



High contrast cartilaginous endplate imaging in spine using three dimensional dual-inversion recovery prepared ultrashort echo time (3D DIR-UTE) sequence

Jiyo S. Athertya¹ · James Lo^{1,2} · Xiaojun Chen¹ · Soo Hyun Shin¹ · Bhavsimran Singh Malhi¹ · Saeed Jerban¹ · Yang Ji¹ · Sam Sedaghat¹ · Hiroshi Yoshioka³ · Jiang Du^{1,2,4} · Monica Guma^{5,6} · Eric Y. Chang^{1,4} · Yajun Ma¹

Received: 17 July 2023 / Revised: 14 October 2023 / Accepted: 29 October 2023 / Published online: 8 November 2023

© The Author(s) 2023

Abstract

Purpose To investigate the feasibility and application of a novel imaging technique, a three-dimensional dual adiabatic inversion recovery prepared ultrashort echo time (3D DIR-UTE) sequence, for high contrast assessment of cartilaginous endplate (CEP) imaging with head-to-head comparisons between other UTE imaging techniques.

Method The DIR-UTE sequence employs two narrow-band adiabatic full passage (AFP) pulses to suppress signals from long T_2 water (e.g., nucleus pulposus (NP)) and bone marrow fat (BMF) independently, followed by multispoke UTE acquisition to detect signals from the CEP with short T_2 relaxation times. The DIR-UTE sequence, in addition to three other UTE sequences namely, an IR-prepared and fat-saturated UTE (IR-FS-UTE), a T_1 -weighted and fat-saturated UTE sequence (T_{1w} -FS-UTE), and a fat-saturated UTE (FS-UTE) was used for MR imaging on a 3 T scanner to image six asymptomatic volunteers, six patients with low back pain, as well as a human cadaveric specimen. The contrast-to-noise ratio of the CEP relative to the adjacent structures—specifically the NP and BMF—was then compared from the acquired images across the different UTE sequences.

Results For asymptomatic volunteers, the DIR-UTE sequence showed significantly higher contrast-to-noise ratio values between the CEP and BMF ($CNR_{CEP-BMF}$) (19.9 ± 3.0) and between the CEP and NP (CNR_{CEP-NP}) (23.1 ± 1.7) compared to IR-FS-UTE ($CNR_{CEP-BMF}$: 17.3 ± 1.2 and CNR_{CEP-NP} : 19.1 ± 1.8), T_{1w} -FS-UTE ($CNR_{CEP-BMF}$: 9.0 ± 2.7 and CNR_{CEP-NP} : 10.4 ± 3.5), and FS-UTE ($CNR_{CEP-BMF}$: 7.7 ± 2.2 and CNR_{CEP-NP} : 5.8 ± 2.4) for asymptomatic volunteers (all P -values < 0.001). For the spine sample and patients with low back pain, the DIR-UTE technique detected abnormalities such as irregularities and focal defects in the CEP regions.

Conclusion The 3D DIR-UTE sequence is able to provide high-contrast volumetric CEP imaging for human spines on a clinical 3 T scanner.

Keywords Cartilaginous endplate · High contrast · Dual adiabatic inversion recovery · UTE · Low back pain

Abbreviation

LBP	Low back pain	UTE	Ultrashort echo time
IVD	Intervertebral disc	OCJ	Osteochondral junction
CEP	Cartilaginous endplate	T_{1w} -FS-UTE	T_1 -weighted fat-saturated UTE

✉ Yajun Ma
yam013@health.ucsd.edu

¹ Department of Radiology, University of California San Diego, San Diego, CA, USA

² Department of Bioengineering, University of California San Diego, San Diego, CA, USA

³ Department of Radiological Sciences, University of California Irvine, Irvine, CA, USA

⁴ Radiology Service, Veterans Affairs San Diego Healthcare System, San Diego, CA, USA

⁵ Department of Medicine, University of California San Diego, San Diego, CA, USA

⁶ Medicine Service, Veterans Affairs San Diego Healthcare System, San Diego, CA, USA

IR-FS-UTE	Inversion recovery-prepared and fat-saturated UTE
DIR-UTE	Dual inversion recovery-prepared UTE
CNR	Contrast-to-noise ratio
BMF	Bone marrow fat
AFP	Adiabatic full passage
TI	Inversion time
VERSE	Variable-rate selective excitation

Introduction

Low back pain (LBP) is one of the major causes of disability, affecting approximately 25% of adults in the United States [1, 2]. The need for accurate diagnoses as well as focused preventive and therapeutic strategies in this condition have become a major public health priority [3]. Intervertebral disc (IVD) degeneration has been recognized as one of the main causes of chronic low back pain [4]. Many of these cases have been associated with abnormality of the cartilaginous endplate (CEP) [5].

Vertebral endplate structure lies at the cranial and caudal interface of the intervertebral disc on one side and vertebral body, which consists of an osseous component and a hyaline cartilage component (known as the CEP), on the other [6]. The CEP comprised of a hydrated proteoglycan matrix, fortified by an interlaced framework of collagen fibrils, which directly attaches to the intervertebral disc via the lamellae within the inner (medial) annulus fibrosus. No direct connection is found between CEP and the osseous structure of the vertebral bodies [7].

The CEP acts as the nutrient transport bridge from blood vessels to the disc cells, which is of critical importance to maintaining disc health [8, 9]. Degeneration, mineralization, and dehydration of the CEP due to age, injury, and degradation reduce the CEP's permeability, leading to a diminished capacity for nutrient transport. Thus, the health of the CEP could be highly related to early IVD degeneration and associated LBP. Moreover, the CEP is also subjected to mechanical loads which are distributed onto adjacent vertebrae as a means to prevent the disc nucleus from bulging under pressure [10]. Damage to this region might present as a painful pathology [11]. Thus, evaluation of the CEP region may be critical for the assessment of LBP and diagnosis of early IVD degeneration.

Magnetic resonance imaging (MRI) is a powerful tool for the diagnosis of IVD degeneration. The most commonly used techniques are the T_1 - and T_2 -weighted sequences [12].

However, because the CEP has a relatively short transverse relaxation time (T_2 : ~18 ms, T_2^* : ~15 ms) [13, 14], these routine clinical sequences cannot detect sufficient CEP signals for direct imaging or quantification, thus limiting the effectiveness of MRI for early evaluation of IVD degeneration.

Ultrashort echo time (UTE) sequences with echo times shorter than 100 μ s are able to detect many short T_2 tissues, such as the CEP, tendons, and bone, positioning this type of sequence as a promising approach to image and evaluate those tissues [15, 16]. Recently, several UTE techniques have been developed for long T_2 suppression and selective imaging of short T_2 tissues, such as dual-echo UTE with subtraction [17], T_1 -weighted fat-saturated UTE (T_{1w} -FS-UTE) [18], inversion recovery-prepared and fat-saturated UTE (IR-FS-UTE) [19], and dual inversion recovery-prepared UTE (DIR-UTE) [20] sequences. The dual-echo UTE is a fast technique but suffers from low contrast-to-noise ratio (CNR) between the CEP and bone marrow fat (BMF) due to the relatively fast T_2^* decay of BMF [21]. The fast T_{1w} -FS-UTE sequence has been applied for high-contrast imaging of the osteochondral junction (OCJ) with better contrast between the CEP and BMF than the dual-echo subtraction method [18]. Moreover, the UTE sequences which incorporate adiabatic inversion, such as IR-FS-UTE, are very efficient in generating high contrast between short and long T_2 tissues [19, 20, 22–24] and have been used for high contrast imaging of the OCJ [25]. Such techniques can aid in imaging the CEP with high resolution [19]. In CEP imaging, the IR-FS-UTE sequence can produce a very high contrast between the CEP and nucleus pulposus (NP), but the contrast between the CEP and BMF is limited because of the chemical fat saturation (FatSat) module that prohibits efficient fat suppression if more acquisition spokes are used in a repetition time (TR). This inefficient fat suppression can be overcome by replacing the FatSat module with another adiabatic inversion pulse that is centered on the fat frequency, i.e., the DIR-UTE technique [20]. The DIR-UTE sequence has been shown to generate a very high image contrast for the OCJ region with efficient suppression of signals from both the long T_2 cartilage and marrow fat [20].

In this study, we aim to image the CEP instead of the bony endplate. Bony endplate and subchondral bone structures are dark in the proposed DIR-UTE images because cortical bone typically has a much lower proton density than cartilage tissues including the CEP [26, 27]. Given the advantages of DIR-UTE sequence in imaging of short T_2 tissues, we proposed to further optimize it for high contrast imaging of CEP in the human lumbar spine, and compare its performance with other established UTE techniques, namely IR-FS-UTE, T_{1w} -FS-UTE, and FS-UTE.

Methods

MR acquisition and pulse sequence

This study was approved by the institutional review board. All sequences were implemented on a 3 T MR750 scanner (GE Healthcare Technologies, Milwaukee, Wisconsin) with a four-channel phased array spine coil utilized for signal reception.

Figure 1 shows the sequence diagrams for the four different UTE techniques used in this study, including DIR-UTE, IR-FS-UTE, T_{1w} -FS-UTE, and FS-UTE sequences. The DIR-UTE sequence utilizes two adiabatic full passage (AFP) pulses to invert long T_2 water (e.g., NP) and BMF with center frequencies of 0 and -440 Hz, respectively [20]. The IR-FS-UTE sequence employs an AFP pulse for inverting long T_2 NP while the FatSat module is utilized to improve CEP contrast against BMF [19]. The T_{1w} -FS-UTE shares the same sequence diagram with the FS-UTE but with much higher flip angles (thus stronger T_1 weighting) [18]. The FatSat module is applied for fat suppression in both T_{1w} -FS-UTE and FS-UTE sequences. The multispoke acquisition strategy is employed in all UTE sequences to

reduce the total scan time. For signal excitation in each spoke, a slab selective half pulse (Shinnar-Le Roux design, duration 1132 μ s, and bandwidth 16 kHz) with variable-rate selective excitation (VERSE) design [28] is utilized. The 3D Cones trajectory enables efficient k-space coverage for all UTE scans.

In the DIR-UTE sequence, the longitudinal magnetizations of NP and BMF are efficiently inverted by two narrow-band AFP (e.g., 500 Hz) pulses. With a proper selection of inversion time (TI) for each inversion, signal nulling for both NP and BMF could be reached at the time of data acquisition simultaneously [22]. On the other hand, the inverted CEP signal recovers quickly because of the very short T_1 property of CEP (\sim 400 ms) [19]. When the data acquisition commences around the signal nulling point of the NP and BMF, the CEP presents with high signal and is therefore highlighted.

In vivo human spine imaging

Twelve volunteers (six asymptomatic subjects and six patients with LBP) were recruited for lumbar spine imaging using both clinical (i.e., T_{1w} - and T_{2w} -FSE) and UTE (i.e., DIR-UTE, IR-FS-UTE, T_{1w} -FS-UTE, and FS-UTE)

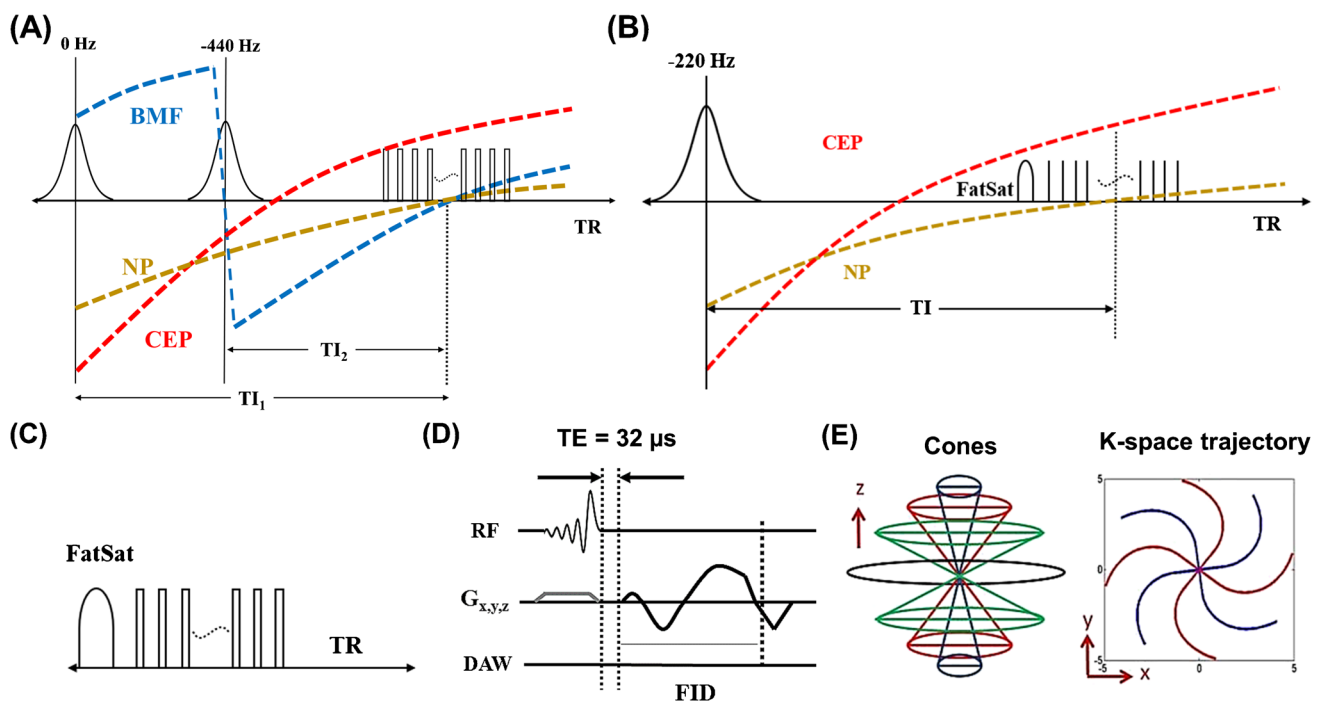


Fig. 1 Sequence diagram for four different UTE techniques, including DIR-UTE (A) IR-FS-UTE (B) T_{1w} -FS-UTE (C), and FS-UTE (C). The DIR-UTE sequence utilizes two AFP pulses to invert long T_2 water (e.g., NP) and BMF with center frequencies of 0 and -440 Hz, respectively (A). The IR-FS-UTE sequence employs an AFP pulse for inverting long NP while the FatSat module is utilized to improve

CEP contrast against BMF (B). The FatSat module is applied for fat suppression in both T_{1w} -FS-UTE and FS-UTE sequences (C). The multispoke acquisition strategy is employed in all UTE sequences to reduce the total scan time. A slab selective half pulse is utilized for signal excitation in each spoke (D). The 3D Cones trajectory enables efficient k-space coverage for all UTE scans (E)

sequences. The age of asymptomatic subjects ranged between 23 and 38 years old, while that of the patient population ranged between 37 and 60 years old. Informed consent was obtained from all participants per institutional review board requirements.

To determine the optimal TIs for high contrast CEP imaging, DIR-UTE scans were performed on a single asymptomatic volunteer with a series of different TI_1 s and TI_2 s (TIs can be seen in Supplemental Information Table 1). The other major parameters for each scan were TR/echo time (TE) = 1500/0.032 ms, flip angle (FA) = 10° , number of spokes per TR (N_{sp}) = 27, field-of-view (FOV) = $28 \times 28 \times 4.8 \text{ cm}^3$, matrix = $220 \times 220 \times 12$, voxel size = $1.273 \times 1.273 \times 4 \text{ mm}^3$, bandwidth = 125 kHz, and scan time = 6 min.

The remaining asymptomatic volunteers and symptomatic patients were scanned with the resultant optimized protocol: (i) DIR-UTE: TR/ TI_1 / TI_2 = 1500/610/150 ms, TE = 0.032 ms, FA = 10° , N_{sp} = 27, FOV = $28 \times 28 \times 4.32 \text{ cm}^3$, matrix = $320 \times 320 \times 12$, voxel size = $0.875 \times 0.875 \times 3.6 \text{ mm}^3$, bandwidth = 125 kHz, and scan time = 10 min; (ii) IR-FS-UTE [19]: TR/TI = 1200/600 ms, TE = 0.032 ms, FA = 10° , N_{sp} = 21, FOV = $28 \times 28 \times 4.32 \text{ cm}^3$, matrix = $320 \times 320 \times 12$, voxel size = $0.875 \times 0.875 \times 3.6 \text{ mm}^3$, bandwidth = 125 kHz, and scan time = 10 min; (iii) T_{1w} -FS-UTE: TR = 120 ms, TE = 0.032 ms, FA = 15° , FOV = $28 \times 28 \times 4.32 \text{ cm}^3$, matrix = $320 \times 320 \times 12$, voxel size = $0.875 \times 0.875 \times 3.6 \text{ mm}^3$, bandwidth = 125 kHz, and scan time = 2 min 20 s; (iv) FS-UTE: TR = 120 ms, TE = 0.032 ms, FA = 6° , FOV = $28 \times 28 \times 4.32 \text{ cm}^3$, matrix = $320 \times 320 \times 12$, voxel size = $0.875 \times 0.875 \times 3.6 \text{ mm}^3$, bandwidth = 125 kHz, and scan time = 2 min 20 s. Clinical T_{1w} - and T_{2w} -FSE sequences were scanned for comparison. In addition, for the patient scan, a proton density weighted-UTE (PDw-UTE) sequence with an isotropic resolution (1.2 mm^3) was also employed for high contrast cortical bone imaging [29]. The sequence parameters used in the other UTE sequences were optimized in previous studies.

Ex vivo human spine imaging

A cadaveric spine sample section (T11-T12) from an 87-year-old female donor was scanned using the DIR-UTE sequence with the following parameters: TR/ TI_1 / TI_2 = 1500/610/150 ms, TE = 0.032 ms, FA = 10° , N_{sp} = 27, FOV = $12 \times 12 \times 4 \text{ cm}^3$, matrix = $320 \times 320 \times 20$, voxel size = $0.375 \times 0.375 \times 2 \text{ mm}^3$, bandwidth = 83.3 kHz, scan time = 16 min. An eight channel-knee coil was used for radiofrequency (RF) transmission and signal reception. Clinical T_{1w} - and T_{2w} -FSE sequences were scanned for comparison.

Data processing

To compare the CEP contrast between the four different UTE techniques on asymptomatic volunteers, the CNRs between the CEP and NP (CNR_{CEP-NP}) and between the CEP and BMF ($CNR_{CEP-BMF}$) were calculated as the mean differences in signals between these tissues divided by the background noise. Representative ROIs of CEP, NP and BMF can be seen in Fig. 2. The noise was estimated as the standard deviation of signals measured from a region of interest (ROI) ($\sim 6 \times 6 \text{ cm}^2$) in an artifact-free background region. Descriptive statistics were performed while mean and standard deviation of the CNR_{CEP-NP} and $CNR_{CEP-BMF}$ were measured to evaluate contrast between CEP and NP and between CEP and BMF for all the UTE sequences. Paired t-test using SPSS software (IBM, Armonk, NY, USA) version 28.0 was utilized to compare CNR values of CEP imaging between all the techniques. P-values lower than 0.05 are considered as significant.

Results

Supplemental Information Fig. 1 shows the DIR-UTE images acquired from a 32-year-old asymptomatic male volunteer at various TI combinations. To optimize the image contrast between CEP and NP, TI_1 varies from 580 to 640 ms while TI_2 is fixed at 150 ms. Similarly, to optimize the contrast between CEP and BMF, TI_2 varies from 125 to 175 ms while TI_1 is fixed at 610 ms. As can be seen from the figure, the images resulting from these different TI combinations all show good CEP contrast visually, proving that the DIR-UTE technique can effectively highlight the CEP region over a wide range of TI_1 s and TI_2 s. This demonstrates the robustness of the DIR-UTE technique for high contrast imaging. Supplemental Information Table 1 lists the measured CNR_{CEP-NP} and $CNR_{CEP-BMF}$ values for the images shown in Supplemental Information Fig. 1. When TI_2 is fixed at 150 ms, the CNR_{CEP-NP} values increase from 17.8 to 19.0, then decrease to 17.9 when TI_1 increases from 580 to 640 ms. In addition, when TI_1 is fixed at 610 ms, the $CNR_{CEP-BMF}$ values increase from 17.0 to 17.6, then decrease to 15.9 when TI_2 increases from 125 to 175 ms. The highest CNR_{CEP-NP} and $CNR_{CEP-BMF}$ values are achieved when $TI_1 = 610 \text{ ms}$ and $TI_2 = 150 \text{ ms}$. These optimized TIs were employed for the subsequent DIR-UTE imaging of both asymptomatic volunteers and patients.

Figure 2 shows the representative lumbar spine images from two asymptomatic volunteers. The CEP signal cannot be efficiently captured in the clinical fat suppressed T_{2w} -FSE sequence owing to its relatively short T_2 relaxation time, whereas it is clearly seen on all UTE images. The DIR-UTE, IR-FS-UTE, and T_{1w} -FS-UTE sequences all produce higher

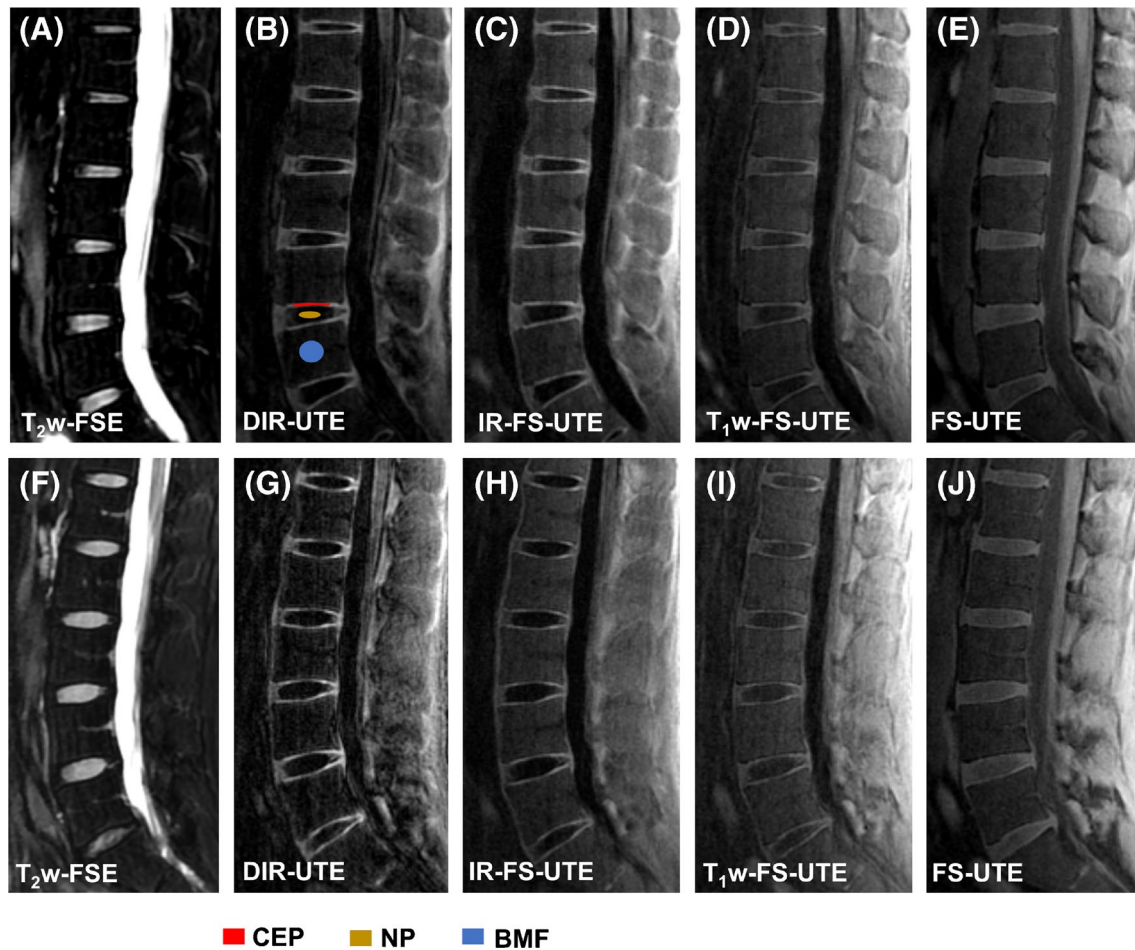


Fig. 2 Lumbar spine imaging for two asymptomatic volunteers. Images from a 32-year-old asymptomatic male subject are shown in (A–E) while (F–J) correspond to images obtained from a 38-year-old asymptomatic male volunteer. The clinical fat suppressed T_{2w} -FSE sequence shown in (A and F) fails to capture CEP signals owing to its long TE relative to the CEP’s short T_2 relaxation. The

DIR-UTE (B and G) provides the best contrast in comparison to other UTE imaging techniques, including IR-FS-UTE (C and H), T_{1w} -FS-UTE (D and I), and FS-UTE (E and J), between CEP and BMF and between CEP and NP. Panel B shows representative ROIs of CEP, NP and BMF to calculate CEP CNRs

Table 1 Summary of the mean CNR_{CEP-NP} and $CNR_{CEP-BMF}$ measurements from the DIR-UTE, IR-FS-UTE, T_{1w} -FS-UTE, and FS-UTE images for six asymptomatic volunteers

CNR	DIR-UTE	IR-FS-UTE	T_{1w} -FS-UTE	FS-UTE
Between CEP and NP	23.1 ± 1.7	19.1 ± 1.8	10.4 ± 3.5	5.8 ± 2.4
Between CEP and BMF	19.9 ± 3.0	17.3 ± 1.2	9.0 ± 2.7	7.7 ± 2.2

BMF Bone marrow fat; NP Nucleus pulposus

CEP contrast than the regular FS-UTE sequence because of their higher T_1 -weighting property. The DIR-UTE images show the best CEP contrast for visual comparison.

Table 1 summarizes the measured CNR_{CEP-NP} and $CNR_{CEP-BMF}$ values of all four UTE sequences for the six asymptomatic volunteers. Of the different sequences used,

the DIR-UTE presents the highest CEP contrast for CEP vs. NP (23.1 ± 1.7) and CEP vs. BMF (19.9 ± 3.0), followed by IR-FS-UTE, T_{1w} -FS-UTE, and FS-UTE. The DIR-UTE showed significantly higher CNRs in both CNR_{CEP-NP} and $CNR_{CEP-BMF}$ compared to IR-FS-UTE ($p = 0.0005$, $p = 0.0109$), T_{1w} -FS-UTE ($p = 0.0017$, $p = 0.0003$), and FS-UTE ($p = 0.0010$, $p = 0.0008$) respectively.

Results from the ex vivo sample study are presented in Fig. 3. As seen from Fig. 3C, signals from long T_2 water and fat are well suppressed by the DIR technique. The high-intensity band of CEP demonstrates a thickness from 0.6 mm to 1.2 mm. In comparison, the CEP is dark in both clinical T_{2w} - and T_{1w} -FSE images due to its short T_2 . The lower level of this disc is particularly interesting, with the presence of a compression fracture deformity and central Schmorl’s node at the superior endplate of the lower spine (orange arrows). This focal CEP fracture is clearly evident as a disruption in

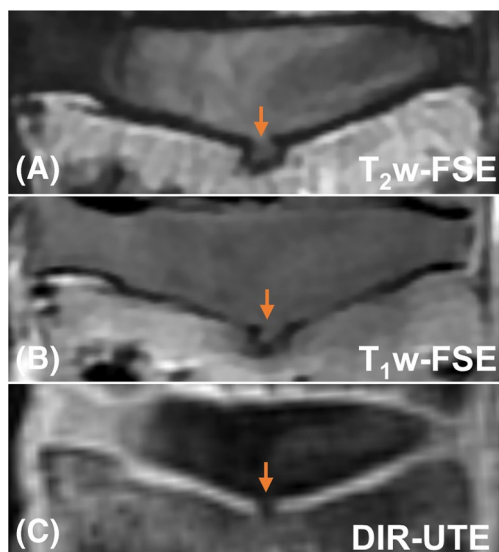


Fig. 3 The clinical T_{2w} - and T_{1w} -FSE (A and B) as well as DIR-UTE (C) imaging for an ex vivo spine sample from an 87-year-old female donor. There is a CEP fracture with herniation of NP through the focal defect as indicated on the DIR-UTE image with an orange arrow

the high signal band as shown on the DIR-UTE image, but is ambiguous in the clinical T_{2w} - and T_{1w} -FSE images.

Figure 4 shows representative lumbar spine images from a patient with low back pain. Similar to the case with the asymptomatic subjects, the clinical fat suppressed T_{2w} -FSE sequence does not capture signals from the CEP region, while the DIR-UTE, IR-FS-UTE, and T_{1w} -FS-UTE images present better CEP contrast than regular FS-UTE. The NP

regions of degenerated discs in the DIR-UTE, IR-FS-UTE, and T_{1w} -FS-UTE images show relatively higher signals than those in normal discs. This may be because of shortened T_1 relaxation time in the NP due to disc dehydration [8, 30, 31].

Figure 5 shows UTE imaging of a patient with psoriatic arthritis (PsA). This patient has features of chronic inflammatory arthritis, including a Romanus lesion at the anterosuperior L2 vertebral body with erosion (shown in PDw-UTE bone image) and fatty change (shown in clinical T_{1w} -FSE image). Numerous bony endplate irregularities as marked by arrows are associated with the signal loss in CEP regions seen vividly in the coronal DIR-UTE image (Fig. 5C). In comparison, continuous CEP signals can be seen in the coronal DIR-UTE image from a asymptomatic subject (Fig. 5D).

Figure 6 presents DIR-UTE imaging of another patient with PsA. The L2-L4 levels show bony endplate remodeling on the PDw-UTE bone image. The CEP irregularities can be clearly seen in the corresponding DIR-UTE image at the inferior endplates of these discs (arrows). In comparison, the clinical sequences do not provide such information on CEP changes.

Discussion

In this work, we have shown that the DIR-UTE sequence can provide high contrast imaging of the CEP region in both ex vivo and in vivo spine studies. Usage of two narrow-band AFP pulses in the DIR-UTE sequence can efficiently suppress long T_2 water and fat simultaneously. Moreover, the CEP region can be imaged with high contrast over a wide

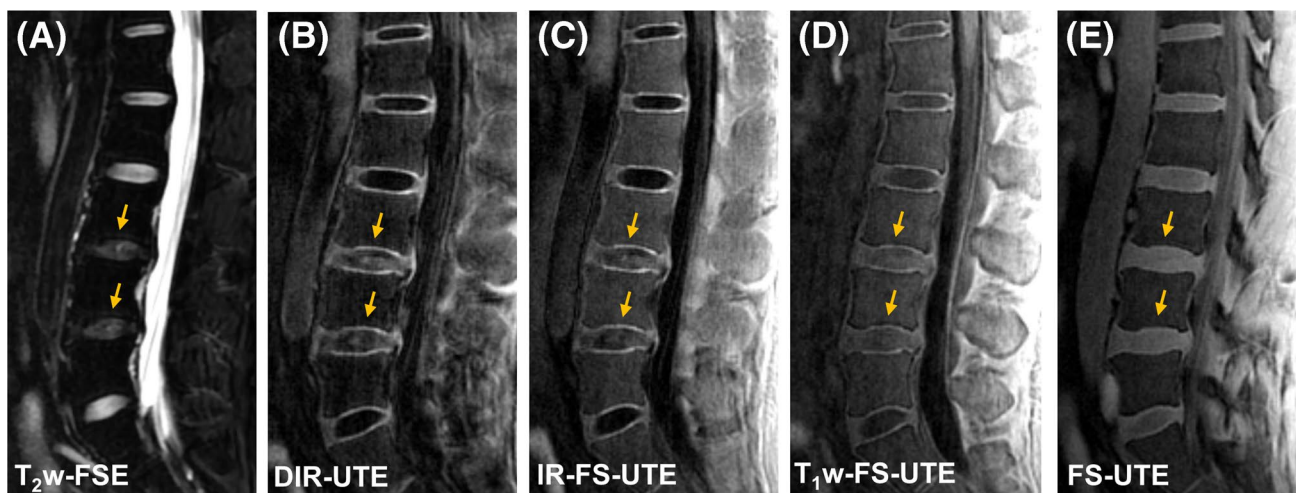


Fig. 4 Representative lumbar spine imaging from a 40-year-old male patient with low back pain. While the CEP region is completely invisible in the clinical fat suppressed T_{2w} -FSE (A) it is well highlighted in the DIR-UTE (B) IR-FS-UTE (C) and T_{1w} -FS-UTE (D). Regular FS-UTE (E) is able to capture the CEP signal, but has relatively low

contrast of the CEP region compared to the other UTE imaging techniques. The NP regions of degenerated discs (indicated by arrows) exhibit stronger signals than those in normal discs for the DIR-UTE, IR-FS-UTE, and T_{1w} -FS-UTE, which can be attributed to the shortened T_1 relaxation due to dehydration in NP

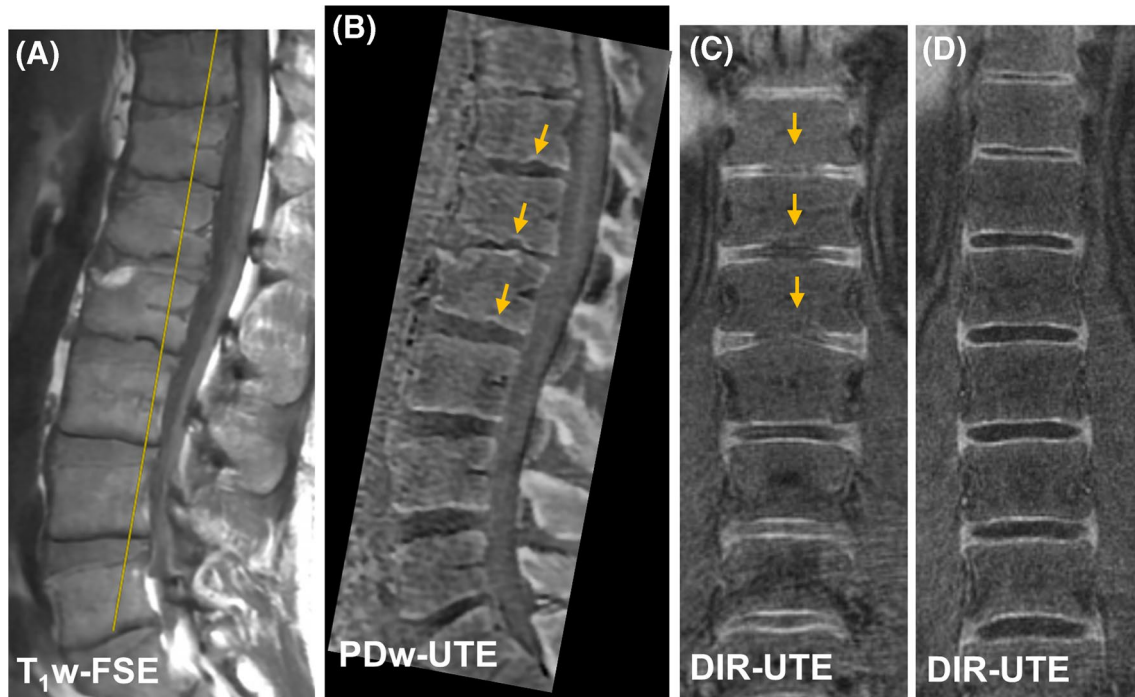
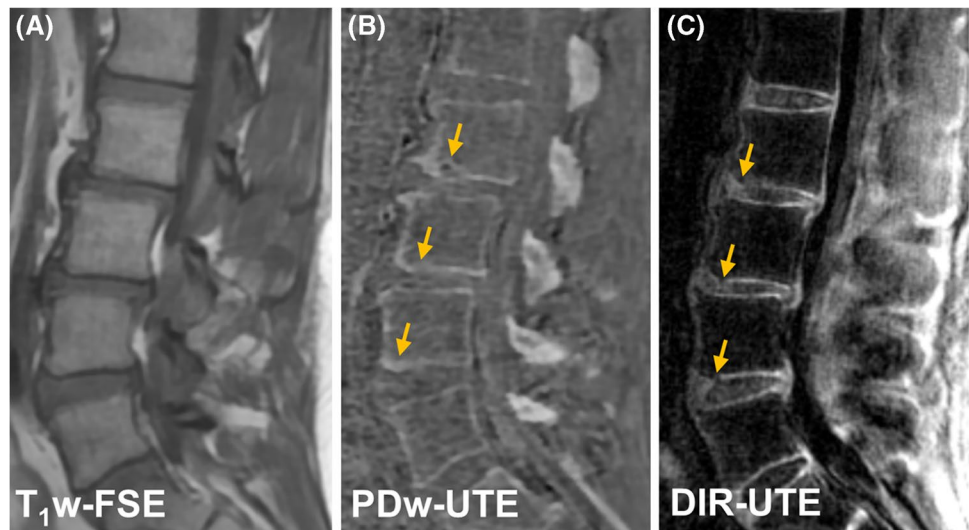


Fig. 5 The clinical T_{1w} -FSE (A) PDw-UTE (B) and DIR-UTE (C) images from a 37-year-old male with known PsA. Yellow line in (A) on the clinical image indicates the cross-section plane for coronal slice in DIR-UTE image. The CEP irregularities as marked by

arrows are depicted in the coronal DIR-UTE image (C). The DIR-UTE image obtained from another asymptomatic subject (24-year-old male) (D) is used for comparison where the CEP region shows continuous and bright signal

Fig. 6 The clinical T_{1w} -FSE (A) PDw-UTE (B) and DIR-UTE (C) images from a 51-year-old male patient with PsA. The CEP irregularities are clearly seen on the inferior endplates at L2, L3, and L4 with the DIR-UTE sequence. These CEP irregularities are associated with bony endplate remodeling shown in the PDw-UTE bone image and were caused by Schmorl's nodes



range of TIs in the DIR-UTE, which indicates the robustness of the proposed technique. In addition, the DIR-UTE sequence highlights the CEP with significant improvements in comparison to IR-FS-UTE, T_{1w} -FS-UTE, and FS-UTE sequences. Ex vivo imaging on a spine sample revealed a compression fracture that was clearly discernible by the DIR-UTE sequence. In vivo study on various asymptomatic and symptomatic subjects has confirmed that DIR-UTE

sequence can capture CEP signal with high contrast and can detect abnormalities such as CEP irregularities and focal defects.

The proposed 3D DIR-UTE sequence provides better CEP contrast in comparison to previously reported dual-echo subtraction UTE and IR-FS-UTE studies, especially with regard to the contrast between the CEP and BMF [19, 32, 33]. To the best of our knowledge, this is the first in vivo

CEP imaging study that uses the 3D DIR-UTE Cones technique; other studies have been focused on the T_2^* quantification of CEP using multi-echo UTE sequences [14, 34, 35]. Our study has demonstrated that this technique has great potential for morphological evaluation of the CEP and IVD health in vivo. The DIR-UTE technique incorporated with multiple-echo acquisition also allows for quantitative evaluation of CEP degeneration or calcification, which could be a new direction for future investigations.

The CEP region is an important facilitator in both biomechanical and nutritional functions of the spine. Given that the CEP is regularly subjected to significant loads during daily activities in order to stabilize spinal posture [36], end plate disruptions are likely to disrupt the uniformity of disc stress distributions [37]. This in turn accelerates matrix degradation in the CEP thereby causing the impediment of nutrient transport to the cells [38]. This phenomenon can be interpreted as the initial stages of disc degeneration and would be highly significant as a marker observable via non-invasive imaging techniques [5]. Given that conventional MRI sequences fail to capture sufficient signals from the CEP region, devising and developing novel high contrast imaging sequences, such as the DIR-UTE presented in this study, are of acute importance.

The IR preparation is much more efficient in signal suppression for the desired tissues with long T_2 relaxations than T_1 weighting (as used in the T_{1w} -FS-UTE sequence) is, with regard to short T_2 imaging. Moreover, since the CEP and BMF both have relatively short T_1 values, it is difficult to distinguish the regions using the T_{1w} -FS-UTE sequence when fat suppression is not complete. In addition, the effectiveness of the FatSat module is limited up to a certain level in the IR-FS-UTE and T_{1w} -FS-UTE imaging. When more spokes are utilized to speed up the imaging acquisition, the effectiveness of fat suppression for the FatSat module diminishes, resulting in a trade-off between scan efficiency and fat suppression. This fat suppression inefficiency can be avoided by replacing the FatSat module with another AFP pulse dedicated specifically to fat suppression by centering the frequency on fat.

The DIR-UTE sequence is able to detect and highlight CEP signals, allowing for more accurate morphological evaluation of CEP health in comparison to clinical MRI. For example, calcification and ossification of the CEP cannot be assessed by clinical MRI because these conditions show a similar dark appearance as normal CEP. In comparison, the DIR-UTE may be able to differentiate normal CEP from calcified and ossified CEPs due to the differences in proton density. In addition to morphological assessment of CEP health, the DIR-UTE with multiple-echo acquisition allows quantitative evaluation (i.e., T_2^* measurement) of CEP biochemistry which clinical MRI cannot. Thus the DIR-UTE technique has the potential to

provide more useful information than clinical MRI in the assessment of CEP and IVD health.

However, the DIR-UTE imaging could be affected by artifacts brought on by the signal fluctuations among various acquisition spokes if there is a significant increase in the number of spokes employed in one TR. For the current study, the number of spokes ($N_{sp} = 27$) was found to provide a high CNR without significant artifacts being introduced. A range of 15 to 30 for N_{sp} is recommended in terms of scan time and image quality.

There are several limitations in this study. First, histology was not performed to validate the signal changes of 3D DIR-UTE imaging. Second, the spatial resolution (e.g., voxel size = $0.875 \times 0.875 \times 3.6\text{mm}^3$) was still a limiting factor for thin CEP imaging considering the signal-to-noise ratio performance of the RF coils and MRI scanners currently available, despite the high CEP contrast in the 3D DIR-UTE imaging. Third, the scan time of 10 min is relatively long in the context of a clinical setting and should be improved with parallel imaging and compressed sensing [39–41]. Fourth, the DIR-UTE sequence is sensitive to background magnetic field inhomogeneity due to narrow bandwidth of the AFP pulses and limited chemical shift of fat at 3T. A high order shimming hardware system could potentially help for uniform signal suppression. Fifth, a relatively small number of subjects were scanned in this study. A future large cohort study will be very useful for exploring the potential of the DIR-UTE technique in clinical use. Sixth, in this study, only the DIR-UTE technique was applied for the patient scans to shorten the total scan time since these patients cannot bear a very long scan due to their moderate low back pain. The comparison of CEP contrast between all the UTE techniques for patient study will be interesting once the scan time of these UTE scans is accelerated.

In conclusion, we showed that 3D DIR-UTE sequence can be used for high contrast CEP imaging both ex vivo and in vivo. The optimized 3D DIR-UTE sequence also showed better CEP contrast than other UTE techniques, including the IR-FS-UTE, T_{1w} -FS-UTE, and FS-UTE sequences, suggesting that the DIR-UTE sequence may facilitate better evaluation of the vital CEP region in clinical practice.

Clinical Relevance The proposed DIR-UTE sequence could potentially assist in characterizing signal variations in CEP owing to its high CNR. Evaluation of abnormalities in the other human spine tissues, such as facet joints, longitudinal ligaments, and intervertebral foramina, are further potential applications.

Supplementary Information The online version contains supplementary material available at <https://doi.org/10.1007/s00256-023-04503-4>.

Acknowledgements The authors acknowledge grant support from the National Institutes of Health (K01AR080257, F32AG082458, R01AR062581, R01AR068987, R01AR075825, and R01AR079484), the VA Clinical Science Research and Development Service (I01RX002604 and I01CX001388).

Data availability Data supporting the reported results can be provided by the corresponding author upon courtesy request.

Declarations

Competing interests The authors have no conflicts of interest to declare.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>.

References

- Nahin RL. Estimates of pain prevalence and severity in adults: United States, 2012. *J Pain* [Internet]. 2015;16:769–80. <https://doi.org/10.1016/j.jpain.2015.05.002>.
- Martin BI, Deyo RA, Mirza SK, Turner JA, Comstock BA, Holtingworth W, et al. Expenditures and health status among adults with back and neck problems. *JAMA*. 2008;299:656–64.
- Hoy D, Brooks P, Blyth F, Buchbinder R. The epidemiology of low back pain. *Best Pract Res Clin Rheumatol* [Internet]. 2010;24:769–81. <https://doi.org/10.1016/j.berh.2010.10.002>.
- Zheng CJ, Chen J. Disc degeneration implies low back pain. *Theor Biol Med Model* [Internet]. 2015;12:1–10. <https://doi.org/10.1186/s12976-015-0020-3>.
- Van Dieën JH, Weinans H, Toussaint HM. Fractures of the lumbar vertebral endplate in the etiology of low back pain: a hypothesis on the causative role of spinal compression in aspecific low back pain. *Med Hypotheses*. 1999;53:246–52.
- Moore RJ. The vertebral end-plate: what do we know? *Eur Spine J*. 2000;9:92–6.
- Roberts S, Menage J, Urban JPG. Biochemical and Structural properties of the cartilage end-plate and its relation to the intervertebral disc. *Spine* 1989;14:166–74.
- Antoniou J, Steffen T, Nelson F, Winterbottom N, Hollander AP, Poole RA, et al. The human lumbar intervertebral disc: evidence for changes in the biosynthesis and denaturation of the extracellular matrix with growth, maturation, ageing, and degeneration. *J Clin Invest*. 1996;98:996–1003.
- Arpinar VE, Rand SD, Klein AP, Maiman DJ, Muftuler LT. Changes in perfusion and diffusion in the endplate regions of degenerating intervertebral discs: a DCE-MRI study. *Eur Spine J*. 2015;24:2458–67.
- Yoganandan N, Maiman DJ, Pintar F, Ray G, Myklebust JB, Sances AJ, et al. Microtrauma in the lumbar spine: a cause of low back pain. *Neurosurgery*. 1988;23(2):162–8.
- Fagan AB, Moore R, Roberts BV, Blumbergs P, Fraser R. ISSLS prize winner: the innervation of the intervertebral disc: a quantitative analysis. *Spine (Phila Pa 1976)*. 2003;28:2570–6.
- Pfirschmann CWA, Metzendorf A, Zanetti M, Hodler J, Boos N. Magnetic resonance classification of lumbar intervertebral disc degeneration. *Spine (Phila Pa 1976)*. 2001;26:1873–8.
- Athertya JS, Lombardi AF, Wong J, Jang H, Jerban S, Du J, et al. Quantitative MR imaging of whole intervertebral disc: a pre-clinical sample study. *ISMRM*. 2022.
- Wang L, Han M, Wong J, Zheng P, Lazar AA, Krug R, et al. Evaluation of human cartilage endplate composition using MRI: spatial variation, association with adjacent disc degeneration, and in vivo repeatability. *J Orthop Res*. 2021;39:1470–8.
- Ma Y, Jang H, Jerban S, Chang EY, Chung CB, Bydder GM, et al. Making the invisible visible—ultrashort echo time magnetic resonance imaging: technical developments and applications. *Appl Phys Rev*. 2022;9(4):041303.
- Afsahi AM, Ma Y, Jang H, Jerban S, Chung CB, Chang EY, et al. Ultrashort echo time magnetic resonance imaging techniques: met and unmet needs in musculoskeletal imaging. *J Magn Reson Imaging*. 2022;55(6):1597–612.
- Du J, Bydder M, Takahashi AM, Carl M, Chung CB, Bydder GM. Short T2 contrast with three-dimensional ultrashort echo time imaging. *Magn Reson Imaging* [Internet]. 2011;29:470–82. <https://doi.org/10.1016/j.mri.2010.11.003>.
- Cai Z, Wei Z, Wu M, Jerban S, Jang H, Li S, et al. Knee osteochondral junction imaging using a fast 3D T1-weighted ultrashort echo time cones sequence at 3T. *Magn Reson Imaging* [Internet]. 2020;73:76–83. <https://doi.org/10.1016/j.mri.2020.08.003>.
- Lombardi AF, Wei Z, Wong J, Carl M, Lee RR, Wallace M, et al. High contrast cartilaginous endplate imaging using a 3D adiabatic inversion-recovery-prepared fat-saturated ultrashort echo time (3D IR-FS-UTE) sequence. *NMR Biomed*. 2021;34:1–11.
- Lombardi AF, Jang H, Wei Z, Jerban S, Wallace M, Masuda K, et al. High-contrast osteochondral junction imaging using a 3D dual adiabatic inversion recovery-prepared ultrashort echo time cones sequence. *NMR Biomed*. 2021;34:1–11.
- Lee YH, Kim S, Song HT, Kim IS, Suh JS. Weighted subtraction in 3D ultrashort echo time (UTE) imaging for visualization of short T2 tissues of the knee. *Acta radiol*. 2014;55:454–61.
- Du J, Takahashi AM, Bae WC, Chung CB, Bydder GM. Dual inversion recovery, ultrashort echo time (DIR UTE) imaging: creating high contrast for short-T2 species. *Magn Reson Med*. 2010;63:447–55.
- Jang H, Ma Y, Searleman AC, Carl M, Corey-Bloom J, Chang EY, et al. Inversion recovery UTE based volumetric myelin imaging in human brain using interleaved hybrid encoding. *Magn Reson Med*. 2020;83:950–61.
- Ma YJ, Jang H, Wei Z, Cai Z, Xue Y, Lee RR, et al. Myelin imaging in human brain using a short repetition time adiabatic inversion recovery prepared ultrashort echo time (STAIR-UTE) MRI sequence in multiple sclerosis. *Radiology*. 2020;297:392–404.
- Ma YJ, Jerban S, Carl M, Wan L, Guo T, Jang H, et al. Imaging of the region of the osteochondral junction (OCJ) using a 3D adiabatic inversion recovery prepared ultrashort echo time cones (3D IR-UTE-cones) sequence at 3 T. *NMR Biomed*. 2019;32:1–13.
- Jerban S, Ma Y, Li L, Jang H, Wan L, Guo T, et al. Volumetric mapping of bound and pore water as well as collagen protons in cortical bone using 3D ultrashort echo time cones MR imaging techniques. *Bone* [Internet]. 2019;127:120–8. <https://doi.org/10.1016/j.bone.2019.05.038>.
- Jerban S, Ma Y, Jang H, Namiranian B, Le N, Shirazian H, et al. Water proton density in human cortical bone obtained from ultrashort echo time (UTE) MRI predicts bone microstructural properties. *Magn Reson Imaging*. 2020;67:85–9.

28. Hargreaves BA, Cunningham CH, Nishimura DG, Conolly SM. Variable-rate selective excitation for rapid MRI sequences. *Magn Reson Med*. 2004;52:590–7. <https://doi.org/10.1002/mrm.20168>.
29. Afsahi AM, Lombardi AF, Wei Z, Carl M, Athertya J, Masuda K, et al. High-contrast lumbar spinal bone imaging using a 3D slab-selective UTE sequence. *Front Endocrinol (Lausanne)*. 2022;12:1–9.
30. Wei Z, Lombardi AF, Lee RR, Wallace M, Masuda K, Chang EY, et al. Comprehensive assessment of in vivo lumbar spine intervertebral discs using a 3D adiabatic T1 ρ prepared ultrashort echo time (UTE-Adiab-T1 ρ) pulse sequence. *Quant Imaging Med Surg*. 2022;12:269–80.
31. Urban JPG, Holm S, Maroudas A, Nachemson A. Nutrition of the intervertebral disc. *Clin Orthop Relat Res*. 1982;170:296–302.
32. Bae WC, Statum S, Zhang Z, Yamaguchi T, Wolfson T, Gamst AC, et al. Morphology of the cartilaginous endplates in human intervertebral disks with ultrashort echo time MR imaging. *Radiology*. 2013;266:564–74.
33. Finkenstaedt T, Siritwanrangsun P, Masuda K, Bydder GM, Chen KC, Bae WC. Ultrashort time-to-echo MR morphology of cartilaginous endplate correlates with disc degeneration in the lumbar spine. *Eur Spine J [Internet]*. 2023;32:2358–67. <https://doi.org/10.1007/s00586-023-07739-9>.
34. Fields AJ, Han M, Krug R, Lotz JC. Cartilaginous end plates: quantitative MR imaging with very short echo times-orientation dependence and correlation with biochemical composition. *Radiology*. 2015;274:482–9.
35. Bonnheim NB, Wang L, Lazar AA, Chachad R, Zhou J, Guo X, et al. Deep-learning-based biomarker of spinal cartilage endplate health using ultra-short echo time magnetic resonance imaging. *Quant Imaging Med Surg*. 2023;0:0–0.
36. Lotz JC, Fields AJ, Liebenberg EC. The role of the vertebral end plate in low back pain. *Glob Spine J*. 2013;3:153–63.
37. Ishihara H, McNally DS, Urban JPG, Hall AC. Effects of hydrostatic pressure on matrix synthesis in different regions of the intervertebral disk. *J Appl Physiol*. 1996;80:839–46.
38. Walsh AJL, Lotz JC. Biological response of the intervertebral disc to dynamic loading. *J Biomech*. 2004;37:329–37.
39. Baron CA, Dwork N, Pauly JM, Nishimura DG. Rapid compressed sensing reconstruction of 3D non-Cartesian MRI. *Magn Reson Med*. 2018;79:2685–92.
40. Athertya JS, Ma Y, Masoud Afsahi A, Lombardi AF, Moazamian D, Jerban S, et al. Accelerated quantitative 3D UTE-cones imaging using compressed sensing. *Sensors*. 2022;22:7459. <https://doi.org/10.3390/s22197459>.
41. Ma YJ, Searleman AC, Jang H, Wong J, Chang EY, Corey-Bloom J, et al. Whole-brain myelin imaging using 3d double-echo sliding inversion recovery ultrashort echo time (DESIRE UTE) MRI. *Radiology*. 2020;294:362–74.

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.