Tissue identification with micro-magnetic resonance imaging in a caprine spinal fusion model

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Abstract Nonunion is a major complication of spinal interbody fusion. Currently X-ray and computed tomography (CT) are used for evaluating the spinal fusion process. However, both imaging modalities have limitations in judgment of the early stages of this fusion process, as they only visualize mineralized bone. Magnetic resonance imaging (MRI) could be of great value as it is able to discriminate between different types of tissue. A feasibility study was performed in nine animals from a goat spinal fusion study, to evaluate the detection capacity of different tissues with micro-MRI. In this study bioresorbable poly-lactic acid cages were used. Six- and 12-months follow-up specimens were scanned in a 6.3 T micro-MRI scanner. After scanning, the specimens were processed for histology. Different types of tissue as well as the degradable cage material were identified in the fusion zone and designated as regions of interest (ROIs). Subsequently, the location of these ROIs was determined on the corresponding micro-MRI image, and average signal intensities of every individual ROI were measured. An excellent match was seen between the histological sections and micro-MRI images. The micro-MRI images showed quantifiable differences in signal intensity between bone with adipose marrow, bone with hematopoietic red marrow, fibrocartilage, fibrous tissue, and degradable implant material. In time the signal intensity of bone with adipose marrow, bone with hematopoietic red marrow, and of fibrous tissue remained relatively constant. On the other hand, the signal intensity of the degradable implant material and the fibrocartilage changed significantly in time, indicating change of structure and composition. In conclusion, in our model using bioresorbable cages the MRI provides us with detailed information about the early fusion process and may therefore, allow early diagnosis of non-union.

Keywords Spine · Interbody fusion · Bioresorbable implant · Micro-MRI · Histology

Introduction Spinal fusion essentially is a process of bone formation, which is influenced by a variety of biomechanical,
biochemical, cellular, and hormonal mechanisms [9]. As in fracture repair, mechanical stability is a prerequisite for successful bone formation at the spinal fusion site. Excessive motion of the segments may lead to complications like pseudoarthrosis or non-union with cartilage and/or fibrous tissue formation [2,4,21]. Non-union, in fact, is one of the most frequent complications of arthrodesis [28]. In the spine, however non-union is difficult to diagnose at an early stage, as half of the patients with a non-union are reported to be asymptomatic [22,26].

Plain radiographs are the most commonly used imaging technique to evaluate the fusion process. Trabecular bone, crossing at the fusion site on antero-posterior and lateral projections is used to assess consolidation of the bone graft [16]. However, plain radiography is accurate in only 59 to 82% of the cases with a high specificity and a low sensitivity [3,6,10,20,26]. Another disadvantage inherent to radiographic imaging techniques is the inability to visualize soft tissue or radiolucent materials that are used in cages [31]. When metallic implants are used, accurate assessment of the fusion zone inside the cage is impossible due to the radiopacity of metallic cages [29,30,32].

During the past several years, computed tomography (CT) has become one of the most important diagnostic modalities in the evaluation of spinal interbody fusion. CT can create 3D images and slices, demonstrating the presence or absence of bone bridging between the vertebral bodies [8,15]. However, none of the previously described disadvantages of X-ray evaluation are overcome by CT. Furthermore, CT comes at the costs of considerable radiation doses [24].

Recent studies show that MRI can detect changes in soft tissues over time [1,17–19,27]. Furthermore, processes inside the cage can be monitored with MRI, especially when bioresorbable cage material is used. Moreover, MRI provides information on the position of the cage after surgery and may show the time dependent degradation of bioresorbable spinal implants [11]. These advantages of MRI, combined with the absence of radiation exposure to the patient, could make MRI scanning a valuable diagnostic tool for monitoring the fusion process.

In the present study we analyzed the feasibility of micro-MRI analysis to study the process of fusion inside a bioresorbable PLDLA cage in a goat model and quantified these micro-MRI measurements on a scale. For this purpose, histological sections were analyzed and compared with corresponding micro-MRI images. The signal intensity of the different tissues in the MRI images was subsequently quantified and tissue specific differences were determined for developing a micro-MRI tissue grading scale.

Materials and methods

Animal model and study design

Micro-MRI images and histological slices were obtained from nine goats used in a previously described caprine spinal fusion model study [11,12]. In this study, the L3-L4 vertebral segments of the lumbar spine were fixated using a rectangular (10 × 10 × 18 mm) bioresorbable PLDLA cage, filled with impacted autologous bone graft. After creating a defect through both endplates, the cage was placed laterally between two vertebrae. The surgical procedure, cage, and implantation procedure have been described in detail elsewhere [11]. The follow-up was 6 (n = 3) and 12 months (n = 6).

Magnetic resonance imaging

At autopsy, the treated motion segment (L3-L4) was excised and trimmed of residual musculature. The transverse and spinous processes were removed. The segment was kept at 4°C and immediately transported to an MR unit.

The MR imaging experiments were performed using a 6.3 T MR imaging scanner with a 9.5 cm diameter horizontal bore, equipped with a Varian VX-R-S imaging console (Varian Associates, Palo Alto, CA). The MR imaging studies were started approximately 3 h after sacrificing the animals. The segments were wrapped in plastic and inserted into a 5.5 cm diameter, linear driven birdcage radio frequency coil. The experiments were performed at room temperature. Depending on the size of the specimen, 21 to 35 slices were recorded; the slice thickness was 1 mm, there was no gap between them. The in-plane field of view for all images was 5.5 × 5.5 cm². High resolution MR images (proton density) were acquired (transverse and sagittal plane) using a standard spin echo sequence with a repetition time of 4 s, an echo time of 15 ms, and 12 signal averages. The matrix size was 256 × 256, zero-filled to 512 × 512, yielding an in-plane resolution of 200 and 100 μm in the original and interpolated images, respectively (Fig. 1b).

Histology

After MR imaging, each segment was sectioned sagitally in a standardized manner, using a water-cooled band saw (EXAKT, Norderstedt, Germany), creating a 5 mm par- amidsagittal section. This section was dehydrated using formaldehyde, embedded in methyl methacrylate and 12 μm sections were obtained. All sections were stained with Goldner Trichrome (GT) and Toluidine Blue (TB) (Fig. 1a).
Four different tissues generally present at a fusion site were defined prior to the histological analysis: bone containing hematopoietic marrow (BHM), bone containing adipose marrow (BAM), fibrous tissue (F) and fibrocartilage (FC). Cage material was identified by the absence of cells. The definitions of the tissues used in the present study are shown in Table 1. Regions of interest (ROIs), each containing only one of the four selected tissues or the cage material, were selected by one of the authors (MPU). Subsequently a second observer (VE) classified the tissue in each ROI on every histological section. When both observers independently agreed on the nature of the tissue in a ROI, it was included in the study. This resulted in tissue specific ROIs in all of the histological sections for all four different tissues as well as cage material.

Magnetic resonance imaging analysis and assessment

For each histological section, the corresponding micro-MRI image was selected using ImageJ, a public domain, Java-based image processing program (National Institutes of Health, Bethesda, MD, USA). Subsequently, the previously identified ROIs were identified in the MRI images (Fig. 1b). The signal intensity of each ROI was determined using ImageJ. Considerable variation in the absolute value of the signal intensities was observed. Therefore, fibrous tissue was selected as baseline because the signal intensity of this tissue was stable and remained stable during the follow-up period. The signals of all tissues were expressed as a ratio of fibrous tissue.

Statistical analysis

Statistical analysis software was used to analyze the data (Instat. Graph pad Software Inc, San Diego, USA). Repeated measures ANOVA with a Tukey–Kramer post-hoc test was used to determine differences between tissue signal intensities. The Student \( t \) test was used to analyze differences between both time-points.

Results

Histology

The selected tissues were identified in all specimens by both observers. The cages were clearly detectable in all specimens. After 6 months, the cages in general showed considerable deformation and fracturing. After 12 months, all six cages had disintegrated, migrated and were deformed (Fig. 1b). At both time points, all cages were surrounded by a thin layer of fibrous tissue, gradually changing into fibrocartilage in the fusion zone and at the load bearing edges of the cage.

The relative signal intensities and their relation to different tissues are shown in Fig. 2. The different tissues

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Table 1  Abbreviations and definitions used for analyzed histological tissues

| Tissue                                | Abbreviation | Definition                                      |
|---------------------------------------|--------------|-------------------------------------------------|
| Bone with hematopoietic marrow       | BHM          | Presence of hematopoietic cells and osteocytes* |
| Bone with adipose marrow             | BAM          | Presence of adipose cells and osteocytes*        |
| Fibrous tissue                       | F            | Presence of fibroblasts and collagenous fibers  |
| Fibrocartilage                       | FC           | Presence of (rows of) chondrocytes              |
| Cage material                        | C            | Absence of cells                                |

* Bone defined as tissue containing osteocytes, hematopoietic marrow defined as tissue containing hematopoietic cells, and adipose marrow defined as tissue containing adipose cells.
In the present study the location of the two types of bone marrow was not determined. However, in all sections (both 6 and 12 months) adipose marrow was observed closest to the fusion zone. Both cranially and caudally of the fusion zone, hematopoietic marrow was seen intruding the fusion zone subsequent to adipose marrow.

### Discussion

Rates of spinal fusion procedures are increasing in recent years in the US [7, 32]. The reported rates of failed union in general are high, varying between 0 and 56% depending on, among others, fixation type and number of levels fused [5, 16]. However, not all patients with a non-union are symptomatic, complicating the process of diagnosis of non-union [22]. At this time, the golden standard in diagnosing non-union is surgical exploration [16]. Ideally, a non-invasive tool should be available for assessing the process of fusion in the patient, and predicting outcome of the process at an early stage. Currently, X-ray and CT are clinically used to evaluate spinal fusion. As the fusion process includes different stages with different sorts of soft tissues, more sophisticated imaging techniques are needed to discern the differences between them. We used samples from a spinal fusion goat study to evaluate the micro-MRI signal intensities of tissues involved in interbody fusion. In this study bioresorbable polyactic acid cages were used. We found that the ratios of the tissues analyzed were reproducible and distinguishable from each other using micro-MRI (Fig. 2). Furthermore, we observed that the signal intensity of fibrocartilage and degradable cage material changed during follow-up. We conclude that MRI provides us with vital and detailed information about the early fusion process and may therefore, allow early diagnosis of non-union. At present the use of MRI for this purpose is limited to non-metallic cages.

Fibrocartilage and cage material signal intensities changed significantly in the goat segments between 6 and 12 months (Fig. 2). The lower signal intensity of the fibrocartilage at 12 months indicates a decrease in the content of water. This may be explained by the ongoing deposition of extracellular matrix and maturation of the fibrocartilage. Clear histological changes, however, were not observed in this tissue. By contrast, the signal intensity of the cage material increased during the follow-up (Fig. 2). After 6 months the cages produced low signal intensity (Fig. 1b). After 12 months, however, the signal intensities increased significantly. This change can be explained by the degradation of the cages, which actually occurs through hydrolysis after water penetrates the polymer. This observation is of interest as it was shown before that these PLDLA cages retain their initial strength for

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**Table 2** P-values of differences between different types of tissue and cage material at 6 months (vertical, bold italic), 12 months (horizontal, italic) and between follow-up time-points within selected tissues (bold)

|          | 12 months | 6 months |
|----------|-----------|----------|
|          | F BAM BHM FC C |
| F        | <0.001 | <0.001 | <0.05 | <0.001 |
| BAM      | <0.001 | NS     | NS    | NS     | <0.01 |
| BHM      | <0.001 | <0.001 | NS    | <0.05  | <0.05 |
| FC       | <0.001 | <0.001 | <0.05 | <0.001 | <0.001 |
| C        | <0.001 | <0.01  | NS    | <0.001 | <0.001 |

NS = not significant.

showed significantly different signal intensities as compared to each other, with three exceptions (Table 2): after 6 months; bone with adipose marrow (BAM) versus bone with hematopoietic marrow (BHM), and bone with adipose marrow versus fibrocartilage (FC); and after 12 months, bone with hematopoietic marrow versus cage material. Comparing the signal intensities of BAM and BHM in time did not result in significant changes. The signal intensity of fibrocartilage was higher after 6 months than intensities of BAM, BHM, and cage material (Fig. 2). However, after 12 months the fibrocartilage signal intensity had decreased significantly ($p = 0.0002$) and was significantly lower compared to all other tissues, including the degradable cage material (Fig. 2).

After 6 months, ROIs classified as bioresorbable PLDLA cage were showing the lowest signal intensities (Fig. 2). At 12 months, the cage material showed more deformation and significant higher signal intensities compared to 6 months ($p = 0.0025$).
approximately 6 months [25, 33]. After 6 months their strength gradually decreases and at 12 months the cages entirely lost their mechanical integrity [25]. In a clinical perspective, the signal intensity of the cage can be used as an indicator of its mechanical integrity; loss of mechanical integrity contributes to instability of the spinal segment, which is a risk factor for non-union.

In the present study, in both 6 and 12 months, adipose marrow was observed closest to the fusion zone. Both cranially and caudally of the fusion zone hematopoietic marrow was seen intruding the fusion zone after adipose marrow. This is in line with previous studies in which red (hematopoietic) marrow was found in later stages, indicating that red marrow is related to a more mature bone type [11, 30]. This is an interesting finding, because adipose marrow is generally found in older persons, whereas, hematopoietic marrow is rather related to bone formation and growth. This study consistently shows a role for adipose bone marrow in the fusion process. The ability to discriminate between red and yellow marrow with MRI also has practical implications, as Lang et al. [14] found that the signal intensity of bone marrow might be related to functional stability of a fused segment.

With respect to future clinical application two methodological aspects of this study need to be addressed. First, the use of a high field MR scanner (6.3 T) and second the use of an internal reference (fibrous tissue) for signal intensities. The high-field MR scanner was used to obtain high-resolution MR images, which resulted in images of very high quality. Because of its small bore size the present scanner is obviously not suitable in a clinical setting. Furthermore, the current scanning times are too long for clinical application. However, high resolution imaging of bone and cartilage becomes readily available in a clinical setting, with the development of high (3 and 7 T) MR scanners and the development of new and faster MR sequences [23]. The signal intensity of fibrous tissue was used as reference because the absolute pixel intensities in the images varied between MRI measurements. Although the use of an external “standard”, e.g., a tube containing water, is preferable, the stable proportions of the different tissues justified the use of this internal “standard”. In future (pre) clinical research an external standard will be used.

In the present model a standardized endplate perforation was used. Polylactic acid implants degrade over time mostly by bulk hydrolysis which produces lactic acid. Transportation of this waste product is essential since accumulation will lead to autocatalysis of the implant and bone resorption. To our opinion, opening of the endplates is not only essential for removal of lactic acid but also for the supply of osteogenic cells. A disadvantage of endplate perforation could be subsidence of the segment. A recent RSA study in the same model has shown that both biodegradable and the titanium control cages induce less than 1.5 mm of segment subsidence after 12 months [13], which is largely or completely due to settling of the cage. Complete perforation of the endplates may not be necessary and could lead to subsidence in combination with stiff and sharp implants. However, we feel that endplate preparation with formation of bleeding subchondral bone is vital when using polylactic acid interbody fusion devices.

In contrast to CT and X-ray, micro-MRI can differentiate between different kinds of soft tissue as well as bone and implant material in our model, using non-metallic cages. The present study demonstrates that there is a clear correlation between micro-MRI signal intensities and the corresponding histological tissues. Our data further show that micro-MRI could be used to analyze the process involved in spinal fusion on tissue level. Moreover, micro-MRI can be used to monitor the process of degradation of biodegradable cage material. MRI could play a major role in preclinical development of biodegradable and other non-metallic implants. Furthermore, the increasing availability of clinical high resolution MR scanners supports a clinical application of this method in the near future. In conclusion, micro-MRI could be a valuable tool to monitor and evaluate early bone healing for both research purposes and future clinical decision making.

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