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Fast in vivo ²³Na imaging and T₂* mapping using accelerated 2D-FID UTE magnetic resonance spectroscopic imaging at 3 T: Proof of concept and reliability study

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Abstract

Purpose: To implement an accelerated MR-acquisition method allowing to map T_2^* relaxation and absolute concentration of sodium within skeletal muscles at 3T.

Methods: A fast-UTE-2D density-weighted concentric-ring-trajectory ²³Na-MRSI technique was used to acquire 64 time-points of FID with a spectral bandwidth of 312.5 Hz with an in-plane resolution of $2.5 \times 2.5 \text{ mm}^2$ in about 15 minutes. The fast-relaxing ²³Na signal was localized with a single-shot, inversion-recovery based, non-echo (SIRENE) OVS method. The sequence was verified using simulation and phantom studies before implementing it in human-calf muscles. To evaluate the 2D-SIRENE-MRSI (UTE = 0.55 ms) imaging performance, it was compared to a 3D-MRI (UTE = 0.3 ms) sequence. Both datasets were acquired within two same-day sessions to assess repeatability. The T₂* values were fitted voxel-by-voxel using a bi-exponential model for the 2D-MRSI data. Finally, intra-subject coefficients of variation (CV) were estimated.

Results: The MRSI-FID data allowed us to map the fast and slow components of T_2^* in the calf muscles. The spatial-distributions of ²³Na concentration for both MRSI and 3D-MRI acquisitions

SUPPORTING INFORMATION

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were significantly correlated (P<0.001). The test-retest analysis rendered high repeatability for MRSI with a CV of 5%. The mean T_{2Fast}^* in muscles was 0.7 ± 0.1 ms (contribution fraction = 37%), while T_{2Slow}^* was 13.2 ± 0.2 ms (63%). The mean absolute muscle ²³Na concentration calculated from the T_2^* -corrected data was 28.6 ± 3.3 mM.

Conclusion: The proposed MRSI technique is a reliable technique to map sodium's absolute concentration and T_2^* within a clinically acceptable scan time at 3T.

Keywords

sodium; MRSI; fast; T₂ mapping; OVS localization; skeletal muscles; quantification

1 | INTRODUCTION

Sodium (²³Na) plays a crucial role in preserving many vital functions in our bodies. Under healthy conditions, ²³Na concentration stays within certain ranges but varies among tissue types.¹ The alterations in these ranges might be a sign of health disorder or physiological changes. In skeletal muscles, the increase in ²³Na concentration was used as a biochemical marker for several diseases such as hypertension², diabetes,³ or muscular channelopathies.^{4,5} It was also used for monitoring early therapeutic responses.⁶ The physiological changes due to exercise have also been observed by using ²³Na-MR. An increase in muscle sodium concentration has been noticed after the exercises.⁷ In a similar study, this has been suggested to be a result of an increase in the transverse relaxation time (T₂).⁸

²³Na possesses 3/2 nuclear spin and its relaxation is determined by the quadrupolar interaction (QI = the interaction between the 23 Na nuclear quadrupole moment and the local electric field gradients), and the containing environment.^{9,10} In an aqueous solution, the time-averaged QI equals to zero, and the spins move freely and tumble very fast. Subsequently, ²³Na exhibits a relatively long monoexponential T₂. However, within a biological system, ²³Na spins interact with the surrounding macromolecules, and a time averaged QI 0 is possible. As the mobility of ²³Na gets more restricted, the transverse relaxation becomes biexponential.¹⁰ The monoexponential relaxation decay within in-vitro aqueous ²³Na solutions is ranging between 20–50 ms.^{11,12} A monoexponential ²³Na relaxation of 55-65 ms was only reported for the CSF.^{1,13} In human muscles, the fast transverse relaxation time (T^*_{2Fast}) ranges between 0.5–3.0 ms, whereas slow transverse relaxation time (T*_{2Slow}) ranges between 12–28.14 This variation in relaxation modes and magnitude between the in-vivo and in-vitro environments can result in a quantification bias. Similarly, the longitudinal relaxation time (T₁) of 23 Na is short. At 3T, the T₁ of 23 Na in skeletal muscles has been estimated to be about 29 ms.⁴ Thus, a careful relaxation measurement is necessary to conduct accurate quantification studies yielding useful physiological information.

Sodium T_2^* mapping and absolute quantification suffer from several challenges originating from lower intrinsic SNR and shorter T_2^* relaxation values compared with ¹H.^{1,14} Thus, to overcome these issues, measurements at ultra-short TE (UTE), increasing the voxel size, taking more averages, and/or scan with ultra-high magnetic fields > 3T have been proposed.

 $^{7,13,15-22}$ Increasing voxel dimensions will be at the expense of large partial-volume artifacts. Alternatively, taking more averages will lengthen the scan duration. 3D MRI methods have been the method of choice to increase SNR and reduce the echo time since it does not require any slice selection. In addition, estimation of the relaxation times and correcting for relaxation effects can further improve the quantitative accuracy. However, this requires acquiring multiple-echo data to perform T₂* with a cost of prolonged acquisition time.

While the 3D-MRI technique is the common method to capture the T*2Fast, conducting relaxometry studies require impractical acquisition scan duration.^{13,15-20} The used ²³Narelaxometry techniques suffer either from a limitation in their spatial precision (large voxel covering different tissue types) or from their impractical acquisition time. The T₂* values of intact skeletal muscles have been measured previously using non-localized FID methods.^{8,21} Such relaxometry methods estimate the averaged value from a large scanned area, which may include different tissues in addition to the muscles. To conduct the relaxometry study with a better spatial resolution, a 3D-UTE acquisition technique has been used with images acquired at different echo times, which consumes a long scan time (9 min per echo), and with a coarse in-plane spatial resolution $(6 \times 6 \text{ mm}^2)$.⁷ Alternatively, a multi-echo GRE sequence has been used to measure the T_2^* over the entire slice of interest with higher resolution and shorter total time (~14 minutes).²² However, since GRE starts with a sliceselective gradient, the minimum possible first echo was 1.9 ms, which may not be short enough to detect the fast decaying component of the T_2^* . To allow shorter TEs and avoid the lengthy slice-selective gradients, outer volume suppression (OVS) methods such as singleshot, inversion-recovery based, non-echo (SIRENE)²³ and FIDLOVS²⁴ have been proposed. However, these methods have not been applied and tested for ²³Na imaging.

Therefore, the goal of this work was to develop an accelerated method with an early acquisition start to estimate in vivo T_2^* values of muscle tissues in a voxel-wise manner at 3T while maintaining high spatial and temporal resolution. To attain this, we are proposing an accelerated density-weighted concentric ring trajectory (DW-CRT) MRSI acquisition to measure in vivo ²³Na relaxation times in the lower leg muscles with a high sampling frequency. Additionally, to mitigate the long TE limitation when using slice-selective gradients, the SIRENE method²³ was used instead to reach UTE. Before conducting an invivo repeatability study, the localization method was tested using simulation and in-vitro experiments. Additionally, to validate the imaging ability of our proposed MRSI, we will compare its results to images acquired by a state-of-the-art UTE-3D-MRI sequence. This comparison would also help us to evaluate the importance of performing T_2^* correction.

2 | METHODS

2.1 | Sequence design

The sequence begins with a pair of slice-selective gradients and adiabatic full passage inversion pulses (hyperbolic secant (HS20) pulses, B_1 peak of 19.2 µT, a pulse duration of 12.8 ms, and a thickness of 100 mm) to invert the magnetization bands outside the desired slice of interest (SOI) along the z direction and followed by spoiler gradients. After a delay

chosen to null the inverted magnetization (TI = $ln[2] \times tissue's T_1$, 20 ms in our case), pairs of wide-bandwidth adiabatic half passage OVS saturation pulses (HS20, B₁ peak of 29 µT, a pulse duration of 2.56 ms, and thickness of 100 mm) and gradients along z-axis were applied. The net effect is to eliminate the magnetization within the bands at each side of the SOI. The FID is then measured from the remaining magnetization within the SOI after a square excitation pulse with a duration of 240 µs (Figure 1). To accelerate data collection, the k-space data is collected by using a fast density-weighted concentric ring trajectory (DW-CRT) acquisition similar to that implemented in Chiew et al.²⁵

2.2 | Outer volume suppression bands performance evaluation

2.2.1 | **Simulation**—The performance of the OVS bands for eliminating the entire ²³Na signal outside the SOI was evaluated by simulation using SpinBench software (HeartVista, Inc. Menlo Park, CA). The identical MRSI experimental parameters were used for the simulation software together with the approximated skeletal muscle sodium NMR properties at 3T (gyromagnetic ratio = 11.25 MHz/T, $T_1 = 29$ ms, fast $T_2^* = 0.5$ ms, and slow $T_2^* = 12$ ms). The resulted thickness, sharpness, and residual signal amplitude over the generated spatial profile were used to judge and optimize the bands' parameters (Figure 2A).

2.2.2 | **Phantom experiment**—To test the optimized OVS bands parameters on the scanner, a phantom study was conducted. Four phantoms (bottles) were prepared with different known 23 Na concentrations (10, 20, 30, 40 mM), which will also serve as a calibration reference for the in vivo signal quantification. To achieve the in-vivo T₁ of sodium, 2.9 g/L CuSO₄ was added to each phantom.

To ensure that no signal is coming from outside the SOI, a scan was acquired while two ²³Na phantoms of the highest concentration (30 and 40 mM) were placed outside the SOI, within the OVS bands. The remaining phantoms of lower concentration (10 and 20 mM) were placed in the center of the SOI (Figure 2B). For comparison, ²³Na images with a 3D-MRI scan covering the same SOI were also acquired.

2.3 | In vivo experiment

2.3.1 | Human subjects—In vivo calf muscle scans were performed in four subjects [1 male and 3 females; age 22–40 years (median = 26 years); body mass index (BMI) = 24 ± 3 kg/m²]. The study was conducted in accordance with the institutional review board of Purdue University. Before being scanned, an informed written consent was obtained from all the subjects.

2.3.2 | **Repeatability study**—In order to evaluate the reliability of the proposed MRSI method, test-retest scans were performed. The subjects were asked to lie in a feet-first supine position with their left leg within a 23 Na coil. The maximum circumference of their lower leg was positioned to be centered in the middle of the coil. Additionally, to make sure that both scans were acquired from the same slice, an ink marker was used to draw a line on the leg region (~10.5–11.5 cm below the knee joint, based on the subject) where the scanner laser marker was centered for the first scan. After a short break (about 5 minutes) outside of the scanner, the repeat scan was acquired using the same scanning protocol.

2.3.3 | **Scanning parameters**—The data were collected using a 3T Siemens Prisma MR system (Siemens Healthineers, Germany) and a frequency-tuned mono-resonant ²³Na-transmit/receive birdcage knee coil (32.6 MHz, Stark-Contrast, Erlangen, Germany).

The ²³Na data were measured through the default shim currents of the MR system ("Tune Up" under shim settings).

For the ²³Na-MRSI study, inversion pulses with TI of 20 ms were applied and followed by the two OVS pulses on the z-direction before the excitation RF pulse. The FID DW-CRT MRSI was implemented with an alpha of 1. The alpha parameter controls the spatial response function, and alpha = 1 was found to be providing a higher SNR with a cost of degrading the spatial response function, as described in Chiew et al.²⁵. The following parameters were used for the DW-CRT MRSI: matrix size = 96×96 , field of view (FOV) = 240×240 mm², resolution = $2.5 \times 2.5 \times 20$ mm³ (nominal resolution = 0.125 mL), flip angle (FA) = 90° , acquisition delay (time from the center of excitation pulse to the first FID time point) = 0.55 ms, repetition time (TR) = 650 ms, temporal samples = 64, number of rings = 48, points-per-ring = 64, spectral bandwidth = 312.5 Hz, spatial interleaves = 4, and readout duration = 204.8 ms. The scan performed with 8 averages, which resulted in a total acquisition time (TA) of 15 minutes and 24 seconds.

To evaluate the spatial mapping performance of the ²³Na-MRSI technique, ²³Na-MRI data were collected with a density-adapted 3D radial acquisition sequence²⁶ with UTE = 0.3 ms, TR = 100 ms, FA = 90°, 1 average, $4 \times 4 \times 4$ mm³ nominal resolution, and FOV = 256 × 256 mm², and TA = 6.66 min. Five axial slices were averaged to cover the same MRSI slice. These 3D-UTE MRI images will serve as a reference to assess the spatial distribution of the MRSI maps. It will also be used to compare the quantification results from data acquired with UTE and T₂*-corrected methods.

For segmentation, high-resolution anatomical images were acquired using the integrated body coil with a T₁-FLASH sequence of TR/TE = 250 ms/2.46 ms, FA = 60°, 2 averages, $0.6 \times 0.6 \times 10 \text{ mm}^3$ resolution, and FOV = $200 \times 200 \text{ mm}^2$.

All the above sequences were planned to collect data from the same axial slice placed at the isocenter. Additionally, shimming using the ¹H coil was done before the sodium measurements.

2.3.4 | **Post-processing**—The reconstruction of the MRSI data were performed in MATLAB (MathWorks, Natick, MA, USA). The gridding and the Fast Fourier Transform were done using the Nonuniform FFT (NUFFT) method.²⁷ In addition to the DW-CRT trajectory²⁵, a Hanning filter was applied in k-space for density compensation.

The B₀ inhomogeneity was corrected by calculating the ²³Na B₀ maps as described in Gast et al.²⁸ Here, the ²³Na B₀ maps were calculated based on the first two ²³Na-MRSI phase-unwrapped images ($\theta_{TE_2,unwrapped}$ and $\theta_{TE_1,unwrapped}$) acquired at TE₁ = 0.55 ms, and TE₂

= 3.75 ms, as follows:

$$\Delta B_0 = \frac{\theta_{TE_2, unwrapped} - \theta_{TE_1, unwrapped}}{TE_2 - TE_1}$$
(1)

These B_0 maps were used to reconstruct field-corrected ²³Na-MRSI images using a fast iterative image reconstruction method.²⁹ An example of B_0 map and correction can be found in Figure 3. Finally, we applied low rank approximations for spatial-spectral denoising (2 for space-time, and 5 for Hankel filtering) of the reconstructed ²³Na MRSI data.³⁰ An example of ²³Na-data without and with the low-rank denoising can be found in Supporting Figure S1.

The 3D-MRI data were reconstructed using a MATLAB tool designed for the radial sequence. To reconstruct the 3D-MRI magnitude images, the k-space data were density compensated before being re-gridded with an oversampling ratio of two using a Kaiser-Bessel kernel,³¹ and Fourier transformed by a conventional fast FFT. The data were filtered with a Hanning filter.

2.3.5 | **Fitting of T₂***—The acquired FID data from the leg region were fitted to a biexponential decay with a nonlinear least squares method to calculate the T_{2Fast}^{*} and T_{2Slow}^{*} relaxation time components:

$$SI_{Leg}(t) = S_{Fast} \quad e^{-t/T_2^* Fast} + S_{Slow} \quad e^{-t/T_2^* Slow} + n \tag{2}$$

Here, S_{Fast} and S_{Slow} are the contributions to the initial signal from the fast and slow components, respectively, *t* indicates the FID points collection time, and *n* represents the offset level.

The initial (undecayed) signal equals the sum of S_{Fast} and S_{Slow} . Thus, their contribution fractions were represented as:

$$F_{Fast} = \frac{S_{Fast}}{S_{Fast} + S_{Slow}} \tag{3}$$

$$F_{Slow} = \frac{S_{Slow}}{S_{Fast} + S_{Slow}} \tag{4}$$

For the reference bottles, Ref, the FID curves were fitted to a monoexponential decay:

$$SI_{Ref}(t) = S_{Ref} e^{-t/T_2^2 \text{Ref}} + n$$
⁽⁵⁾

2.3.6 | **Quantification**—The ²³Na concentration maps in mM were reconstructed by calibrating the signals of the reference bottles to their corresponding known concentrations. The resulting signal-to-concentration linear equation was used to calibrate the signals within the leg and to estimate the concentration in mM.

2.3.7 | **Muscle segmentation**—The high-resolution T1-weighted image was used to manually draw regions of interest (ROIs) over each of the seven main large muscles (Figure 4). The borders of these ROIs were determined by tracing the boundaries of their corresponding muscle. The ROIs were then down-sampled and co-registered to each sodium map to evaluate the ²³Na spatial distribution. Voxels close to the main blood vessels were avoided. The subcutaneous fat region was segmented similarly.

2.3.8 | **Statistical analysis**—The Spearman regression analysis was performed to assess the signal spatial distribution agreement between the two used acquisition techniques within the segmented ROIs. The MRSI signals were extrapolated to their estimated values at 0.3 ms to match the TE of 3D-MRI.

To evaluate the repeatability of the MRSI and 3D-MRI methods, coefficient of variance (CV), intraclass correlation coefficient (ICC), and Bland–Altman analyses were utilized.

3 | RESULTS

3.1 | Outer volume suppression bands performance evaluation

The result of the simulation (Figure 2A) showed that using a TI of 20 ms provides a sharp suppression profile. The spatial profile in Figure 2A represents the ²³Na magnetization after the 90° excitation. While the signal within the OVS bands is totally suppressed, the magnetization within the SOI has full transverse magnitude in a slice thickness of 20 mm.

In-line with the simulation, the phantom study has demonstrated that the signals from the high ²³Na concentration phantoms (30 and 40 mM) were totally suppressed by the OVS bands (Figure 2B). Thus, the signals of the low concentration phantoms (10 and 20 mM) within the SOI do not have any contamination and are in agreement with the 3D-MRI sequence (Figure 2B).

3.2 | In vivo experiment

The results of the biexponential T_2^* values (T_{2Fast}^* and T_{2Slow}^*) and their fractions are summarized in Table 1 together with the T_2^* -corrected concentration within each ROI. Fast and slow T_2^* -maps and their corresponding signal fraction maps are illustrated in Figure 5. The T_2^* correction was demonstrated by showing sodium maps before and after correction with their difference (Figure 6A–C). Additionally, Figure 6D shows an example of two signals with similar initial MR signals diverging with time due to their different transverse relaxation in different environments. The ²³Na spins decay monoexponentially within the reference phantom (blue) and biexponentially within the muscle (green).

As shown in Figure 7, the MRSI quantitative data was compared to the UTE 3D-MRI at three conditions: 1- the MRSI data at 0.55 ms, 2- extrapolated values to the 3D-MRI TE of 0.3 ms, and 3- at 0 ms using the estimated T_2^* values. Before T_2^* correction, the comparison slope between the 3D-MRI and MRSI was 0.88, which increased to 1 at 0.3 ms. After the T_2^* correction, the comparison slope increased to 1.27. For all conditions, the regression analysis resulted in significant correlation (P < 0.001).

A representative set of baseline and repeated scan maps of the MRSI ²³Na concentration (before and after T_2^* correction) and their corresponding 3D-MRI maps are provided in Figure 8. The repeatability of the muscle and subcutaneous fat ROIs from all subjects are shown in Figure 9. The CV was calculated as 4.2, 5.2, and 5.9 %, and the ICC as 0.98, 0.95, and 0.89 for the MRSI data before correction, after correction, and the 3D MRI data, respectively.

4 | DISCUSSION

In this work, we have introduced a novel method to acquire ²³Na 2D-FID data with short acquisition delay (0.55 ms) with a good spatial-quality comparable to a well-established 3D acquisition method. The proposed method provides additional time points in the FID to calculate both of T_2^* and B_0 , and correct for their effects without additional scans.

To reduce the delay time while performing a 2D-measurement, we utilized a single-shot, inversion-recovery based, non-echo (SIRENE) method based on OVS bands²³. In this study, we implemented the technique with a novel accelerated k-space trajectory, DW-CRT,²⁵ and showed its feasibility for the ²³Na acquisition. The simulation and phantom studies showed that the proposed technique could be used as an alternative to the slice-selective gradient approach in which achieving an ultra-short TE is difficult. On the other hand, the in-vivo study demonstrated that the proposed MRSI method could be a faster option to provide ²³Na-T₂* maps compared to multi-echo 3D-MRI approaches that usually require lengthy acquisition times.^{7,16–20}

Although MRSI and the 3D-MRI sequences have different acquisition and reconstruction parameters, the estimated ²³Na spatial distributions were in a good agreement with a regression line close to unity and a small bias of 0.67 mM (Figure 7B). This variability between the two sequences results is expected due to their different spatial resolution (MRSI: $2.5 \times 2.5 \text{ mm}^2$; 3D-MRI: $4 \times 4 \text{ mm}^2$), and spatial response function.^{25,26} Moreover, the field inhomogeneity correction was only performed on the MRSI reconstruction. A recent study with a similar 3D sequence, but with double echo to measure B₀ and correct for its inhomogeneity at 3T, showed a small enhancement in the results.³²

In this work, we provided an accelerated technique to conduct voxel-wise T_2^* measurements within an acceptable scan time (15 minutes) while maintaining the spatial resolution. It is worth mentioning that recently a number of studies have shown the feasibility of voxel-wise T_2^* mapping in the brain using UTE-3D acquisition techniques.^{16–20} However, these techniques were conducted with fewer time points (8–38 echoes), and require long acquisition times (26 min - 1 h). Moreover, many of these methods were either done with lower spatial resolution, at a higher magnetic field, or done with both. If the proposed MRSI method was applied under these conditions, further reduction in scan time can be achieved.

The estimated mean muscle relaxation times ($T_{2Fast}^* = 0.7 \pm 0.1$ ms of $F_{Fast} = 37 \pm 2$ %; $T_{2Slow}^* = 13.2 \pm 0.2$ ms of $F_{Slow} = 63 \pm 2$ %) are in line with previously reported values measured using single voxel ²³Na MRS ($T_{2Fast}^* = 0.8 \pm 0.2$ [32 ± 7 %], $T_{2Slow}^* = 12.4 \pm 1.8$ ms [68 ± 7 %]).²¹ The average muscle absolute concentration (after T_2^* correction) was

 28.6 ± 3.3 mM, which is also in agreement with reported biopsy results.⁷ In terms of T₂* spatial distribution, the slow fraction is larger and slower in areas with large blood vessel, which makes sense. In blood vessels, the motion is less restricted compared to muscles and spins move more freely that resulting in longer T₂* values and the time averaged QI is minimal, which means that the fast component fraction is also minimal.

One can conclude at least two important reasons for performing a relaxation study. First, relaxation corrected data provides a better estimation of absolute concentration. In Figure 7, the estimated concentration at 0.3 ms increased by 27% after the T_2^* correction (the comparison slope increased from unity to 1.27). This enhancement in concentration (Figure 6C&7) indicates that T_2^* correction might be useful for more accurate quantification results even with UTE measurements, as T_2^* -corrected data showed a better agreement with biopsy results (see the preceding paragraph). Additionally, one can see that even with the UTE measurement, a relaxation bias might be present due to the difference in decay mode between the reference bottles and muscle tissue (Figure 6D). The second use of T_2^* mapping is to provide extra information that can be utilized to assess potential physiological changes. However, the voxel-wise T_2 correction may result in overestimating ²³Na concentrations in certain regions, where partial volume effect exists. This was seen in this study within areas where the main blood vessels start bifurcating to smaller vessels (see the regions of very large difference in Figure 6C).

According to the repeatability analysis, the proposed method showed high repeatability (CV = 4%, ICC = 0.98). The repeatability CVs and ICCs after the T_2^* correction were close to the 3D MRI results (CV = 5%, ICC = 0.95 vs CV = 6% and ICC = 0.89). Thus, the proposed MRSI might be a potential method to estimate the T2* values and the absolute sodium concentration (free from T_2^* bias) within a reasonable scan time. Since ²³Na signal suffers from low SNR, scanning with large voxels or at high fields is performed to maintain good SNR within a reasonable scan time. Compared to other techniques at 3T, the proposed MRSI method acquires 64 data points with an acceptable spatial resolution (2.5 mm², nominal inplane) and acquisition time (15 minutes and 24 seconds). Since the FID data were collected at different time points with different phases, B₀ correction is also feasible without extra scans. Although scanning at the isocenter reduces the effect of B₀, the proposed approach can be crucial for studies conducted off the isocenter and/or at higher field strengths. Additionally, in this study, we performed the B₀ correction and low-rank filtering in separate steps. Using a combined B₀-corrected low-rank filtering method is expected to improve our results.³³ Moreover, even after B_0 correction, some artifact (shading across the phantoms) is still present. The potential influence of this small artifact on quantification results has been minimized by using the average signal within carefully placed ROIs that covers the entire phantom (without their walls) that reduces any potential spatial variation. This resulted in a calibration line with high linearity and allowed an excellent fitting with R^2 almost = 1 (see example in Supporting Figure S2). Thus, we believe that this artifact does not cause a major quantification issue. Additionally, to ensure a proper biexponential fitting for T_2^* estimation, the prolonged acquisition sampling duration allows for acquisition of the entire FID. This is important for areas of longer T₂* values like reference phantoms.

A limitation of this method is its large dependency on the T_1 value of the scanned tissues. In order for the OVS to perform correctly, the sequence needed to be applied with $TI = ln(2) \times I$ tissue's T_1 (close approximation when TR>> T_1). In this study, we used a TI of 20 ms because muscles have a ²³Na T₁ of 29 ms, as previously measured at 3T.⁴ To suppress tissues with shorter T₁, such as cartilages, the TI has to be reduced, which may be technically challenging. However, one can make sure that no such tissues are within the active region of the receiving coil. For instance, we avoided getting residual signals from the knee by keeping it outside the ²³Na-coil field. Here, we evaluated the technique in healthy muscles. However, T₁ might change with diseases. Although it is expected that this OVS technique would still achieve good suppression even with slightly T1 deviations, future studies might be needed to confirm this. In this study, no B₁ correction was implemented. However, the coil B₁ mapping was performed using a GRE sequence with the double angel method.³⁴ Since the B_1 map was very homogeneous, no B_1 correction was conducted. The 23 Na-coil B₁ map can be found in Supporting Figure S3. Although the sample size may be a limitation in this study, there is no reason to anticipate a large difference in our results with larger sample size.

For this study, we used a volume coil placed along the z-axis of the scanner. Thus, the spatial B_1 variation is expected to increase along the z-direction, where the OVS bands are applied. Together with inversion pulses, OVS bands make the reduction of the longitudinal magnetization depends mainly on the T_1 and is independent of spatial B_1 variations. Additionally, the broad-BW OVS pulses were applied with a B_1 peak of 29 μ T, 2.56 ms duration, and a thickness of 100 mm that resulted in no SAR issues.

The considerable regional variability within T_2^* , their fraction maps, and the increased accuracy of concentration maps demonstrate the potential for future characterization of ²³Na in conditions such as muscle diseases, diabetes, cancers, strokes, cartilage degeneration, and to evaluate physiological interventions. The reduction in scan time will increase the technique's availability and reduce motion artifacts. Technically, while still using a long enough TR (4–5 times T₁), a shorter TR than that used in this study can be used, which will accelerate the acquisition further. In addition to ensuring a full T₁ recovery and maintaining a low SAR, we used a very long TR because we wanted to study the existence of any long decaying species. However, no such decay has been noticed within the scanned area. Thus, we would suggest using a TR of about half of the value used here. Additionally, scanning with shorter TEs, and with higher sampling frequency could enhance the fitting quality further. Beyond ²³Na, the proposed MRSI technique might be useful for ¹H applications such as imaging dental tissues, ³⁵ which characterized by a very fast T_2^* . Measurements of other fast decaying nuclei such as ¹³C, ³¹P, or ¹⁵N may benefit from the proposed technique as well.

5 | CONCLUSIONS

The proposed method allows fast data collection for measuring in vivo sodium concentrations and T_2^* values at high reliability. This may facilitate evaluating pathological and physiological changes related to the ²³Na concentration and T_2^* values within a clinically feasible time, and with a good spatial resolution at 3T.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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FIGURE 1.

Pre-readout MRSI pulse sequence. To suppress the unwanted region outside the slice of interest (SOI), two OVS bands are assigned before the FID excitation. These OVS bands along the z-direction are applied after two selective 180° inversion recovery pulses covering the area outside the SOI at inversion time (TI) = $\ln(2) \times 29$ ms = 20 ms, where 29 ms is the T₁ of muscles.⁴ Directly after the OVS band pulses, a nonselective 90° excitation pulse is applied before starting the FID-MRSI readout



FIGURE 2.

SIRENE-MRSI localization performance evaluation by simulation and phantom studies. A, Simulated data showing a sharp spatial profile along the z-direction when using an inversion time of 20 ms. B, Phantom evaluation of the OVS localization. Two ²³Na phantoms (10 and 20 mM) were placed at the center of the slice of interest (SOI), which were located between the 2 OVS bands. Two additional phantoms of higher concentrations (30 and 40 mM) were placed outside the SOI, within the OVS bands. As shown in the axial images, the MRSI signal is obtained only from the phantoms within the SOI, resulting in an image that is very similar to that produced by the 3D-MRI sequence

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FIGURE 3.

A, An example of a ²³Na-MRSI map without B_0 correction. B, The same ²³Na-MRSI map with B_0 correction. C, The difference in concentration resulted from applying the B_0 correction. D, ²³Na- B_0 map used to correct the ²³Na map



FIGURE 4.

Representative segmented regions of interest (ROIs). Calf muscle and subcutaneous fat (SF) ROIs were drawn based on their high-resolution T1 axial ¹H-image, which allows clear anatomical features for segmentation. Seven ROIs were drawn to cover the main human calf muscles: SOL, Soleus; FBL, Fibularis; EXT, Extensor longus; TA, Tibialis anterior; GM, Gastrocnemius medialis; GL, Gastrocnemius lateralis; and TP, Tibialis posterior muscles. The shown ROIs are presented in the MRSI resolution and overlaid over their corresponding high-resolution T1 anatomical image



FIGURE 5.

Representative relaxation maps. Top panel: fast and slow T_2^* maps, and their corresponding signal fraction (F_{Fast} and F_{Slow}). Their mean values from the entire leg slice (without the bottles) are listed above their maps. Bottom panel: the same maps overlaid on their anatomical images



FIGURE 6.

Illustrations of the importance of the relaxation correction. The difference between the data acquired at 0.55 ms (A) and the data after correction (B), shows a large improvement in concentration estimation. In the difference map (C), about 5 mM difference in muscles was found. (The red star, D) represents the spin density-weighted signal (\approx the initial signal before decay) corresponding to 30 mM absolute concentration. The fitting example of leg voxel (2.5 × 2.5 mm² green box, B and D) and quantification reference voxel (2.5 × 2.5 mm² blue box, B and D) with this absolute concentration shows how their signal can diverge with time before fully decaying. Such diverging can result in T₂*-weighted signals across the FOV with time, leading to underestimated quantification results

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FIGURE 7.

Results of the regression analysis comparing the normalized mean signal (normalized to the phantom signal) of the MRSI and the 3D MRI within 64 regions of interest from all subjects data (7 muscles and one subcutaneous fat ROIs × 4 subjects × 2 scans). A, MRSI data before T_2^* correction (at 0.55 ms) compared to the 3D-MRI data. B, The MRSI data extrapolated (based on the estimated T_2^* values) to represent data at 0.3 ms compared to the 3D-MRI data, which was acquired at TE = 0.3 ms. The regression line almost matches the unity line (slope \approx 1). C, Comparing MRSI data after T_2^* correction to the 3D-MRI data. The increase in slope (from A to C) represents the T_2^* correction impact

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FIGURE 8. Data example of baseline and repeat scans



FIGURE 9.

Evaluation of repeatability by Bland–Altman analysis. A, The results from the MRSI data before T_2^* correction. B, The results from the MRSI data after the T_2^* correction. C, The 3D-MRI data results. The graphs represent the variability of the measured data (muscles and subcutaneous fat ROIs from all subjects, 7 muscles and one subcutaneous fat ROIs × 4 subjects) between 1^{st} and 2^{nd} scans. The coefficient of variation (CV), reproducibility coefficient (RPC), systematic bias, and intraclass correlation coefficient (ICC) values are listed on each plot

TABLE 1

Regions of interest absolute $^{23}\mathrm{Na}$ concentration, $\mathrm{T_2}^*$ values, and signal fractions

ROI	Concentration [mM]	T [*] _{2Fast} [ms]	F _{Fast} [%]	T [*] _{2Slow} [ms]	F _{Slow} [%]
SF	15.7 ± 1.9	0.71 ± 0.04	38.9 ± 1.5	13.2 ± 0.1	61.1 ± 1.5
GM	25.0 ± 2.8	0.67 ± 0.03	38.3 ± 2.1	13.2 ± 0.4	61.7 ± 2.1
TA	25.3 ± 2.1	0.61 ± 0.09	34.9 ± 2.1	12.9 ± 0.4	65.0 ± 2.1
EXT	26.9 ± 3.5	0.62 ± 0.19	35.8 ± 2.8	13.0 ± 0.5	64.2 ± 2.8
GL	28.6 ± 3.1	0.73 ± 0.06	38.6 ± 0.9	13.1 ± 0.2	61.4 ± 0.9
TP	29.5 ± 2.3	0.73 ± 0.21	37.8 ± 3.4	13.3 ± 0.5	62.2 ± 3.4
FBL	31.1 ± 3.4	0.59 ± 0.14	34.8 ± 2.8	13.3 ± 0.6	65.2 ± 2.8
SOL	34.1 ± 2.2	0.76 ± 0.11	38.4 ± 2.8	13.3 ± 0.4	61.6 ± 2.8
Phantom	10 - 40			29.5 ± 0.3	100.0

Abbreviations: SF, Subcutaneous fat; GM, Gastrocnemius medialis muscle; TA, Tibialis anterior muscle; EXT, Extensor longus muscles; GL, Gastrocnemius lateralis muscle; TP, Tibialis posterior muscle; FBL, Fibularis muscles; SOL, Soleus muscle. *F_{Fast}*. The fraction of fast decaying signal; *F_{Slow}*, The fraction of slow decaying signal.