Respiratory Motion-Registered Isotropic Whole-Heart T₂ Mapping in Patients With Acute Non-ischemic Myocardial Injury

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Background: T₂ mapping is a magnetic resonance imaging technique that can be used to detect myocardial edema and inflammation. However, the focal nature of myocardial inflammation may render conventional 2D approaches suboptimal and make whole-heart isotropic 3D mapping desirable. While self-navigated 3D radial T₂ mapping has been demonstrated to work well at a magnetic field strength of 3T, it results in too noisy maps at 1.5T. We therefore implemented a novel respiratory motion-resolved compressed-sensing reconstruction in order to improve the 3D T₂ mapping precision and accuracy at 1.5T, and tested this in a heterogeneous patient cohort.

Materials and Methods: Nine healthy volunteers and 25 consecutive patients with suspected acute non-ischemic myocardial injury (sarcoidosis, n = 19; systemic sclerosis, n = 2; acute graft rejection, n = 2, and myocarditis, n = 2) were included. The free-breathing T₂ maps were acquired as three ECG-triggered T₂-prepared 3D radial volumes. A respiratory motion-resolved reconstruction was followed by image registration of the respiratory states and pixel-wise T₂ mapping. The resulting 3D maps were compared to routine 2D T₂ maps. The T₂ values of segments with and without late gadolinium enhancement (LGE) were compared in patients.

Results: In the healthy volunteers, the myocardial T₂ values obtained with the 2D and 3D techniques were similar (45.8 ± 1.8 vs. 46.8 ± 2.9 ms, respectively; P = 0.33). Conversely, in patients, T₂ values did differ between 2D (46.7 ± 3.6 ms) and 3D techniques (50.1 ± 4.2 ms, P = 0.004). Moreover, with the 2D technique, T₂ values of the LGE-positive segments were similar to those of the LGE-negative segments (T₂LGE− = 46.2 ± 3.7 vs. T₂LGE+ = 47.6 ± 4.1 ms; P = 0.49), whereas the 3D technique did show a significant difference (T₂LGE− = 49.3 ± 6.7 vs. T₂LGE+ = 52.6 ± 8.7 ms, P = 0.006).

Conclusion: Respiratory motion-registered 3D radial imaging at 1.5T led to accurate isotropic 3D whole-heart T₂ maps, both in the healthy volunteers and in a small patient cohort.
BACKGROUND

The T2 relaxation time is one of the physiology-dependent properties of a tissue in a magnetic field that governs the image contrast in magnetic resonance imaging (MRI). In the myocardium, it increases in the presence of edema, which makes the T2 relaxation time a useful indicator of acute myocardial injury irrespective of its etiology (e.g., inflammatory, toxic, or ischemic) (1). T2 mapping, i.e., quantifying the T2 relaxation time in every pixel, has therefore seen increased use for the diagnosis of acute myocardial injury in recent years (2). T2 mapping has been shown to be an effective complementary tool in inflammatory diseases such as myocarditis (3), systemic sclerosis (4), and sarcoidosis (5).

Most current T2 mapping techniques (6, 7) involve the acquisition of several thick 2D slices of the left-ventricular myocardium, which is largely adequate in the case of diseases that affect the entire myocardium or that have a well-defined pattern, such as acute ischemic myocardial injury. However, in a spectrum of inflammatory myocardial injuries such as viral myocarditis and sarcoidosis, the inflammation pattern is essentially irregular and unpredictable, despite typical patterns of segmental and transmural distribution. This may render the standard 2D T2 mapping technique suboptimal, since the disease foci can be missed due to insufficient coverage. Moreover, the thick slices may mask the disease foci through partial volume effect by including healthy and injured tissue in the same voxels. On the other hand, scanning would need to be prolonged to a large series of breath holds to cover the entire myocardium. To address these challenges, free-breathing high-resolution 3D T2 mapping techniques have been proposed in recent years (8–11), and have for example been applied in patients with graft rejection (12) and myocarditis (13).

Among these techniques, T2 mapping based on a self-navigated 3D radial acquisition (14, 15) can make use of the intrinsic robustness of 3D radial imaging against undersampling and motion artifacts, but it faces a challenge in its low effective signal-to-noise ratio (SNR), which leads to a loss in precision of the T2 maps. This 3D radial T2 mapping has therefore mostly been applied at a magnetic field strength of 3T (8, 12), and not at 1.5T, which may be more commonly used for cardiac magnetic resonance (CMR). However, recently several new techniques have been developed that can be used to increase the precision of a 3D radial T2 mapping, including resolving the motion instead of correcting it (16) in order to reduce motion streaking artifacts, and using compressed sensing (17) in order to reduce undersampling artifacts and to denoise the source images.

In this study, we therefore aimed to enable 3D radial T2 mapping at 1.5T by improving the image reconstruction, and to demonstrate the efficacy of this reconstruction method in healthy volunteers as well as in a small cohort of patients with suspected acute non-ischemic myocardial injury. To this end, the T2 maps were generated by first reconstructing respiratory motion-resolved source images, which were then registered to one another to decrease noise and motion artifacts, and thus to improve 3D T2 mapping precision and accuracy at 1.5T. These 3D T2 maps were then compared to routine 2D maps acquired in the same subjects.

MATERIALS AND METHODS

Study Participants

This study was approved by the Institutional Review Board of the Medical University of Gdansk (#NKBBN/72/2019). All participants provided written informed consent prior to the procedure and none of them had contraindications for MRI.

TABLE 1 | Subject characteristics.

<table>
<thead>
<tr>
<th></th>
<th>Healthy volunteers</th>
<th>Patients</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>(n = 9)</td>
<td>(n = 25)</td>
</tr>
<tr>
<td>Age, mean (SD) [years]</td>
<td>43(7)</td>
<td>49(11)</td>
</tr>
<tr>
<td>Gender, F (%)</td>
<td>5 (56%)</td>
<td>9 (36%)</td>
</tr>
<tr>
<td>Heart rate, mean (SD) [bpm]</td>
<td>68 (11)</td>
<td>67 (13)</td>
</tr>
<tr>
<td>BMI, mean (SD) [kg/m2]</td>
<td>24.7 (1.0)</td>
<td>27.2 (4.2)</td>
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<tr>
<td>Hypertension, n(%)</td>
<td>0</td>
<td>15 (60)</td>
</tr>
<tr>
<td>Diabetes, n(%)</td>
<td>0</td>
<td>3 (12)</td>
</tr>
<tr>
<td>Hyperlipidemia, n(%)</td>
<td>0</td>
<td>6 (24)</td>
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<tr>
<td>CAD, n(%)</td>
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<td>3 (12)</td>
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<tr>
<td>LV end-diastolic volume index, mean (SD) [ml/m2]</td>
<td>71(10)</td>
<td>85(30)</td>
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<tr>
<td>LV end-systolic volume index, mean (SD) [ml/m2]</td>
<td>26(7)</td>
<td>39(23)</td>
</tr>
<tr>
<td>LV ejection fraction, mean (SD) [%]</td>
<td>65(5)</td>
<td>55(9)</td>
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<tr>
<td>LV mass index, mean (SD) [g/m2]</td>
<td>59(8)</td>
<td>72(18)</td>
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<tr>
<td>Referral diagnosis, n(%)</td>
<td>Sarcoïdosis</td>
<td>NA (76)</td>
</tr>
<tr>
<td></td>
<td>Systemic sclerosis</td>
<td>NA (28)</td>
</tr>
<tr>
<td></td>
<td>Acute graft rejection</td>
<td>NA (28)</td>
</tr>
<tr>
<td></td>
<td>Myocarditis</td>
<td>NA (28)</td>
</tr>
</tbody>
</table>

BMI, body mass index; CAD, coronary artery disease; LV, left ventricular; SD, standard deviation.
To study the baseline relaxation times, healthy volunteers (*n* = 9, age = 43 ± 7 y, 5 (56%) women, Table 1) without any history or symptoms of cardiovascular disease were recruited.

Consecutive patients with suspected acute non-ischemic myocardial injury (*n* = 25, 19 cardiac sarcoidosis, 2 acute graft rejection, 2 systemic sclerosis, 2 myocarditis; 9 (36%) women, age 49 ± 10 y; Table 1) were recruited.

**MR Acquisition**

All MR scanning was performed on a 1.5T clinical scanner (MAGNETOM Aera, Siemens Healthcare, Erlangen, Germany). All participants underwent routine bSSFP cine imaging to assess cardiac function (18), routine breath-held 2D T2 mapping (19), and the prototype free-breathing 3D T2 mapping. In addition, the patients underwent routine late gadolinium enhancement (LGE).
imaging 7–15 min after injection of 0.1 mmol/kg of gadobutrol (Gadovist, Bayer AG, Leverkusen, Germany).

The routine $T_2$ maps were acquired as ECG-triggered Cartesian 2D $T_2$-prepared bSSFP images (6) with repetition time $TR = 2.5$ ms, echo time $TE = 1.1$ ms, flip angle $= 70^\circ$, pixel bandwidth $= 1.18$ Hz/px, field of view $= 360 \times 288$ mm$^2$, slice thickness $= 8$ mm, acquired pixel size $= 2.49 \times 1.88$ mm$^2$ interpolated to $1.88 \times 1.88$ mm$^2$, $T_2$ prep duration $= 0/25/55$ ms, breath-hold duration nine heartbeats (data acquired every three heartbeats), and GRAPPA acceleration factor 2. Images were acquired in a short-axis (SAX) orientation at the basal and mid-ventricular level. Since the reconstructed 2D maps were immediately available on the scanner, visibly corrupted maps were re-acquired as per routine protocol.

The free-breathing $T_2$ maps were acquired as three ECG-triggered 3D radial bSSFP volumes with a phyllotaxis trajectory (14), $TR = 2.6$ ms, $TE = 1.3$ ms, flip angle $= 35^\circ$, pixel bandwidth $= 908$ Hz/px, field of view $= (220)$ mm$^3$, isotropic voxel size $= 1.6$ mm$^3$, $T_2$ prep duration $= 0/30/60$ ms, and interleaves of 49 k-space lines acquired every other heartbeat preceded by a superior-inferior line that could be used for self-navigation (15). This resulted in a total acquisition time of 112 interleaves $\times$ 3 $T_2$preps $\times$ 2 heartbeats/interleave $= 672$ heartbeats, or 11.2 min at 60 bpm.

Blokch equation simulations of the abovementioned proposed pulse sequence were performed for a heart rate range from 40 to 90 bpm with an assumed myocardial $T_1$ relaxation of 1,050 ms and a true $T_2$ of 50 ms to assess the influence of the heart rate on the estimated $T_2$ relaxation time.

$T_2$ Map Reconstruction

The routine 2D $T_2$ maps were reconstructed on the scanner (Siemens IDEA, Erlangen, Germany): the source images were non-rigidly registered and a pixel-wise $T_2$ fit was performed with the standard two-parameter exponential decay without offset, which resulted in maps in the DICOM format.

The respiratory motion-resolved reconstruction of 3D radial volumes (Supplementary Figure 1) was performed in MATLAB (the Mathworks, Natick, USA) on a workstation equipped with two Intel Xeon CPUs, 512 GB of RAM, and an NVIDIA Tesla K40 GPU. Here, a principal component analysis (PCA) was performed on the superior-inferior profiles in order to partition the dataset into four different respiratory states. 4D (x-y-z-respiratory dimensions) images were then reconstructed with a parallel imaging and compressed sensing algorithm that exploits sparsity along the respiratory dimension (16, 21, 22), resulting in separate images for all $T_2$ preparation times and respiratory states. The compressed sensing optimization problem was solved with the conjugate gradient technique (17) using the finite difference operator as a sparsifying transform over the respiratory dimension with a weight $\lambda$. All respiratory bins were translationally and then non-rigidly registered to the end-expiration bin with Elastix (23), and were subsequently averaged in order to increase the SNR of each $T_2$-prepared volume. After a second, similar, registration of the resulting three averaged $T_2$-prepared volumes, voxel-wise $T_2$ mapping with an offset factor to account for $T_1$ recovery (24–26) was performed. Since the motion is no longer resolved after these registrations, we named the resulting 3D maps “motion-registered” $T_2$ maps. The total reconstruction time from raw data to $T_2$ map was recorded.

The regularization weight was optimized by comparing $T_2$ map sharpness in maps reconstructed with $\lambda = 0.01$, $\lambda = 0.05$ (the commonly used value at this spatial resolution), and $\lambda = 0.25$ in a subgroup of $n = 6$ patients. $T_2$ map sharpness was assessed by fitting a line from the middle of the septal myocardium to the left-ventricular blood pool with a parametrized sigmoid function ($T_2$ $= a/(1+e^{-k(b+c)})+c$), where $a$, $b$, and $c$ are scaling variables and $k$ is the sharpness (in px$^{-1}$ or mm$^{-1}$; higher is better) (27). The sharpness assessment was repeated for a total of five adjacent lines and the average sharpness $k$ was reported. To ascertain that there is no significant difference between the proposed map reconstruction and alternatively first mapping the $T_2$ relaxation time in each respiratory-resolved bin and then averaging these four bins, the $T_2$ map sharpness was also quantified in this alternative reconstruction in these $n = 6$ patients.

In order to visually demonstrate that a self-navigated reconstruction (15) results in non-diagnostic maps at 1.5T due to too low SNR, a self-navigated reconstruction was performed in a single healthy volunteer, since the resulting maps were often too noisy for segmentation and quantitative analysis. This reconstruction was made with the same 3D radial data mentioned above. Here, the 1D displacement of the left-ventricular blood pool along the superior-inferior readouts acquired at the start of each interleave was used to correct each interleave for respiratory motion in k-space prior to image reconstruction (15, 28). The resulting three 3D images were translationally and then non-rigidly registered with Elastix, and voxel-wise $T_2$ mapping (8) was performed.
FIGURE 3 | Segmental analysis of the myocardial $T_2$ values in the healthy volunteers. Basal and mid-ventricular segmental $T_2$ values and CoVs are shown according to the standard AHA segmentation. (A,B) The segmental $T_2$ values obtained with the 2D and 3D techniques are highly similar. (C,D) The CoV is consistently lower for the 2D technique than for the 3D technique. Significantly different CoV values between the respective segments are marked in red.

Map Analysis and Statistics

The visible myocardium in the routine 2D maps and their matching single slices in the 3D volumes were segmented in MATLAB. The $T_2$ values of the entire visible myocardium and the regional segments defined by the American Heart Association (AHA) were then measured in all volunteers and patients by two independent observers (JF and AS, with 7 and 10 years of experience with cardiac MRI, respectively). For the whole myocardium and 12 out of 16 AHA segments of each subject (2D apical segments were not included as they are generally considered prone to partial volume effects that may compromise the measurement accuracy) the $T_2$ values, coefficients of variation (CoV, the standard deviation divided by the average), and the inter-subject standard deviation obtained with the two techniques were calculated. Since segmental values can strongly vary both due to difference in local disease patterns and precision of the technique, segmental $T_2$ values were only directly compared in the healthy volunteers. In the patients, the segments that were LGE-positive were grouped for comparison with the LGE-negative segments. These values were then compared between the 2D and 3D techniques with paired Student's $t$-tests with a Bonferroni correction.
**FIGURE 4** Motion-registered 3D $T_2$ maps of the heart of three patients. (A–C) Routine $T_2$-prepared 2D bSSFP short-axis (SAX) $T_2$ maps. (D–F) Matching single slices from the motion-registered 3D $T_2$ maps; $T_2$ values closely match those of the routine maps. (G–L) Perpendicular long-axis (LAX) maps from the same 3D datasets as in (D–F). Patient 1 (female, 37 y.o.) and patient 2 (male, 44 y.o.) had myocarditis, while patient 3 (male, 65 y.o.) had sarcoidosis. Local minor $T_2$ fluctuation can be observed in all three orientations of the 3D $T_2$ maps and all three patients, and might be caused by small misregistrations, as well as by local $T_2$ variations that show up in the 1.6 mm-thick 3D slice, but are averaged out in the 8 mm-thick 2D slice. However, global $T_2$ values did not significantly differ in any of the patients. The color bar shows the $T_2$ relaxation time in ms.

when appropriate, with $P < 0.05$ considered statistically significant when two quantities were compared. When multiple quantities were compared to one another a one-way ANOVA with a post-hoc Tukey correction for multiple comparisons was used.

The total number of segments that was not considered of diagnostic quality (i.e., not clearly defined or too thin for segmentation) by an experienced reader (KD) was counted for the 3D technique; this analysis could not be performed for the 2D technique, since visually corrupted maps were re-acquired, resulting in analyzable 2D $T_2$ maps in all study subjects. The visual quality of the maps was assessed by two experienced CMR specialists (KD, AG) on a continuous scale from 1 to 10 with a visual analogue (30).

Bland-Altman analyses were performed to assess the $T_2$ differences between the 2D and 3D techniques in the patients, as well as to assess the inter- and intra-observer agreement for the 3D technique. Trends in these Bland-Altman plots were tested for significance with Spearman’s rank correlation.

**RESULTS**

The respiratory motion-resolved reconstruction resulted in visibly well-separated motion states in the source images (Supplementary Animated Figure 1), while motion-registered isotropic 3D $T_2$ maps of the heart were successfully obtained in all subjects (Figure 1). Several features in these 3D maps were visually more blurred than their equivalents in the 2D maps. The self-navigated reconstruction of the source images did not lead to diagnostic maps (Figures 1E–G). The total reconstruction time from raw data to $T_2$ map was 1 h 19 ± 2 min.

The sharpness measurements in the patient subgroup resulted in $k = 1.56 ± 0.22$ mm$^{-1}$ for the proposed technique, i.e., respiratory-registered $T_2$ maps with $\lambda =$
Bland-Altman analyses of the 2D vs. the 3D T$_2$ mapping technique in the patients. The bias is indicated with a solid line, while the upper and lower limits of the 95% confidence interval are indicated with a dotted line. (A) The comparison of the 2D and 3D techniques shows a small bias, but no significant trend (P = 0.11 for a linear correlation). (B) The interobserver comparison for the routine 2D technique indicates a very small bias and small confidence interval. (C) The interobserver comparison of the 3D technique has a similarly small bias and a slightly larger confidence interval than the 2D technique. Neither interobserver plot shows a significant trend (P > 0.21). The intra-observer analysis performed in a subset of patients (n = 15) showed no bias or trend for the 2D (D) and 3D (E) techniques, although the confidence interval for the 3D technique was twice as large as that of the 2D technique.

0.05 (Supplementary Figure 2). The motion-registered mapping with $\lambda = 0.01$ and $\lambda = 0.25$ resulted in $k = 1.67 \pm 0.34$ mm$^{-1}$ (P = 0.32) and $k = 1.49 \pm 0.29$ mm$^{-1}$ (P = 0.34), respectively. Mapping each respiratory-resolved bin first and then averaging these maps as an alternative reconstruction resulted in $k = 1.63 \pm 0.29$ mm$^{-1}$ (P = 0.40) for the motion-registered T$_2$ mapping.
Conversely, the entire patient group resulted in $k = 1.71 \pm 0.29$ mm$^{-1}$ and $k = 1.89 \pm 0.40$ mm$^{-1}$ ($P = 0.038$) for the 3D and 2D techniques respectively, demonstrating the higher sharpness in the 2D technique despite the larger pixel size. This also held true in the healthy volunteers at $k = 1.71 \pm 0.31$ vs. $k = 1.98 \pm 0.38$ mm$^{-1}$ ($P = 0.037$).

In the healthy volunteers, the myocardial T$_2$ values obtained with the 2D and 3D techniques were highly similar at $45.8 \pm 1.8$ and $46.8 \pm 2.9$ ms ($P = 0.33$, Figure 2), respectively, while the CoV was lower in the 2D technique at $4.5 \pm 0.8$ vs. $8.2 \pm 1.5\%$ for the 3D technique ($P \leq 0.001$). The segmental T$_2$ values did not significantly differ between the two techniques ($P \geq 0.09$ for all, Figures 3A,B), while the CoV differed in 4 out of 12 segments (Figures 3C,D). Out of 108 analyzed segments in the 3D T$_2$ maps in healthy volunteers, 6 (5.6\%, of which 4 [3.7\%] in one subject) were deemed non-diagnostic due to inaccurately registered thin myocardium.

In the patients, the average myocardial T$_2$ relaxation times did differ between the 2D ($46.7 \pm 3.6$ ms) and 3D techniques ($50.1 \pm 4.2$ ms, $P = 0.004$, Figures 2, 4). The CoV was again lower in the 2D technique than in the 3D technique at $6.8 \pm 1.5$ vs. $10.4 \pm 1.8\%$ ($P < 0.001$). When the myocardium was segmented according to the AHA guidelines, the two techniques resulted in significantly different T$_2$ values in the basal-anterior, basal-inferior, mid-inferoseptal and mid-inferior segments (segment numbers 1, 4, 9, and 10; $P \leq 0.001$). Conversely, the segmental CoV in the patients was significantly lower for the 2D technique in all segments ($P < 0.001$) except in the mid-inferoseptal segment. Out of the 300 myocardial segments analyzed in the patients, 14 (4.6\%, of which 8 [2.6\%] in one patient) were deemed non-diagnostic. A Bland-Altman analysis of the 2D vs. the 3D technique demonstrated that there was a small bias of $-3.0$ ms (Figure 5A). Although a slight trend can be observed for the difference to become more negative as the average increases, this trend was not significant ($\rho = -0.33, P = 0.11$). The patient T$_2$ values as measured with the 2D technique were not significantly different from those in healthy volunteers ($P = 0.50$), while the difference was significant when measured with the 3D technique ($P = 0.04$).

Inter-observer analyses of the 2D and 3D techniques showed very small biases between the observers, no visible or significant trends ($\rho \leq 0.27, P \geq 0.20$), and similar confidence intervals (Figures 5B,C). The intra-observer analyses of a subset of patients ($n = 15$) showed a higher confidence interval for the 3D technique (Figures 5D,E) and no significant trend for either technique ($\rho < 0.35, P > 0.2$). The visual quality score was higher for the 2D maps than for the 3D maps at 9.6 ± 0.4 vs. 7.2 ± 2.1 ($P < 0.001$, Supplementary Figure 3).

The LGE-positive segments did not have a significantly higher T$_2$ relaxation when compared to their LGE-negative counterparts as quantified with the 2D technique (T$_{2\text{LGE}-} = 46.2 \pm 3.7$ vs. T$_{2\text{LGE}+} = 47.6 \pm 4.1$ ms, $P = 0.49$, Figures 6, 7). Conversely, the 3D technique did result in a significant difference (T$_{2\text{LGE}-} = 49.3 \pm 6.7$ vs. T$_{2\text{LGE}+} = 52.6 \pm 8.7$ ms, $P = 0.006$) despite its larger spread in individual T$_2$ values. Both 2D segment groupings were also significantly different from their 3D equivalents ($P < 0.001$). The Bloch equation simulations indicated that the proposed 3D mapping technique will moderately underestimate (<5\%) the T$_2$ relaxation times for high heart rates (80–90 bpm, Supplementary Figure 4).

**DISCUSSION**

In this study, we proposed a novel method to obtain high-resolution respiratory motion-corrected 3D T$_2$ maps of the heart at 1.5T by extracting, resolving, and then registering the respiratory motion states, and tested this technique in patients and healthy volunteers. The respiratory motion was detected and visually resolved well in all subjects, and the resulting isotropic 3D T$_2$ values matched their routine 2D counterparts while also enabling the visualization of other views of the heart.

The T$_2$ CoV (i.e., the inverse of precision) agreed well with that obtained in previous 3D T$_2$ mapping studies (8, 12), but was higher than commonly reported for 2D T$_2$ mapping (31, 32). The main two causes of this are most likely the ($1.9 \times 1.9 \times 8)/1.6^3 = 7$ times smaller voxel size and the radial acquisition itself, although this is partially compensated by the denoising effect of the compressed-sensing reconstruction and the 3D nature of the acquisition. The higher CoV values that are seen in some segments might result from these particular segments being more prone to small misregistrations due to the neighboring epicardial fat or small myocardial thickness. If a higher precision is desired, it can most likely be achieved by increasing the voxel size, the acquisition time, or the regularization parameters of the reconstruction. Similarly, the observed residual blurring in the short-axis plane could for example be caused by incorrect motion extraction or misregistration. The slightly (but not significantly) different T$_2$ averages in the healthy volunteers with the 2D and 3D techniques

![Figure 6](image-url)
FIGURE 7 | A comparison of the 2D and 3D T$_2$ mapping techniques in slices with lesions detected with LGE imaging. (A) LGE shows intramyocardial areas of irreversible damage in several basal regions of the LV in a 56 y.o. male patient with suspected cardiac sarcoidosis. (B) No T$_2$ elevation is visually apparent on the 2D T$_2$ map. As an example, the measured segmental ROI average T$_2$ in the basal inferolateral segment was 46 ms. (C) Conversely, local increases of the T$_2$ relaxation time can be spotted on the 3D T$_2$ map in areas corresponding to the LGE regions (white arrows). The measured segmental ROI average T$_2$ in the respective inferolateral segment was 56 ms, suggestive of an area of acute injury or ongoing inflammation. (D) An intramyocardial area of irreversible damage as shown by LGE (left, white arrow) in the basal inferoseptal segment of the LV in a 66 y.o. male patient with suspected cardiac sarcoidosis. The pink arrowhead indicates an ischemic scar in the anterior wall. (E) Again, no T$_2$ elevation was visually apparent on the 2D T$_2$ map in the respective segment with the segmental ROI average T$_2$ = 47 ms. (F) Local (white arrow) increase of the T$_2$ relaxation time (segmental ROI average T$_2$ = 55 ms) can be spotted in this region on the 3D T$_2$ map, suggestive of an area of acute injury. White arrowhead points to another area of elevated T$_2$ that does not correspond to any LGE in (D). The institutional reference range for 2D T$_2$ mapping is 44 (2.4) ms, i.e., 39–49 ms. LGE, late gadolinium enhancement; LV, left ventricle; ROI, region of interest.

most likely also have several origins, such as a different interplay with the T$_1$ relaxation time during the acquisition (24), and a residual noise floor that promotes increased T$_2$ relaxation time measurements (25). The observed map sharpness agrees well with previous map sharpness quantifications (33), while the minor (and non-significant) differences between the reconstructions with different regularization weights suggest a relatively broad optimum for the regularization parameter. The alternative reconstruction where T$_2$ maps were obtained for the four respiratory motion bins, followed by averaging of these bins, also led to similar map sharpness, though at the cost of 4-fold longer map fitting time.

It is well-known that T$_2$ reference values in healthy subjects may vary and that they are therefore not easily comparable between studies (34–36). In addition to magnetic field strength, pulse sequence type, and specific parameters (32), other factors play a role, including gender and cardiac physiology (34). Therefore, the assessment of institution-specific reference ranges with established scanner- and sequence-specific T$_2$ values in healthy volunteers are strongly recommended (2). With scanner- and sequence-specific reference values as a prerequisite, it was demonstrated that T$_2$ values in studies of specific inflammatory conditions such as myocarditis (3), systemic sclerosis (37), and sarcoidosis (35) can unequivocally be used to identify myocardial injury. Therefore, the significant difference in T$_2$ relaxation time that was found in patients using the 3D but not the 2D T$_2$ mapping merits special attention, as this might suggest better sensitivity to detect tissue alterations with the proposed high-resolution motion-registered radial 3D T$_2$ mapping technique at 1.5T. However, confounders such as the observed blurring, SNR, irregular breathing patterns of the patients, and the heart-rate dependence of the pulse sequence should also be taken into consideration. Of note, the patient population in our study was a fairly “healthy” patient population that included consecutive patients with suspected myocardial inflammation, where no large elevation of T$_2$ was observed by conventional protocol. Therefore, the significantly higher T$_2$ values in 3D maps in patients that were not seen in conventional 2D maps and were higher than 3D T$_2$ values in controls suggest potential for the detection of more subtle changes or earlier stages of
disease. Further study will be required in a broader spectrum of disease severity to shed more light on these findings. The proposed 3D technique indicated a small but significant $T_2$ difference between the LGE+ and LGE− myocardial segments, while the routine 2D technique did not. This may again have a technical origin such as the smaller voxels in the 3D maps having less partial volume effect (especially through the much thinner slices) with healthy tissue that blunts the signal change and therefore reduces the sensitivity. Even though LGE presence by itself does not directly indicate edema or inflammation, it potentially colocalizes with patches of active disease in patients with an ongoing disease process. Thus, the slightly increased $T_2$ value in LGE+ segments suggests that smaller and isotropic voxels may be more appropriate in patients with potentially focal non-ischemic injury. Nevertheless, a higher sensitivity of 3D $T_2$ mapping would need to be confirmed with a clinical standard method such as positron emission tomography (38).

While we did not evaluate short-axis apical segments in this study, it may be of interest to include these in future studies to take advantage of a reduced partial volume effect consistent with smaller voxel sizes.

The acquisition itself could be improved in several ways in future studies. By using a different radial k-space trajectory for each $T_2$-prepared volume (39), the sparsity in the relaxation time dimension could also be exploited for a compressed sensing reconstruction (11), which should result in a higher precision. Calculating the $T_2$ in each pixel based on an individually patient-specific simulated dictionary (40, 41) could also be used to remove the constraint of acquiring every other heartbeat, which would significantly accelerate the acquisition, and would remove the heart-rate dependence of the $T_2$ relaxation time.

The higher variation in quality of the 3D maps as compared to the 2D maps is not surprising, as the 2D $T_2$ maps were immediately re-acquired during the scanning sessions if they were visually of unsatisfactory quality, which was not feasible for 3D $T_2$ maps because of time constraints. Of note, most of the segments that were termed non-diagnostic belonged to a single patient and a single volunteer, which suggests that there might have been a subject-specific challenge involved, such as particularly thin myocardium or a sub-optimal ECG triggering. Therefore, considering the inherent complexity of the isotropic 3D sequence and multiple reconstruction steps required in the proposed technique, the observed quality score scatter is an expected trade-off, especially with the improvement potential outlined above. Studies with a combination of pseudo-spiral Cartesian trajectory (instead of the 3D radial used here), a 2D respiratory motion correction (instead of the 1D correction), and the addition of patch-based denoising (which improves the apparent SNR) did result in consistent $T_2$ precision (11, 13). The patch-based denoising (42) in particular could improve the 3D $T_2$ mapping presented here, since it might improve the apparent SNR of the source images, resulting in better image registration as well as improved map precision.

In conclusion, respiratory motion-registered 3D radial imaging at 1.5T led to accurate isotropic 3D whole-heart $T_2$ maps, both in the healthy volunteers and in a small patient cohort with suspected inflammatory myocardial injury. With the 3D technique, significantly higher $T_2$ values were found in patients as compared to controls, as well as in LGE-positive as compared to LGE-negative segments, both of which were not observed with the routine 2D technique. These findings are suggestive of the technique's potential to increase the sensitivity of CMR for localized inflammatory myocardial injury. Further study will be needed in a broader spectrum of disease severity to demonstrate its clinical utility.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Institutional Review Board of the Medical University of Gdansk. The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

KD and RBvH designed the study. LD, CR, DP, MS, JY, and RBvH developed and implemented the pulse sequence and image reconstruction algorithm. KD, JF, and HJ recruited the volunteers and patients and performed their diagnoses. AS and AG performed the MR scanning. RBvH supervised the analysis of MRI results and the manuscript draft. KD, KG, AS, ES, and RBvH analyzed the results and performed the statistical analysis. All authors participated in manuscript drafting and editing, approved the final version of the submitted manuscript, and agreed to the submission of the manuscript to Frontiers in Cardiovascular Medicine.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcvm.2021.712383/full#supplementary-material

Supplementary Figure 1 | A flowchart of the proposed 3D T2 mapping method. Data are indicated in colored boxes, while manipulations are mentioned as unboxed text.

Supplementary Figure 2 | Sharpness measurement and various reconstructions of a 3D T2 map. (A) A T2 map in grayscale with a display range from 0 to 300 ms to better visualize the blood pool, with red crosses indicating the start- and end-points of the assessed transition. Four horizontal pixel lines below it were also assessed. (B) Normalized T2 values in the pixel (blue circles) and the parametrized sigmoid fit (red curve). The sharpness of the transition was characterized by $k = 2.6 \, \text{px}^{-1}$, which at the 1.6 mm spatial resolution translated to $k = 1.6 \, \text{mm}^{-1}$. This value was averaged with those obtained from the lines below it. (C) The proposed 3D T2 mapping method in another patient. (D) An alternative reconstruction method in which the individual respiratory bins are all mapped first, and these bin maps are then averaged. While the papillary muscle appears to resolve better, the sharpness of the blood-myocardium interface remains constant. (E,F) The proposed reconstruction method, but with $k = 0.01$ and 0.25. The blood-myocardium interfaces appear marginally sharper and more blurred, respectively, but this is balanced by respectively increased and decreased T2 variation in the myocardium.

Supplementary Figure 3 | A paired comparison of the visual quality scores as agreed by two experienced readers. The 3D T2 maps have a much higher variation in quality than the 2D T2 maps ($7.2 \pm 2.1 \text{ vs. 9.6} \pm 0.4, P < 0.001$), although it should be noted that the 2D T2 maps were immediately re-acquired during the scanning sessions if they were visually of unsatisfactory quality.

Supplementary Figure 4 | A Bloch equation simulation of the influence of the patient’s heart rate on the estimated T2 value. Since sampling occurs every other heartbeat, T1 relaxation will cause a gradual underestimation of the T2 value with the increase of the heart rate.

Supplementary Animated Figure 1 | An animation of the respiratory-resolved reconstruction of a source image with T2prep duration of 30 ms in a patient. The breathing motion is well-resolved, and results in a non-rigid translation of the heart.

REFERENCES


Conflict of Interest: DP and KG are full-time employees of Siemens Healthcare.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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