appropriate choice in ACL animal studies. ACLR surgery has also become a common choice for the treatment of ACL injury.

Daily observation, though not performed in the present study due to resource limitation, may be carried out prior to and following the ACLR surgery. Items listed in the daily observation include, but are not limited to, the animal's weight, hair color, wound healing condition, knee joint flexion and extension angle regarding range of motion and active activity level (19). These indicators partially reflect the animal's quality of life following surgery and, to a certain extent, the graft functions in daily movement. However, difficulties in quantitative analysis with the majority of these indicators impede further progress of indicator analysis (20). The most frequently used assessment system of ACLR may be the adjusted Oswestry Arthroscopy Score (OAS) on a macroscopic assessment. Items listed on the evaluation form include levels at the insertion site with the surrounding cartilage, integration at the insertion site with the surrounding bone, stiffness of the tendon graft, appearance of articular surface and color of the articular surface, with a total score of 10 (21). Gross evaluation provides investigators with an initial semi-quantitative analysis of the samples and set a rational foundation for further study. Cheng et al (18) recently used the OAS in the assessment of fresh specimens harvested from ACLR rabbit models. As an assessment scale output results in ordinal data, some evaluation bias is inevitable due to results at boundaries between the two classes. Anatomic study following the sample harvest provides an opportunity to further understand general anatomical structure following graft transplantation and its association with structures adjacent to it (22,23). Like the daily observation, those studies failed to offer quantitative data for assessment. Molecular biology study refers to the testing of certain biomedical markers associated with ligament integration processes based on the serum or synovium extracted during or following caging (24-26). By conducting molecular biology study, researchers may determine whether certain enhancement methods induce a concentration rise in certain growth factors and proteins that have been renowned for their cartilage or bone incorporation capacity. Functional biological networks, however, should not be neglected, as it is believed that signals and factors function through networks instead of a simple downstream signal or factor. Thus, a confounding bias should be noted. Biomechanical testing has been emphasized in recent years in ACLR research such as testing dynamic stress loads and stiffness with certain fixing devices including a Biopuls prosthetic limb fixture test and a three-point bending/breaking device (27,28). The boundedness of biomechanical testing is a result of the improper design and use of a fixation device at the beginning of a test. Certain vehicles, including prosthetic limb fixture testing devices may fail to stretch the knee joint in a functional way, resulting in misleading data (27,28). Histology study is crucial in ACLR preclinical study. Slides stained with hematoxylin and eosin, toluidine blue, Masson's trichrome stain, and safranin O help researchers observe grafts, newly born bone, graft-bone interface and types of collagen specifically and clearly (29-32). Examination via histology also has some disadvantages. The mechanical and biological properties of the whole ACL structure are evaluated using localized sections. Furthermore, tissue wastage during slicing hinders continuous observation of the graft bone complex.

Imaging study methods used previously in ACLR preclinical study primarily involve X-ray and micro computed tomography (micro-CT) scans. The former is beneficial in knee joint arthritis detection based on the Kellgren and Lawrence system and IKDC scale, offering second-hand ACL graft healing information (29). Micro-CT scan has been used in the study of bone architecture and mineral density during preclinical investigations of osteoporosis in several animal models (31,32). High contrast quality enables a non-invasive method in evaluating changes in bone stereology and bone trabecular structure following ACLR (30,31). However, the ligament grafted in the

Figure 7. Representative histologic images of the graft bone interfaces at (a) 4, (b) 8, and (c) 16 weeks following surgery (hematoxylin-eosin staining; magnification, x200). Scale bar=100 μm. IF, interface; HB, host bone; G, artificial ligament graft.
surgery is not markedly developed under screening as ligament grafts are typically made with materials that are low density and are close to the air background.

Micro-MRI is a novel study solution tool in ACLR preclinical studies. With the use of small high-efficiency coils and high field imagery, micro-MRI is superior in soft tissue imaging. In osteoarthritis animal models, non-invasive quantitative high field micro-MRI is more specific for quantitatively detecting menisci and cartilage lesions at different stages when compared with the results used with common 3T MRI devices (11,16,30).

With the use of micro-MRI, the present study demonstrated that graft tunnel diameters decreased over time following ACLR surgery. The preliminary result was rationalized as a micro-bone fracture due to drilling in the tunnel formation, which was unavoidable, and the following callus formation and remodeling are natural courses for fracture healing, resulting in decreases in the diameters. On further examination, however, at 16 weeks following surgery it was observed that the diameter of the whole femoral and partial tibial parts were <3 mm, which was the diameter of the Kirschner wire used in drilling. A difference of 0.4 mm between the observed and estimated diameter at the distal site in the femoral part may be attributed to bone incorporation from the surrounding area, as reported recently in the use of micro-CT scanning on ACLR in animal models (2,30). Inward growth decreased the graft bone interference distance in micro-scale assessment. An alternative explanation is required for the variance in diameters among proximal, middle and distal sites in both femoral and tibial parts. This is primarily due to the technique used in drilling the tunnel. The tunnel was initially drilled from the native ACL remnants within the knee joint towards femoral and tibial insertion sites. To feed the PET ligament through the tunnel, extra drilling was performed to expand the tunnel size at the femoral and tibial sites of the tunnel. SNR analysis provided another perspective in evaluating graft conditions following ACLR. The graft used in ACLR was set to present a relatively fixed signal intensity. The rise in SNR between 4 and 16 weeks following surgery in both femoral and tibial parts of the PET ligament graft indicated potential tissue growth on the surface or inside the ligament graft, which is consistent with the aforementioned decreases in diameters. Based on the hematoxylin-eosin staining results, it was demonstrated that the results were consistent with micro-MRI findings at 4, 8 and 16 weeks. The interface decreased over time, which was in accordance with micro-MRI findings. These findings suggest that micro-MRI is an effective tool and possible supplement to the histology, biomechanical study, gross evaluation, and micro-CT scans in the ACLR preclinical research field.

However, there are several limitations to the present study. First, a small sample size of 12 New Zealand white rabbits was investigated. Further studies with larger sample amounts are required for better analysis of micro-MRI. Second, due to the limitation of the coil size, in vivo micro-MRI scanning, which is accessible for rats and mice in preclinical studies, was not available in the current study, as reported above. With in vivo observation, a single knee joint may be continuously observed at different time points. Despite the short interval between euthanasia, harvesting and micro-MRI scanning, the in vitro micro-MRI scan may not fully reflect the true structural circumstance inside the knee joint. A more suitable MRI coil may solve this problem. Third, a further limitation is the imaging of the graft bone complex samples with micro-MRI. The T2-Turborare Sequence used in the present study may detect the ligament and cartilage lesion more easily than T1 sequences, but specific scan protocols, such as T2 mapping, fat saturated gradient echo, and 3D gradient echo, may be more feasible in subsequent research on ACLR preclinical studies (31,32). Furthermore, CT scanning or Micro-CT scanning would be helpful in tunnel diameter evaluation in our future study.

The present study applied micro-MRI scans in an animal model that underwent ACLR surgery with a PET ligament, and determined that micro-MRI is promising in quantitatively observing graft bone healing processes directly, with a focus on graft tunnel distances and the SNR.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions
FC and FW performed the experiment and wrote the manuscript. JJ conceived the idea and designed the study with SC. In addition, SC reviewed and rewrote the manuscript.

Ethics approval and consent to participate
The present study was reviewed and approved by the Animal Experiment Ethics Committee of the College of Pharmacy, Fudan University (Shanghai, China).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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