Detection of early cartilage degeneration in the tibiotalar joint using 3 T gagCEST imaging: a feasibility study

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Abstract
Objective To establish and optimize a stable 3 Tesla (T) glycosaminoglycan chemical exchange saturation transfer (gagCEST) imaging protocol for assessing the articular cartilage of the tibiotalar joint in healthy volunteers and patients after a sustained injury to the ankle.

Methods Using Bloch–McConnell simulations, we optimized the sequence protocol for a 3 T MRI scanner for maximum gagCEST effect size within a clinically feasible time frame of less than 07:30 min. This protocol was then used to analyze the gagCEST effect of the articular cartilage of the tibiotalar joint of 17 healthy volunteers and five patients with osteochondral lesions of the talus following ankle trauma. Reproducibility was tested with the intraclass correlation coefficient.

Results The mean magnetization transfer ratio asymmetry (MTR asym), i.e., the gagCEST effect size, was significantly lower in patients than in healthy volunteers (0.34 ± 1.9% vs. 1.49 ± 0.11%; \( p < 0.001 \) [linear mixed model]). Intra- and inter-rater reproducibility was excellent with an average measure intraclass correlation coefficient (ICC) of 0.97 and a single measure ICC of 0.91 (\( p < 0.01 \)).

Discussion In this feasibility study, pre-morphological tibiotalar joint cartilage damage was quantitatively assessable on the basis of the optimized 3 T gagCEST imaging protocol that allowed stable quantification gagCEST effect sizes across a wide range of health and disease in clinically feasible acquisition times.

Keywords Cartilage · Magnetic resonance imaging · Proteoglycans · Osteoarthritis · Molecular imaging

Introduction
To this day and age, several magnetic resonance imaging (MRI) techniques have emerged that go beyond mere morphological depiction of joint cartilage. Such compositional MRI techniques allow the detection of early degenerative changes of the articular cartilage, e.g., loss of proteoglycans, that precede morphological damage and hence are considered an early, and more importantly, reversible, stage of osteoarthritis (OA) [1]. Because of its proteoglycan-specificity, the gold-standard technique of compositional MRI of cartilage is delayed gadolinium-enhanced MRI of cartilage (dGEMRIC) [2, 3]. However, due to recent restrictions imposed on gadolinium-based contrast agents, alternative compositional MRI techniques that do not rely on the administration of contrast agents have received ever-increasing scientific and clinical attention [4]. Among these techniques, glycosaminoglycan chemical
exchange saturation transfer (gagCEST) imaging assesses the specific GAG content in human articular cartilage and its depletion, which is considered an early sign of cartilage degeneration [5].

GagCEST imaging is based upon the chemical exchange of water protons between GAG and bulk water molecules. To induce a CEST effect, solute protons are saturated by a frequency-specific radiofrequency (RF) pulse and then transferred to bulk water by chemical exchange, which consequently reduces its signal. The normalized signal can then be used to quantify the CEST effect at a GAG-specific frequency range of 0.9–1.9 ppm via analysis of the magnetization transfer ratio asymmetry (MTR asym), which correlates with the GAG concentration [5, 6]. For additional details on the basic principles of CEST imaging, the interested reader is referred to earlier excellent reviews [7, 8]. Several studies showed promising results using gagCEST imaging at the spine [9–12]. However, data on the joints of the lower extremity with substantially thinner cartilage are sparse. In 2016, our group demonstrated promising results for the application of gagCEST at the knee joint [13]. Kogan et al. applied gagCEST imaging on a 7 T MRI scanner to assess the ankle joint of healthy volunteers [14]. Even though these results were promising, gagCEST imaging of the ankle joint has not yet been established on a 3 T MRI scanner. To achieve a more widespread scientific and clinical adaptation of the technique, the clinical utility has to be demonstrated on a broader scale, which -given the limited availability of 7 T MRI scanners- necessitates the technique’s implementation on more widely available 3 T MRI scanners.

Tibiotalar joint injuries are common [15]. Osteochondral lesions of the talus (OLT), defined as an injury of the cartilage layer and the underlying subchondral bone, are frequent injuries in active populations that can be seen in up to 73% of all traumatic ankle injuries [16]. OLTs may predispose the joint to premature OA and ought to be diagnosed in an early and reliable manner as a timely diagnosis is a pre-requisite for appropriate treatment [17].

The aim of this study was (a) to develop and optimize a gagCEST imaging protocol for the articular cartilage of the tibiotalar joint that is clinically feasible and fits into diagnostic workflows and (b) to apply this imaging protocol to a population of healthy volunteers and patients with OLT after an ankle injury to prove clinical utility and validity. We hypothesized that -based on the developed and optimized gagCEST imaging protocol- (a) imaging of the articular cartilage of the tibiotalar joint would be possible in a clinical population and in clinically feasible time frames and (b) patients after variable ankle injuries (representative of the patient population undergoing MRI diagnostics in the clinic) demonstrate lower gagCEST effects compared to healthy volunteers.

### Methods

#### Simulations

In a first step, simulations using the two-pool (water and GAG (–OH and –NH) Bloch–McConnell equation [18, 19] and a customized script (implemented in MATLAB R2018a, The MathWorks, MA, USA) and to be downloaded at https://github.com/cest-sources/BM_sim_fit/) [20] were applied for the optimization of a pulsed gagCEST sequence [20–22]. The equations were solved analytically [19]. Based on this script, the CEST effect was simulated without the application of a saturation pulse. The radiofrequency field strength B1, the pulse duration t p and the number of CEST saturation pulses n p were varied using a constant duty cycle (DC) of 0.5. To keep the specific absorption rate (SAR) within the safe range, local SAR was restricted accordingly. Therefore, the maximum pulse duration was secondarily restricted by their scanner to a maximum of 300 ms. For water, simulations were performed with relaxation times as reported earlier, i.e., T1 = 1.2 s and T2 = 0.039 s and a concentration of 88 M [23, 24]. The following parameters were used for GAG-OH protons: exchange rate = 1000 Hz, concentration 0.3 M, T1 = 1 s, T2 = 0.01 s and chemical shift = 1 ppm, and for GAG-NH protons: exchange rate = 50 Hz, concentration = 0.1 M, T1 = 1 s, T2 = 0.01 s and chemical shift = 1 ppm [24, 25]. The different variations of the parameters used in the simulation are displayed in Table 1; output parameters were z-spectra and MTR asym curves. For each parameter, the maximum MTR asym value was analytically determined at a step size of 0.01, 0.02 and 0.05 ppm at frequency offsets of 0.9–1.9 ppm, 0.5–1.5 ppm and 1–1.5 ppm. The optimized protocol in terms of the largest gagCEST effect at a reasonable acquisition time was used for the subsequent in-vivo studies.

| Experiment | n p | t p (ms) | B1 [µT] |
|------------|-----|----------|----------|
| 1          | 6   | 100      | 0.2; 0.4; 0.6; 0.8; 1.0; 1.2; 1.4 |
| 2          | 6   | 100; 200; 300 | 1.0 |
| 3          | 2; 4; 6; 8; 10; 12; 14 | 100 | 1.0 |

In each experiment, one of the three parameters (number of pulses n p, pulse duration t p, and radiofrequency-field strength B1) was systematically varied.

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[1-12] Citing sources...
In-vivo study

Study population

19 healthy volunteers (mean age 23.0 ± 3.8, range 20–37 years, 11 males, 8 females) and six patients (mean age 31.7 ± 9.3, range 20–44 years, two males, four females) after earlier ankle injury were recruited from 06/2018 to 01/2019 via dedicated specialist consultations at our Department of Orthopedic and Trauma Surgery. The predefined inclusion criterion for patients was an isolated traumatic OLT lesion as diagnosed in earlier MRI studies. At the time of recruitment, patients were graded according to the Anderson modification of the Berndt and Harty classification and four patients had grade 1 and two patients grade 2b OLT lesions [15, 16]. Predefined exclusion criteria for healthy volunteers included all forms of primary or secondary OA of the ankle as well as other bone and joint disorders such as OLT, rheumatoid arthritis, avascular necrosis, gouty arthritis, septic arthritis, Paget disease or osteochondritis dissecans. Volunteers were also excluded if they had acute or chronic ankle pain or a history of serious trauma or surgery to the index ankle joint.

The MRI data sets of one patient and two healthy volunteers had to be excluded from image analysis due to excessive motion artifacts. The mean disease duration of patients was 22 ± 30 months (range 1–60 months). Written and informed consent was obtained from all patients before the initiation of the study. The study was approved by the local ethics committee (Ethical Committee of the University Hospital Düsseldorf, study number: 3980).

MRI studies

All imaging studies were performed on a 3 T MRI scanner (Magnetom Prisma, Siemens Healthineers, Erlangen, Germany) using a dedicated receive-only 16-channel foot–ankle coil (Foot/Ankle 16, Siemens Healthineers). Patients and volunteers were scanned in the supine position with a neutral ankle position of 90° dorsiflexion. Positioning aids, sandbags and medical tape were used to reduce motion artifacts.

The MRI protocol included standard morphological sequences, i.e., sagittal (sag) and coronal (cor) Proton Density-weighted (PDw) fat-saturated (fs) sequences, transversal (tra) T2-weighted turbo-spin echo (TSE), and cor T1-weighted TSE sequences. In addition to the actual gagCEST sequence as detailed below, water saturation shift referencing (WASSR), T1 mapping gradient echo (GE) and T2 multi-spin- echo (SE) mapping sequences with five different echo times (13.8, 27.6, 41.4, 55.2 and 69 ms) were acquired. Of note, the latter two sequences were only acquired in the healthy volunteers and not in the patients. GagCEST imaging was performed using a two-dimensional (2D) radiofrequency (RF)-spoiled GE sequence with a pulsed CEST pre-saturation module consisting of 8 Gaussian-shaped RF pulses with a duty cycle of 0.5. 26 images with pre-saturation pulses at different offset frequencies around the bulk water resonance were obtained. Among these images was one reference image with a frequency offset of 300 ppmThe maximum frequency offset (Δω) was 4 ppm with a step size of 0.33 ppm. In a fraction of the healthy volunteer cohort (n = 10, mean age 22.4 ± 1.8, range 20–25 years, seven males, three females) radiofrequency field strengths and pulse durations were systematically varied to optimize the protocol at the beginning of the study. More specifically, three different radiofrequency field strengths (B1 = 0.6, 0.8 and 1.0 µT) and three different pulse durations (τp = 100, 200 and 300 ms) were used. Based on the results of the simulations, i.e., the largest measured MTRasym values, we used a radiofrequency field strength of B1 = 0.8 and a pulse duration τp = 300 ms in the remaining healthy volunteer and patient cohorts. For the WASSR sequence, 22 images with pre-saturation and a reduced radiofrequency field strength (B1 = 0.25 µT) were obtained. The maximum frequency offset was decreased to Δω = 1 ppm with a step size of 0.1 ppm. For WASSR and CEST sequences, motion correction was applied. The acquisition time was 5:01 min for the CEST and 2:22 min for the WASSR sequence. The total acquisition times for the compositional MRI sequences were: 24:21 min for the initial 10 healthy volunteers (3 × 5:05 min CEST, 1 × 2:22 min WASSR, 6 × 1:14 min T1) and 7:27 min for the remaining 7 healthy volunteers and the 5 patients (1 × 5:05 min CEST and 1 × 2:22 min WASSR). The acquisition time for the morphological sequences was 18 min, resulting in a total scan time of 42:21 min for the initial 10 volunteers and 25:27 min for the consecutive 7 volunteers and the 5 patients.

Detailed parameters of the morphological and compositional sequences are given in Tables 2 and 3.

Image analysis

All images were independently analyzed by two radiologists (DBA, 3 years of training in musculoskeletal imaging; CS, 8 years of training in musculoskeletal imaging) who were blinded to the volunteers’ or patients’ data. First, all studies were read to determine the individual joint’s overall status with a particular focus on the integrity of tibiotalar cartilage. Also, OLTs were -if present- classified according to Hepple et al. [26]. Second, using the unsaturated WASSR image, both readers independently identified the cartilage layers of the tibiotalar joint and quantified its biophysical properties in a standardized manner by placing an ellipsoid-shaped region-of-interest (ROI) in the median plane onto both cartilage layers at the central load-bearing region of the tibiotalar joint. Each ROI was placed distant to the tibial and talar bone cortex and the anterior and posterior joint.
areas to reduce partial volume artifacts due to the presence of cortical bone and potentially excessive amounts of joint fluid (Fig. 1). The second reader repeated the ROI placement at a different time point to allow for the assessment of inter-rater reliability.

For the analysis of the MTR asym curve, i.e., the CEST effect, we used an in-house script implemented in Matlab (MATLAB R2018a, The MathWorks, Inc., MA, USA). Prior to further evaluation, $B_0$ field inhomogeneities were corrected by the WASSR maximum-symmetry algorithm with the calculation of a pixel-wise frequency offset curve [27, 28]. These offset-corrected CEST-curves divided by the signal without pre-saturation ($S_0$) were defined as the so-called z-spectrum ($Z(\omega)$). The maximum frequency offset of each z-spectrum was $\Delta \omega = 3$ ppm. Next, we used the magnetization transfer asymmetry (MTR asym) (defined as $\text{MTR}_{\text{asym}}(\Delta \omega) = Z(-\Delta \omega) - Z(\Delta \omega)$) for the evaluation of the gagCEST effect [29]. MTR asym maps were calculated using the average value of MTR asym in the GAG-specific range of $\Delta \omega = 0.9 - 1.9$ ppm [30]. In addition, the $B_0$-corrected and -normalized spectra were fitted using Lorentzian function analysis to account for the GAG-OH, GAG-NH, water pools at $-1$ ppm, the nuclear Overhauser effect at $-1$ and $-2.8$ ppm and the magnetization transfer pool at $-2.43$ ppm [31, 32]. In the following, the Lorentzian-fitted gagCEST effect is given as GAG-OH amplitude.

T1 and T2 relaxation times calculations in ten healthy volunteers were also performed in Matlab. In a pixel-wise manner, acquired data was fitted and calculated based on the following equations:

Table 2  Detailed sequence parameters of morphological MRI sequences

| Imaging parameter | Sagittal fs PDw | Coronal fs PDw | Transversal T2w TSE | Coronal T1w TSE |
|-------------------|----------------|----------------|---------------------|----------------|
| FOV (mm)          | 160×160        | 160×160        | 160×160             | 160×160        |
| Slice thickness (mm) | 3             | 3              | 3                   | 3              |
| TE (ms)           | 40             | 40             | 78                  | 17             |
| TR (ms)           | 4000           | 4000           | 4600                | 700            |
| Resolution (mm/pixel) | 0.31×0.42     | 0.31×0.42      | 0.31×0.39           | 0.28×0.4       |
| Flip angle (°)    | 150            | 150            | 150                 | 140            |
| Acquisition matrix | 512×384        | 512×384        | 512×410             | 576×403        |

Field of view (FOV), slice thickness, echo time (TE), repetition time (TR), resolution, flip angle, and acquisition matrix are given for sagittal and coronal fat-saturated proton-density-weighted (fs PDw), transversal T2-weighted turbo spin echo (T2w TSE) and coronal T1-weighted TSE (T1w TSE) sequences.

Table 3  Detailed sequence parameters of compositional MRI sequences

| Imaging parameter | WASSR | gagCEST | T1 map | T2 map |
|-------------------|-------|---------|--------|--------|
| FOV (mm)          | 160×160 | 160×160 | 160×160 | 160×160 |
| Slice thickness (mm) | 5      | 5       | 7      | 3 mm   |
| TE (ms)           | 3.5   | 3.5     | 11     | 13.8/27.6, 41.4/55.2/69 |
| TR (ms)           | 7.2   | 7.2     | 6000   | 1000   |
| TI (ms)           | 25/50/100/500/1000/2000 | 0.6×0.6 | 0.6×0.6 | 0.6×0.6 |
| Resolution (mm/pixel) | 0.6×0.6 | 0.6×0.6 | 0.6×0.6 | 0.4×0.4 |
| Flip angle (°)    | 15    | 15      | 180    | 180    |
| Pulsed CEST saturation module | – 1 to 3 | – 3 to 3 | – 1 to 3 | – 3 to 3 |
| Frequency range (ppm – ppm) | 21+1 | 25+1 | 25+1 reference image |
| Number of Dynamic Scans | 1 | 8 | 8 |
| Number of saturation pulses | 54 | 300 (100, 200) | 300 |
| Pulse Duration $t_p$ (ms) | 0.2 | 0.8 (0.6, 1.0) |

In healthy volunteers, pulse duration $t_p$ and $B_1$ amplitude were evaluated at 100, 200, and 300 ms and at 0.6, 0.8, and 1.0 µT, respectively, while in patients, the following parameter settings were used: 300 ms and 0.8 µT.

WASSR water saturation, gagCEST glycosaminoglycan chemical exchange saturation transfer imaging, FOV field of view, TE echo time, TR repetition time, TI inversion time.
with $T_1$ and $T_2$ being the sought relaxation times, $M_z(t)$ the total magnetization in the $z$-direction, and $M_{xy}(t)$ the total magnetization in the $xy$-plane at time point $t$.

Statistical analysis

SPSS software (IBM, version 22, Armonk, NY, USA) was used for all statistical analyses by KLR and DBA. For descriptive analysis, mean gagCEST values ± standard deviation, median, and range (minimum–maximum) were calculated for healthy volunteers and patients. For optimization of the imaging protocol radiofrequency field strength and pulse duration were systematically varied and then compared using a multivariate analysis of variance (MANOVA) and a post-hoc Scheffé-test. For the comparison of gagCEST values between both cohorts, a multivariable statistical analysis was performed using a linear mixed model (LMM). The established model included a subject-specific random intercept, the factors healthy volunteer/patient, age, gender and the interaction of these factors assuming a fixed linear effect on the gagCEST values. Results of this model are given in Table 1 of the Supplementary Material. The LMM was fitted using a restricted maximum likelihood approach (REML). Based on this final model, the mean differences of gagCEST values were calculated and evaluated for significance. For correlation analyses of MTRasym values and GAG-OHAmplitudes, Pearson’s correlation was determined and quantified using the correlation coefficient $r$. Correlation strength was graded as suggested by Cohen [33]: small (0.1–0.3), moderate (0.3–0.5), and large (> 0.5). $p$ values < 0.05 were considered significant. For the evaluation of inter- and intrainter reliability, single and average measure intraclass correlation coefficients (sICC and aICC) were calculated based on the ROIs drawn by the two raters.

Results

Simulations

The results of the systematic simulations are illustrated in Figs. 2 and 3.

a. Variation of $t_p$.

Maximum MTRasym values were 1.33 % at 0.9–1.9 ppm with $t_p = 200$ ms, 1.07 % at 0.5–1.5 ppm with $t_p = 100$ ms and 1.17 % at 1.0–1.5 ppm with $t_p = 100$ ms (Fig. 3a).

b. Variation of $n_p$.

The CEST effect increases with the number of applied saturation pulses ($n_p$) (Fig. 3b). Eight applied pulses reach 98% of the maximum gagCEST effect that could be obtained with 14 pulses at all ranges (0.9–1.9 ppm, 0.5–1.5 ppm and 1.0–1.5 ppm). Maximum MTRasym values with eight applied pulses were 1.33 % at 0.9–1.9 ppm, 1.02 % at 0.5–1.5 ppm and 1.33 % at 1.0–1.5 ppm.

c. Variation of $B_1$.

The CEST effect increases with increasing $B_1$ until it reaches a maximum (Fig. 3c). Due to the spillover effect, MTRasym values decrease beyond the maximum. Maximum MTRasym values were 1.33 % at 0.9–1.9 ppm and a $B_1$ of 1 $\mu$T, 1.17 % at 0.5–1.5 ppm and a $B_1$ of 0.8 $\mu$T and 1.37 % at 1.0–1.5 ppm and a $B_1$ of 1 $\mu$T.

In-vivo studies

Morphological MRI of patients and healthy volunteers

Apart from the presence of OLTs as outlined below and a moderate joint effusion, the overall joint status of three of
Fig. 2 Simulations results detailing the effects of variations in CEST framework conditions. Pulse durations $t_p$ (100, 200, and 300 ms) (a), number of applied pulses $n_p$ ($n = 2$ – 14) (b), and radiofrequency field strengths $B_1$ (0.2–3 μT) (c) were systematically varied. Each colored curve represents a simulated parameter value and gives the $z$-spectrum at different offset frequencies (0–4 ppm).

Fig. 3 Simulations results detailing the effects of variations in CEST framework conditions. Pulse durations $t_p$ (100, 200, and 300 ms) (a), number of applied pulses $n_p$ ($n = 2$ – 14) (b), and radiofrequency field strengths $B_1$ (0.2–1.4 μT) (c) were systematically varied. Each colored curve represents a simulated parameter value and gives the maximum magnetization transfer ratio asymmetry at different offset frequencies (0–4 ppm).
five patients was unremarkable. In them, we did not find any bone marrow lesions, subchondral thickening, osteophytes or joint space narrowing. In two patients, we noted signs of secondary OA with osteophytes, joint space narrowing, subchondral sclerosis, and moderate joint effusion. The joint status of healthy volunteers was unremarkable without any structural alterations. Within the entire study population, the following accessory ossicles were found: Os tibiale externum in six individuals, Os trigonum in three individuals, Os supralaterale in one individual.

Staging of OLTs was performed according to the Heppner classification (stages 1–5, 1: articular cartilage damage only, 2a: cartilage injury with underlying fracture and surrounding edema, 2b: 2a without surrounding edema, 3: detached, but undisplaced fragment, 4: detached and displaced fragment, 5: subchondral cyst). The following stages were observed in the patient cohort: one individual with stage 2a, one individual with stage 3, one individual with stage 4 and two individuals with stage 5.

Implementation of the optimized protocol in 10 healthy volunteers

Table 4 gives the details of the MTRasym values in 10 healthy volunteers as a function of systematically varied parameter settings of $B_1$ (0.6, 0.8, and 1.0 µT) and $t_p$ (100, 200, and 300 ms).

| Offset frequency [ppm] | $B_1$ (µT) | $t_p$ (ms) | MTRasym (%) | $p$ value |
|------------------------|------------|------------|-------------|-----------|
| 0.9–1.9                | 0.6        | 100        | 0.37 ± 0.78 |           |
|                        | 200        | 0.75 ± 0.65|            |           |
|                        | 300        | 1.34 ± 1.05|            |           |
| 0.8                    | 100        | 0.12 ± 0.47|            |           |
|                        | 200        | 0.71 ± 0.81|            |           |
|                        | 300        | 1.67 ± 1.35|            |           |
| 1.0                    | 100        | 0.27 ± 0.78|            |           |
|                        | 200        | 0.94 ± 1.02|            |           |
|                        | 300        | 0.49 ± 0.95|            |           |

MTRasym values are given as mean ± standard deviation

Means were compared using a multivariate analysis of variance (MANOVA) followed by a post-hoc Scheffé test

$p$ values < 0.05 were considered significant and are given in bold type
Discussion

The most important finding of this study is that following comprehensive and systematic sequence optimization-gagCEST imaging of the tibiotalar joint is feasible using a clinical standard 3 T MRI scanner, fits into clinical workflows with an acquisition time of less than 07:30 min, and yields stable and reproducible results that allow compositional cartilage assessment. In addition, we demonstrated that the tibiotalar joint cartilage of patients with known tibiotalar joint injury, especially OLT, have significantly lower gagCEST values than healthy volunteers.

Compositional MRI exceeds the mere morphological depiction of cartilage and allows for the detection of early cartilage changes that precede morphological alterations, i.e., loss of proteoglycans, as an early, potentially treatable stage of OA. GagCEST can be used for the detection and treatment monitoring of very early OA [34]. Despite this great clinical need, research on gagCEST imaging in general has been limited by the numerous technical complexities involved such as homogeneous magnetic field properties, long scan times, low SNR, and high field strengths (optimally ≥ 7.0 T) that are considered necessary for cartilage imaging [35]. Moreover, with the majority of imaging studies focusing on the knee joint, data on the tibiotalar joint is sparse [14]. This is mainly due to the joint’s limited cartilage thickness, measuring only about 2 mm in healthy individuals and the known limited spatial resolution of gagCEST imaging [36, 37]. In this study, we set out to establish and optimize a gagCEST imaging protocol with reasonable scan times, sufficient SNR, and high reproducibility at 3.0 T for the potential implementation in the clinical setting.

GagCEST imaging can be modified by altering the number of applied saturation pulses, pulse durations and radiofrequency field strengths. To find the optimal setting of these parameters that allow for both a high gagCEST effect size and reasonable acquisition time, we used the Bloch–McConnell simulation before proceeding with the in-vivo measurements [38]. The simulation experiments showed a maximum effect size at a radiofrequency field strength of 0.8 µT.
The effect size decreased at higher field strengths due to the ‘spillover effect’: With an increasing $B_1$ amplitude, the spillover effect leads to direct saturation of the water pool instead of the soluble proton pool and hence results in decreases of the gagCEST effect [39]. When tested in healthy volunteers, we noted numerically higher $\text{MTR}_{\text{asym}}$ values and GAG-OH amplitudes at 0.8 than at 1.0 $\mu$T, but not than at 0.6 $\mu$T. The effect size increased with the applied number of pulses with a $\text{MTR}_{\text{asym}}$ of 0.98% at 14 pulses; however, at eight applied pulses, the $\text{MTR}_{\text{asym}}$ reached 0.98% of the maximum effect size. To keep the acquisition time as short as possible at a maximum gagCEST effect size, we decided to use eight pulses. Moreover, the effect size was found to be increased with increasing pulse durations. Due to limitations secondary to the specific absorption rate (SAR); however, the maximum pulse durations to be used in vivo were limited to 300 ms [40]. By trend, we found higher $\text{MTR}_{\text{asym}}$ values in vivo at a pulse duration of 300 ms (as compared to 100 and 200 ms)—even though these differences were only partially significant.

After simulations and in-vivo experiments, our final gagCEST protocol consisted of 8 applied pulses with a pulse duration of 300 ms at a radiofrequency field strength of 0.8 $\mu$T and a constant duty cycle of 0.5 aiming for a minimized scan time. We used WASSR to improve the differentiation of the water and GAG peak as well as to correct for $B_0$ field inhomogeneities [27]. Using this protocol, we found excellent reproducibility of gagCEST values as measured by one individual rater and between two independent raters ($\text{aICC} = 0.97$ and $\text{sICC} = 0.91$). These values for reproducibility were even higher than presented in previous studies focusing on gagCEST of peripheral joints [34]. A good reproducibility is beneficial not only for future studies, but also for the perspective of clinical implementation of the technique [41].

The acquisition time of the optimized gagCEST sequence was 5:01 min, followed by an additional 2:22 min for the WASSR sequence. Thus, the sequence requires 7:23 min. Hence, our scan time is comparable to the one presented...
by Kogan et al., who conducted the only previous study on gagCEST imaging of the ankle joint, and even shorter than several gagCEST studies focusing on the knee joint [13, 14, 42]. Additionally, the gagCEST imaging protocol was designed for 3 T scanners, which is the commonly used field strength for musculoskeletal imaging in clinical practice [43]. Thus, our protocol may be applied in both research and clinical contexts to further advance the clinical utility of gagCEST imaging of the tibiotalar joint. However, it still has to be considered less sensitive at detecting early cartilage changes than imaging protocols applied at 7 T scanners, especially if the latter are designed as volumetric multi-slice approaches [14]. Volumetric protocols have been implemented at 3 T scanners for gagCEST imaging of the knee joint and generally allow for better localization of cartilage changes. Consequently, future adaptation of volumetric protocols for gagCEST imaging of the tibiotalar joint seems of great scientific and clinical interest.

In addition to providing a stable and reproducible protocol, we observed significant differences between healthy volunteers and patients with OLTs. Since this study was the first of its kind comparing healthy individuals with patients using gagCEST at the ankle joint, we chose a patient cohort with morphologically damaged cartilage to demonstrate feasibility of this technique. In the future, we intend to study patients after ankle trauma without morphological apparent cartilage lesions to assess the presence of pre-morphological tissue damage.

Despite its strengths, our study has limitations. Our measured T1 and T2 relaxation times were shorter than the ones used for the simulations, but were overall comparable to the current literature [44].

Synovial fluid in general and joint effusion in particular are known to interfere with gagCEST imaging due to the presence of GAGs [1, 45]. Therefore, we placed our ROIs in the center of the tibiotalar joint at a distance to the anterior and posterior anatomical recesses, where joint fluid may collect and distort our measurements. A priori, we excluded patients with manifest joint effusion as visible in the morphological sequences. However, since we included both cartilage layers, i.e., both tibial and talar, in one single ROI, the odds are high that synovial fluid might have contaminated our gagCEST measures. Future studies should, therefore, use sequences that use fluid suppression. Moreover, our study population was small, which may be explained by the fact that we set out to implement a clinically applicable imaging protocol for gagCEST imaging. Nonetheless, future studies need to be conducted to corroborate our findings in larger patient numbers. Furthermore, we did not compare our findings to the gold-standard technique dGEMRIC. Since dGEMRIC relies on gadolinium-based contrast agents and its use is restricted due to ethical reasons, we consider this only a minor limitation. Last, we used a two-pool exchange model considering only the water- and the GAG-OH pool for the simulation. This model might be partially inaccurate for in-vivo applications, because of other influencing factors such as the GAG-NH pool, the nuclear Overhauser effect (NOE), and the magnetization transfer (MT) that were not included in our simulation because of lacking application-specific-framework fitting parameters for the NOE and MT. However, for the eventual quantification of the in-vivo measurements we used both the MTRasym values and the Lorentzian fit analyses. While the former accounts only for the water and the GAG-OH pool the latter also takes the GAG-NH, NOE and magnetization transfer pools into consideration. As both were strongly correlated, we consider the more simple two-pool exchange model to be sufficient for in-vivo quantification purposes.

In this feasibility study, pre-morphological tibiotalar joint cartilage damage was quantitatively assessable on the basis of an optimized 3 T gagCEST imaging protocol that allowed a stable gagCEST effect quantification both in normal and degenerated cartilage in clinically feasible acquisition times.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Ethical standards All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki Declaration and its later amendments or comparable ethical standards.
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