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## Free-Breathing Multitasking Multi-Echo (MT-ME) MRI for Whole-Liver Water-Specific T1, Proton Density Fat Fraction, and R2\* Quantification

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## Abstract

**Purpose:** To develop a three-dimensional Multitasking multi-echo (MT-ME) technique for the comprehensive characterization of liver tissues with 5-minute free-breathing acquisition, whole-liver coverage, a spatial resolution of  $1.5 \times 1.5 \times 6 \text{ mm}^3$ , and simultaneous quantification of T1, water-specific T1 (T1<sub>w</sub>), PDFF, and R2\*.

**Methods:** Six-echo bipolar spoiled GRE readouts following inversion recovery (IR) preparation was performed to generate T1, water/fat and R2\* contrast. MR Multitasking was used to reconstruct the MT-ME images with 3 spatial dimensions, one T1 recovery dimension, one multi-echo dimension, and one respiratory dimension. A basis-function-based approach was developed for  $T1_w$  quantification, followed by the estimation of R2\* and T1-corrected PDFF. The intra-session repeatability and the agreement against references of MT-ME measurements were tested on a phantom and 15 clinically healthy subjects. In addition, 4 patients with confirmed liver diseases were recruited and the agreement between MT-ME measurements and references was assessed.

**Results:** MT-ME produced high-quality, co-registered T1,  $T1_w$ , PDFF, and R2\* maps with good intra-session repeatability and substantial agreement with references on phantom and human

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studies. The ICCs of T1, T1<sub>w</sub>, PDFF, and R2\* from the repeat MT-ME measurements on clinically healthy subjects were 0.989, 0.990, 0.999, 0.988, respectively. The ICCs of T1, PDFF, and R2\* between the MT-ME and reference measurements were 0.924, 0.987, 0.975 in healthy and 0.980, 0.999, 0.998 in patients. The T1<sub>w</sub> was independent to PDFF (R=-0.029, P=0.904).

**Conclusion:** The proposed MT-ME technique quantifies T1, T1<sub>w</sub>, PDFF, and R2\* simultaneously and is clinically promising for the comprehensive characterization of liver tissue properties.

#### Keywords

liver T1/PDFF/R2\* mapping; MR Multitasking; low-rank tensor; water-specific T1; free-breathing acquisition

### 1. Introduction

Chronic liver disease is increasing in prevalence and has become a major cause of mortality worldwide in recent years.<sup>1,2</sup> Nonalcoholic fatty liver disease (NAFLD) is one of the major types, which is estimated to affect 25% of the world population.<sup>2</sup> The progressive form of NAFLD is nonalcoholic steatohepatitis (NASH), which is typically accompanied by liver inflammation and fibrosis<sup>2</sup>. Liver fibrosis can advance into cirrhosis, an end-stage disease that invariably leads to death<sup>3,4</sup>. Understanding the complex interplay between liver fat, iron concentration, and fibrosis is of great significance to the management of liver disease. Currently, liver biopsy is the gold standard for the diagnosis and staging of liver disease. However, it is invasive, prone to sampling error, associated with risks of complications, and subject to inter-observer variability<sup>5–7</sup>. Therefore, an objective, non-invasive tool for comprehensive evaluation of liver tissues is highly desired.

In recent years, quantitative MRI with multiparametric mapping has been a promising approach for the objective, noninvasive assessment of liver diseases<sup>7–12</sup>. Proton density fat fraction (PDFF) is a widely used MRI marker to quantify tissue fat concentration<sup>13–15</sup>. R2\*, the reciprocal of T2\*, is proportional to iron content in the liver<sup>10,16</sup>. MR elastography (MRE) is a widely-accepted approach to measure liver stiffness as well as fibrosis but requires complicated setups and additional external hardware<sup>17</sup>. Recently, the role of T1 in the assessment of liver fibrosis and inflammation has gained increased interest. Prior investigations have reported that the T1 relaxation time is significantly lengthened in fibrotic and cirrhotic livers in both animal and human studies with histological confirmation<sup>12,18–21</sup>. Some works showed that T1 values were significantly correlated with MRE measurements; it also had reduced inter-reader variability and improved reproducibility compared to MRE<sup>20,22</sup>. These studies indicate that T1 may have the potential to serve as a surrogate biomarker of MRE in the longitudinal monitoring of liver disease with easier setup and improved reliability.

Despite all the encouraging results, a standard clinical protocol including the measurement of PDFF, R2\*, and T1 for liver diseases has yet to be established due to demanding technical challenges. First, while PDFF and R2\* are measured simultaneously in one 3D acquisition, T1 usually needs to be measured with several repeated 2D acquisitions. Well-validated

3D T1 mapping techniques for liver are lacking. The most frequently used sequences for T1 measurement are 2D modified Look-Locker inversion recovery (MOLLI)<sup>23</sup>, shortened MOLLI (shMOLLI)<sup>24</sup>, or saturation-recovery single shot acquisition (SASHA)<sup>25</sup>, which mandates a number of breath-holds to cover multiple slices. This can cause extra burden on patients, reduced reliability, and misregistration across breath-holds. Second, the breath-hold sequences force compromises between spatial coverage, spatial resolution, and scan time (limited by breath-hold duration). Third, the presence of liver fat confounds the quantification of water T1 and reduces the sensitivity of T1 in the estimation of fibrosis<sup>26,27</sup>. Previous works have demonstrated that MOLLI/shMOLLI or SASHA produce large and complex T1 errors when applied to mixed water-fat systems<sup>28</sup>. Some recent publications have developed approaches that can quantify T1, PDFF, and R2\* simultaneously within a single scan, and correct T1 confounding from fat by measuring water-specific T1 (T1<sub>w</sub>)<sup>8,9</sup>. However, these techniques still require breath-holds and have limited spatial coverage.

To address the above-mentioned limitations, we propose a novel MR technique, Multitasking multi-echo (MT-ME), for the comprehensive characterization of liver tissues. MR Multitasking<sup>29</sup> uses a low-rank tensor (LRT) imaging model to exploit the high correlation along and across different image dimensions for a vastly accelerated acquisition, and has been demonstrated for 2D multiparametric mapping in moving organs and 3D multiparametric mapping in the brain<sup>30–32</sup>. MT-ME is the first Multitasking technique for 3D multiparametric mapping on moving organs. The aim of this study was to describe and validate the MT-ME technique for whole-liver free-breathing acquisition and simultaneous quantification of water-specific T1 (T1<sub>w</sub>), PDFF, and R2\* in a phantom and in vivo.

#### 2. Methods

#### 2.1 Sequence and sampling

A 3D continuous-acquisition sequence was designed based on spoiled gradient echo (spoiled GRE) readouts, consisting of periodic non-selective inversion recovery (IR) preparations and 6-echo bipolar readouts to generate and acquire T1, water/fat, and R2\* contrasts, as shown in Figure 1A. Both RF- and gradient-spoiling are used to reduce the residual transverse magnetization at the end of each readout. The TEs were chosen as 1.25, 2.50, 3.75, 5.00, 6.25, 7.50 ms according to Zhong et al<sup>14</sup>. An axial stack-of-stars sampling pattern with golden-angle rotation in  $k_x$ - $k_y$  plane and Gaussian-density randomized Cartesian reordering in the  $k_z$  dimension was used to incoherently sample k-space. A perpendicular center k-space line in the head–foot direction ( $k_x = k_y = 0$ ) was acquired every 8 readouts (every 74 ms) to form the training data, which were used to determine LRT temporal factors. Positioning the training center k-space lines along the  $k_z$  direction better captures respiratory motion<sup>33</sup>. The sampling pattern was demonstrated in Figure 1B.

#### 2.2 Image reconstruction

The target MT-ME images have three spatial dimensions *x*, *y*, *z*, and three temporal dimensions: IR time  $T_I$ , multi-echo dimension  $T_E$ , and respiratory motion bin *r*. It can be represented as a 4-way tensor  $\mathscr{A}$  by collapsing the three spatial dimensions into voxel index  $\mathbf{x} = [x \ y \ z]^T$ . The strong correlation along and across multiple dimensions of  $\mathscr{A}$  induces it

to be low-rank, so that it can be decomposed into the product of different basis matrices representing each dimension<sup>29,34–36</sup>:

$$\mathscr{A} = \mathscr{G} \times_1 \mathbf{U} \times_2 \mathbf{V} \times_3 \mathbf{W} \times_4 \mathbf{Q},\tag{1}$$

where the columns of **U**, **V**, **W** and **Q** contain the basis functions for spatial, IR, multi-echo, and respiratory dimensions, respectively,  $\mathscr{G}$  is the core tensor, and  $\times_i$  denotes the *i*th mode product. The reconstruction of the final images can be framed as the recovery of each basis matrix, which significantly reduces the total degrees of freedom and requires a much shorter scan than without the LRT model.

The first step was to determine **V** from a pre-generated dictionary describing IR according to the Bloch equation. The dictionary consisted of 101 T1 values logarithmically spaced from 100 ms to 3000 ms, 17 flip angles from  $1^{\circ}$  to  $9^{\circ}$  in half-degree increments, and 21 inversion pulse flip angles linearly spaced from  $90^{\circ}$  to  $180^{\circ}$ . The IR basis matrix **V** was directly obtained from the singular value decomposition (SVD) of this dictionary.

The second step was to identify respiratory motion and divide data into 6 respiratory bins using an automatic respiratory binning method described in our previous work<sup>29</sup>. The prior-generated IR basis matrix V was used to compensate the T1 contrast of the training data. Then a modified k-means clustering algorithm was performed to assign each training data readout to a specific respiratory bin. All the data at different respiratory bins were used for image reconstruction. After the reconstruction, images at all 6 bins are available.

The third step was to recover **W**, **Q**, and  $\mathscr{G}$  through the completion of the training data tensor. The acquired (k,t)-space training data  $\mathbf{d}_{tr}$  were reshaped into a 4-way tensor  $\mathscr{D}_{tr}$  with one k-space readout dimension, and three temporal modes  $T_F$ ,  $T_E$ , and r. The training data were not available from every combination of  $T_F$ ,  $T_E$ , and respiratory state, resulting in an undersampled  $\mathscr{D}_{tr}$ . A small-scale low-rank tensor completion algorithm was applied to recover the training data tensor using:

$$\widehat{\mathscr{D}}_{\text{tr}} = \arg \min_{\mathbf{D}_{\text{tr},(2)} \in \text{range}(\mathbf{V})} \|\mathbf{d}_{\text{tr}} - M(\mathscr{D}_{\text{tr}})\|_2^2 + \lambda \sum_{i=1,3,4} \|\mathbf{D}_{\text{tr},(i)}\|_* + R(\mathscr{D}_{\text{tr}}),$$
(2)

where  $M(\cdot)$  is the training data sampling pattern,  $\|\cdot\|_*$  denotes the nuclear norm,  $\mathbf{D}_{tr,(i)}$  denotes the mode-*i* matricization of the tensor  $\mathcal{D}_{tr}$ , and  $R(\cdot)$  is an optional additional regularization functional, which was chosen as temporal total variation (TV) along the respiratory dimension in this work. Following the recovery of  $\widehat{\mathcal{D}}_{tr}$ , the core-tensor  $\mathscr{G}$ , multi-echo basis functions  $\mathbf{W}$ , and respiratory basis functions  $\mathbf{Q}$  were obtained using the higher-order SVD (HOSVD)<sup>37</sup>.

The final step was to recover the spatial basis functions  $\mathbf{U}$  by fitting the temporal basis functions and core tensor into the acquired imaging data  $\mathbf{d}$ :

$$\widehat{\mathbf{U}} = \arg\min_{\mathbf{U}} \|\mathbf{d} - \mathbf{F}_{\Omega}(\boldsymbol{\Phi} \times_1 \mathbf{S}\mathbf{U})\|_2^2 + R(\mathbf{U}),$$
(3)

where  $\Phi = \mathscr{C} \times_2 \mathbf{V} \times_3 \mathbf{W} \times_4 \mathbf{Q}$  is the product of the core tensor and temporal basis functions, **S** is the coil sensitivity operator,  $\mathbf{F}_{\Omega}$  is the nonuniform fast Fourier transform (NUFFT) on the undersampling set  $\Omega$ , and  $R(\cdot)$  is an optional regularization functional chosen as a spatial wavelet regularizer. In this work, gpuNUFFT<sup>38</sup> was used to compute nonuniform fast Fourier transform.

#### 2.3 Multiparametric quantification

**2.3.1 Joint T1**—In this work, "Joint T1", denoted as the T1 with both water and fat components, was quantified based on the first-echo MT-ME images at the end-expiration state using the following equation<sup>39</sup>:

$$s(A, B, T1) = A \frac{1 - e^{-T_{\rm R}/T1}}{1 - e^{-T_{\rm R}/T1} \cos(\alpha)} \Big[ 1 + (B - 1) \Big( e^{-T_{\rm R}/T1} \cos(\alpha) \Big)^n \Big] \sin(\alpha), \tag{4}$$

with amplitude *A*, inversion effect *B*, flip angle  $\alpha$ , spoiled GRE readout interval *T<sub>R</sub>*, and recovery time point *n* = 1, 2, …, *N*(*N*=192 per IR period) such that  $\tau = nT_R$ . In the quantification, *A*, *B*, and *T*1 were estimated jointly from the signal evolution. The effect of the RF transmit field (B<sub>1</sub><sup>+</sup>) inhomogeneity on the T1 estimation was analyzed and is presented in the Supporting Information Section D.

**2.3.2** Water-specific T1 (T1<sub>w</sub>)—One straightforward way to map T1<sub>w</sub> would be to perform water/fat separation on the multi-echo images at each inversion time, and then perform the T1 mapping using the water images only. However, for 3D images with 192 inversion times, this approach would be time-consuming. Therefore, we performed a basis-function-based approach<sup>8,40</sup>. Specifically, a partially-recomposed tensor  $\mathcal{Y}$  was generated based on model basis factors:

$$\mathscr{Y} = \mathscr{G} \times_1 \mathbf{U} \times_3 \mathbf{W} \times_4 \mathbf{Q} = \mathscr{A} \times_2 \mathbf{V}^+, \tag{5}$$

where the superscript "+" denotes the pseudoinverse.  $\mathcal{Y}$  is composed of all model factors except the IR basis **V**, therefore keeping T1 information embedded and reducing the number of water/fat separations that must be performed: for any given respiratory bin, water/fat separation only needs to be done 5 times (the dimension of **V**), rather than on each of the 192 inversion times.  $\mathcal{Y}$  is a three-way tensor with a spatial dimension, a multi-echo dimension, and a respiratory dimension. The water-fat separation was then performed on the end-expiration images of  $\mathcal{Y}$  using a multiple-fat-peak single-R2\* model:

$$\mathscr{Y}(\mathscr{W},\mathscr{F},f_{B0},R_2^*) = \left(\mathscr{W} + \mathscr{F}\sum_l C_l e^{i2\pi f_l T_E}\right) e^{if_{B0}T_E - R_2^* T_E},\tag{6}$$

where  $\mathscr{W}$  and  $\mathscr{F}$  are the respiratory-motion-resolved water and fat components, respectively,  $C_I$  and  $f_I$  are the weighting and the resonance frequency offsets of the *I*th fat peak,  $T_E$  is the echo time, and  $f_{B0}$  represents B0 inhomogeneity. A multi-peak fat model was employed in this work according to Yu et al<sup>41</sup>. The water-fat separation was performed using an in-house algorithm using variable projection (VARPRO) according to Hernando et al<sup>42</sup>.

$$\mathscr{A}_{\mathsf{W}} = \mathscr{W} \times_2 \mathsf{V} \,. \tag{7}$$

T1 quantification was performed  $\mathscr{A}_w$  using Equation 4, yielding T1<sub>w</sub> value. The fat-specific T1, denoted as T1<sub>f</sub>, was also generated similarly on the fat images  $\mathscr{A}_f$  with multiple inversion times:

$$\mathscr{A}_{\mathrm{f}} = \mathscr{F} \sum_{l} C_{l} \times_{2} \mathbf{V} \,. \tag{8}$$

To validate the feasibility of the basis-function-based approach, a comparison was performed between the  $T1_w$  values from the basis-function-based approach and from the straightforward approach where the T1 mapping was conducted on the water images of 192 inversion times, as described in Supporting Information Section A.

**2.3.3 PDFF and R2\* mapping**—Following the estimation of  $T1_w$  and  $T1_f$ , PDFF and R2\* were fitted together using the multi-echo images of MT-ME at end-expiration at the last TI. The T1 effect of water and fat components<sup>43</sup> were also taken into consideration:

$$s(\rho_{\rm w}, \rho_{\rm f}, f_{B0}, R_2^*) = \left(M_{\rm w, last}\rho_{\rm w} + M_{\rm f, last}\rho_{\rm f}\sum_l C_l e^{i2\pi f_l T_E}\right) e^{if_{B0}T_E - R_2^* T_E},\tag{9}$$

where  $\rho_w$  and  $\rho_f$  are the proton density of water and fat, and  $M_{w,last}$  and  $M_{f,last}$  represent the spoiled GRE signal intensity at the last inversion time of water and fat component, which can be calculated with the estimated T1<sub>w</sub> and T1<sub>f</sub> according to Equation 4:

$$M_{\rm w,last} = \frac{1 - e^{-T_{\rm R}/T_{\rm I}}w}{1 - e^{-T_{\rm R}/T_{\rm I}}w_{\cos(\alpha)}} \bigg[ 1 + (B - 1) \Big( e^{-T_{\rm R}/T_{\rm I}}w_{\cos(\alpha)} \Big)^{192} \bigg] \sin(\alpha),$$

$$M_{\rm f,last} = \frac{1 - e^{-T_{\rm R}/T_{\rm f}} f_{\rm cos(\alpha)}}{1 - e^{-T_{\rm R}/T_{\rm f}} f_{\rm cos(\alpha)}} \Big[ 1 + (B - 1) \Big( e^{-T_{\rm R}/T_{\rm f}} f_{\rm cos(\alpha)} \Big)^{192} \Big] \sin(\alpha)$$

The same in-house algorithm as in Section 2.3.2 was used for the estimation of PDFF and R2\*. The flow chart in Figure 2 illustrates the entire process of the reconstruction and multiparametric mapping of MT-ME.

#### 2.4 Imaging experiments

Experiments were performed on a phantom, clinically healthy subjects (N = 15), and patients with diagnosed liver diseases (N = 4). The in vivo studies were approved by the local institutional review board. All subjects provided written informed consent before scanning. The phantom and volunteer studies were performed on a 3T system (MAGNETOM Vida, Siemens Healthcare, Erlangen, Germany) with an anterior 18-channel flexible array coil in combination with 12–16 elements of the table-mounted spine

array. The patient study was performed on another 3T scanner (MAGNETOM Skyra, Siemens Healthcare, Erlangen, Germany) with the same coil settings. The detailed imaging parameters of the sequences used in the experiments are included in Supporting Information Table S1.

**2.4.1 Phantom study**—A commercial phantom (Calimetrix, Madison, WI) was used in this work with 15 vials, each of which has a unique combination of T1, PDFF, and R2\* values. A prototype single-slice inversion recovery spin echo (IR-SE) without and with water-selective excitation were both acquired at the center slice as references for T1 and T1<sub>w</sub>, respectively. A product q-DIXON sequence with 3D volumetric interpolated breath-hold examination (VIBE) acquisition and 6-echo bipolar readout was acquired as the reference for PDFF and R2<sup>\*14</sup>. Following the reference scans, the proposed MT-ME sequence was performed twice to assess its feasibility and intra-session repeatability.

In addition, a custom phantom was designed<sup>44</sup> to specifically test the ability of MT-ME to map the  $T1_w$  in the samples with T1 values that match the liver T1 at 3T (700 to 1200 ms<sup>9,20,45</sup>) in the presence of fat. The details of the phantom study are described in the Supporting Information Section C.

**2.4.2** Volunteer study—Fifteen clinically healthy subjects (3 females; age 38±12 years, BMI 24.8±3.4 kg/m<sup>2</sup>) without a record of liver diseases were recruited as the clinically healthy group. All the scans were performed in transversal orientation. A product 2-echo VIBE-DIXON sequence was performed to generate anatomical water and fat images with following parameters: FOV =  $308 \times 380$  mm<sup>2</sup>, matrix =  $195 \times 320$  mm<sup>2</sup>, TE = 1.29/2.52 ms, TR = 3.60 ms, slice thickness = 6 mm. The prototype 2D MOLLI sequence was collected under breath-hold at end-expiration for 3 slices at the top, middle and bottom of the liver. The parameters for each slice are: FOV =  $306 \times 360 \text{ mm}^2$ , matrix =  $204 \times 240 \text{ mm}^2$ , TE = 1.17 ms, TR = 2.63 ms, slice thickness = 8 mm. The product q-DIXON VIBE sequence with 6-echo bipolar readout was conducted to provide reference PDFF and R2\* values under breath-hold at end-expiration with the parameters:  $FOV = 359 \times 359 \text{ mm}^2$ , matrix =  $128 \times 128 \text{ mm}^2$ , TE = 1.25/2.50/3.75/5.00/6.25/7.50 ms, TR = 9.28 ms, slice thickness = 6mm. Following the breath-hold scans, the proposed free-breathing MT-ME sequence was performed covering the whole liver with the parameters:  $FOV = 288 \times 288 \text{ mm}^2$ , matrix =  $192 \times 192 \text{ mm}^2$ , TE = 1.25/2.50/3.75/5.00/6.25/7.50 ms, TR = 9.20 ms, slice thickness = 6mm, number of slices = 40. The entire protocol was performed twice in the same imaging session half-an-hour apart to assess the intra-session repeatability of MT-ME.

**2.4.3 Patient study**—Four patients (1 female; age  $65\pm4$  years, BMI  $31.5\pm2.3$  kg/m<sup>2</sup>) previously diagnosed with biopsy-confirmed chronic liver diseases were recruited for the study: two with NAFLD and two with NASH. The patients first received a set of clinical standard-of-care MRI scans including 2-echo VIBE-DIXON and 6-echo q-DIXON. Then, MOLLI and MT-ME were performed with the same settings as in the volunteer study. MT-ME was performed one time on each patient due to time limitations.

#### 2.5 Analysis

Image reconstruction and quantitative mapping of T1, T1<sub>w</sub>, PDFF, and R2\* were performed off-line in MATLAB (R2019a, Mathworks, MA, USA) for all MT-ME data. An imaging physician (AK) assessed all the images, including 1) water images, fat images, T1, T1<sub>w</sub>, PDFF, and R2\* maps from MT-ME; 2) water images, fat images, PDFF and R2\* maps from q-DIXON; 3) water and fat images from VIBE-DIXON; and 4) T1 maps from MOLLI. For each set of images, the subjective SNR, artifacts, and sharpness were evaluated and a score of overall image quality was given using a 5-point scale: 1-worst (anatomic details were impossible to identify, and there were extremely high noise and artifacts); 2-bad (anatomic details were difficult to identify, and there were significant noise and artifacts); 3-moderate (anatomic structures were visible but not clearly shown, and there were acceptable noise and artifacts); 5-excellent (the anatomic structures and details were highly visible and there were no significant noise and artifacts).

The T1, T1<sub>w</sub>, PDFF, and R2\* were measured on phantom and in vivo. In phantom, the ROIs were drawn on a center slice within each vial avoiding the edge. For the in vivo study, the ROIs of MT-ME images were manually drawn on the three slices matching the locations of MOLLI. As much liver tissue as possible was included while avoiding blood vessels, nodules/cysts (if any), and edges. The same ROIs were applied to the maps of T1, T1<sub>w</sub>, PDFF, and R2\*. For reference scans, the ROIs were also drawn on three matched slices avoiding blood vessels, nodules/cysts (if any), and edge. The mean value throughout all pixels was reported for each parameter of each measurement.

The agreement of T1, T1<sub>w</sub>, PDFF, and R2\* measurements from MT-ME and references were evaluated by the sample Pearson correlation coefficient R and intra-class coefficient (ICC) in phantom and in vivo studies. A paired t-test was used to identify the significant difference of parameters measured with the MT-ME against references. A P value less than 0.05 is considered as a significant difference. The repeatability of the MT-ME measurements was assessed by the Bland-Altman plots, ICC and coefficient of variation (CoV) on phantom and volunteer studies.

#### 3. Results

#### 3.1 Phantom study

Vials 1 to 15 are labeled on the gray-scale image in Figure 3A. Vial 14 is a fat-only phantom, whose  $T1_w$  value is not available from water-excited IR-SE. The quantitative map of T1, T1<sub>w</sub>, PDFF, and R2\* from MT-ME and from references were displayed in Figure 3A. The Bland-Altman plots in Figure 3B shows good repeatability of MT-ME for all the four parameters. The ICCs of T1, T1<sub>w</sub>, PDFF, and R2\* were 0.999, 0.999, >0.999, >0.999, respectively, and the CoVs were 1.4%, 0.5%, 1.1%, 0.9%, respectively. Figure 3C demonstrates the agreement of parameters measured by MT-ME and references. The Pearson Correlation Coefficients (*R*) were 0.990, 0.996, 0.999, >0.999, respectively, and the ICC were 0.988, 0.994, 0.998, 0.999, respectively, representing good agreements. The paired t-test indicated that the measurement of MT-ME and references showed no significant

differences for all the four parameters with P = 0.126, 0.097, 0.319, 0.067. The results of the custom phantom are illustrated in the Supporting Information Figure S3.

#### 3.2 In vivo study

The in vivo protocols were successfully applied to all healthy volunteers and patients, and MT-ME images were reconstructed and processed for all subjects. The reconstructed MT-ME images possess 192 TIs from 9.2 ms to 1766.4 ms with an increment of 9.2 ms, 6 echoes, and 6 respiratory bins, yielding a total of  $192 \times 6 \times 6 = 6912$  images, each corresponding to a 3D image volume. Figure 4 is an illustration of the images displayed in either coronal or axial orientation at 3 echoes (echo 1, echo 2, and echo 6), 2 representative TIs (266.8 ms and 1766.4 ms) and 2 respiratory states (end-expiration and end-inspiration). The coronal view with reference lines shows the respiratory motion of the liver, while the transversal view provides clear liver anatomy and contrast. Supporting Information Video S1 further demonstrates the multiple contrasts from MT-ME.

In clinically healthy subjects, the mean, median, minimum, and maximum values for T1 (ms) are 851.7, 841.0, 718.5, 998.2; for T1<sub>w</sub> (ms) are 797.6, 814.5, 688.3, 875.4; for PDFF are 7.5%, 4.6%, 1.9%, 19.8%, and for R2\* (s<sup>-1</sup>) are 51.8, 50.5, 37.3, 72.9. In patient group, the mean, median, minimum, and maximum for T1 (ms) are 911.2, 911.8, 792.6, 990.3; for T1<sub>w</sub> (ms) are 829.8, 869.3, 679.2, 901.5; for PDFF is 10.1%, 9.8%, 4.9%, 15.9%; and for R2\* (s<sup>-1</sup>) is 60.6, 53.3, 44.2, 90.5.

Figure 5 displays the example multiparametric maps from MT-ME and references from a clinically healthy subject. The T1, PDFF, and R2\* maps generated by MT-ME are high-quality, co-registered and match well with the available reference maps (an in vivo  $T1_w$  reference was not available). Figure 6 shows the example multiparametric maps from a 64-year-old patient diagnosed with NASH. Elevated PDFF was detected by q-DIXON and MT-ME.

The mean and standard deviation of the scores of water images from MT-ME, VIBE-DIXON, and q-DIXON are  $4.7\pm0.5$ ,  $4.8\pm0.4$ , and  $3.8\pm0.5$ , respectively; the scores of fat images in the same order are  $4.8\pm0.4$ ,  $4.8\pm0.3$ , and  $4.1\pm0.3$ , respectively. MT-ME produced water and fat images with similar quality compared to VIBE-DIXON, and improved image quality and sharpness compared to q-DIXON. The scores of PDFF maps from MT-ME and q-DIXON are  $4.7\pm0.4$ ,  $4.1\pm0.2$ , respectively, indicating superior PDFF maps from MT-ME over q-DIXON. The scores of R2\* maps from MT-ME and q-DIXON are  $3.7\pm0.5$ ,  $3.8\pm0.3$ , respectively, indicating that the R2\* maps from MT-ME were similar quality to the maps from q-DIXON. The scores of T1, T1<sub>w</sub> maps from MT-ME and T1 maps from MOLLI are  $4.3\pm0.5$ ,  $4.4\pm0.5$  and  $4.9\pm0.3$ , respectively, which are rated good to excellent. Detailed image quality scores are listed in Supporting Information Table S2 and Table S3.

Supporting Information Figure S1 shows the comparison of the  $T1_w$  measurements using the proposed basis-function-based approach and using the straightforward approach applying the water/fat separation to the images of each inversion time. For the in vivo data, the ICC of the two measurements is 0.993 with a mean difference at 4.6 ms, indicating that the basis-function-based approach produced similar results compared to the straightforward approach.

However, the straightforward approach requires 3D water-fat separation on 192 inversion time images, while the basis-function-based approach requires 3D water-fat separation per each of 5 basis functions, which is 38 times faster.

The example maps of fat-specific T1 (T1<sub>f</sub>) are displayed in Supporting Information Figure S2. The mean and standard deviation of T1<sub>f</sub> is  $375.8\pm18.1$  in the clinically healthy group, and is  $379.5\pm11.2$  in the patient group, which are consistent with literature values<sup>45</sup>.

The importance of performing T1 correction on the PDFF estimation for the MT-ME technique was demonstrated in Figure 7. Figure 7A and 7B display two representative in vivo cases. Without T1 correction, the water proton density was underestimated due to the longer T1, resulting in overestimated PDFF. After performing T1 correction, the PDFF estimated by MT-ME matched well with the reference from q-DIXON. For all the in vivo datapoints (15 clinically healthy subjects and 4 patients), the T1 corrected PDFF from MT-ME showed increased Pearson correlation coefficient R and ICC with reference values (R = 0.979 and ICC = 0.935 without T1 correction; R = 0.993 and ICC = 0.990 with T1 correction).

The in vivo intra-session repeatability of T1, T1<sub>w</sub>, PDFF, and R2\* measurements of MT-ME were illustrated by the Bland-Altman plots in Figure 8A. The ICC of T1, T1<sub>w</sub>, PDFF, and R2\* were 0.989, 0.990, 0.999, 0.988, respectively, and the CoV were 0.9%, 0.7%, 1.3%, 1.4%, respectively. The high ICC and low CoV values of all parameters indicated a good intra-session repeatability. Figure 8B shows the regression analysis of MT-ME measurements against in vivo references of the clinically healthy cohort with the Pearson correlation coefficients R = 0.925, 0.992, 0.984 and ICC = 0.924, 0.987, 0.975 for T1, PDFF, and R2\*, respectively. The paired t-test showed that there was no significant difference for either PDFF (P= 0.853) or R2\* (P= 0.337) measurements. A significant difference was found in T1 measured from MT-ME and MOLLI (P<0.001). The overall T1 from MT-ME was 4.6% higher than the T1 of MOLLI, as listed in Table 1.

The regression analysis for the multiparametric measurements of patient cohort were shown in Figure 8C, with the Rs = 0.991, > 0.999, >0.999, and ICCs = 0.980, 0.999, 0.998 for T1, PDFF, and R2\*, respectively; the *P*-values from paired t-test were 0.027, 0.252, 0.075, indicating statistical difference in T1 measurement but no differences in PDFF and R2\*. Table 1 summarizes the mean and standard deviation measurements of all the parameters from MT-ME and reference for clinically healthy and patient cohorts.

The differences between T1 and T1<sub>w</sub> for each subject were analyzed using a dot-line diagram, as shown in Figure 9A, including all in vivo datapoints (15 clinically healthy volunteers and 4 patients). The joint T1 were estimated on the first echo images, whose TE = 1.25 ms is close to the out-of-phase of water and fat. Results showed that joint T1 was larger than the T1<sub>w</sub> by a different portion, which is consistent with previous findings<sup>28</sup>. The differences between T1 and T1<sub>w</sub> presented a significantly strong correlation with PDFF with the Pearson correlation coefficient R = 0.945 and *P*< 0.001, as shown in Figure 9B. On the contrary, the water-specific T1<sub>w</sub> appeared to be independent to PDFF (R = -0.029, *P*= 0.904), agreed with the expectation.

4.

The potential of multiparametric MRI in the characterization of liver tissues has been demonstrated in a variety of investigations. However, technical challenges, including the requirements for multiple breath-holds, misregistration among separate scans, lack of reliable 3D T1 mapping techniques, and confounder factors among the biomarkers have limited the clinical applications of multiparametric MRI. In this study, a novel

technique, MT-ME, was developed, achieving 5-min free-breathing acquisition with wholeliver coverage, a spatial resolution of  $1.5 \times 1.5 \times 6 \text{ mm}^3$ , and simultaneous, co-registered quantification of T1, T1<sub>w</sub>, PDFF, and R2\*.

A major advantage of the proposed technique is the free-breathing acquisition and motionresolved reconstruction. In patient populations with breath-hold difficulties such as seriouscondition, pediatric, sedated, or elderly patients, a free-breathing technique robust to respiratory motion is a desirable and useful clinical option. There are free-breathing techniques available to quantify PDFF and R2\* with a stack-of-star sampling pattern<sup>46,47</sup> and respiratory averaging. However, the unresolved respiratory motion can lead to reduced image quality and estimation error for PDFF and R2\*<sup>46,47</sup>. In our work, the respiratory motion was captured by training data acquired at the center k-space every 72 ms, and modeled by the Multitasking framework, yielding high-quality images and quantitative maps comparable to breath-hold references.

Another advantage of MT-ME is the 3D coverage with high spatial resolution, which can potentially capture the spatial distribution of fibrosis stage, PDFF and R2\* between different segments of liver<sup>48–50</sup>, which can be used for disease differentiation<sup>50</sup>. Moreover, the association between the fibrosis and fat concentration and the development of hepatocellular carcinoma are gaining increased attention in recent years<sup>51,52</sup>. The MT-ME technique with full-liver coverage and high spatial resolution has the potential to capture the nodules and lesions, which can be a promising tool to study this association.

The phantom experiments on a commercial 15-vial fat and R2\* phantom showed that the quantification of MT-ME matched well with the gold-standard references. The T1 of vial 13, which is around 2300 ms by IR-SE, was underestimated by MT-ME. This may result from the fact that the sequence parameters were designed for the mapping of liver T1, whose range was typically reported from 700 to 1200 ms on 3T systems<sup>9,20,45</sup>.

The in vivo experiments demonstrated that T1, PDFF, and R2\* from MT-ME were consistent with references for healthy and patient cohorts. q-DIXON is a well-validated product technique on commercial MRI systems from a main MRI vendor for PDFF and R2\* quantification with good reproducibility and clinical utility<sup>49,53–55</sup>. The PDFF and R2\* measured by MT-ME were consistent with q-DIXON results with no significant differences. The T1 estimated by MT-ME was 4.6% and 4.8% higher than MOLLI in healthy and patient cohorts with P < 0.001, = 0.028, respectively. However, MOLLI itself can only serve as a suboptimal reference for in vivo T1 measurement due to its tendency to underestimate T1<sup>56,57</sup>. A major factor is the magnetization transfer effect between the free pool and bound pool, which can lead to the underestimation of T1 at 100 to 200 ms<sup>58</sup>. The high quality of

the maps produced by MT-ME and consistency with references demonstrated its capability to achieve respiratory-motion-resolved quantification of T1, PDFF, and R2\*, which has a great clinical potential to relief the burdens of breath-hold from patients and improve the mapping quality and reproducibility.

The PDFF estimation in MT-ME used the multi-echo images at the last TI. Given the IR period of 1800 ms, water proton in liver with the T1 around 800–900 ms cannot reach the steady-state at the last TI, while fat proton can almost achieve steady-state. This T1 effect can result in the underestimation of water proton density, leading to the overestimation of PDFF. By introducing T1 correction, the agreement between PDFF of MT-ME and reference was substantially improved with increased Pearson Correlation Coefficient R (from 0.979 to 0.993) and ICC (from 0.935 to 0.990).

Intra-session repeatability is an important indicator of the precision of the quantitative techniques<sup>59,60</sup>. The phantom studies produced excellent intra-session repeatability for T1, T1<sub>w</sub>, PDFF, and R2\* with high ICC (>=0.999 for all parameters) and low CoV (<1.5% for all parameters). In the volunteer studies, high intra-session repeatability was also demonstrated with ICC > 0.980 and CoV < 1.5% for all parameters with the presence of respiratory motion. As observed in the study, the respiratory pattern can change over time within the same imaging session. The high intra-session repeatability strongly supported that MT-ME is reliable for respiratory-motion-resolved quantification of T1, T1<sub>w</sub>, PDFF, and R2\* and has the potential to serve as a clinical tool for diagnosis, staging, and longitudinal monitoring of liver diseases. Future studies to evaluate the inter-session repeatability and inter-vendor reproducibility of MT-ME will be conducted.

The estimation of liver T1 is affected by the presence of fat. As reported previously, when TE of MOLLI is close to the out-of-phase point of water/fat, the measured T1 tends to be longer than the true T1 of water pool; the bias grows with the increase of PDFF<sup>28</sup>. This can impair the reliability of T1 in the assessment of fibrosis. It is possible to correct the fat effect on the MOLLI/shMOLLI T1, but at the cost of separate mapping of off-resonance frequency and fat content in combination with numerical simulations<sup>26</sup>. In this work, the estimation of water-specific T1<sub>w</sub> was performed efficiently on the reconstructed basis functions. The validation on the phantom study showed that the T1<sub>w</sub> agreed well with water-excited IR-SE with R = 0.996, ICC = 0.994, and *P* = 0.097. For the in vivo study, the joint T1 was estimated on the first echo images, the TE of which is 1.25 ms, close to the out-of-phase TE for water and fat. The observation indicated that the joint T1 was longer than T1<sub>w</sub> for all cases, consistent with the trend reported in Mozes et al<sup>28</sup>. The differences between joint T1 and T1<sub>w</sub> showed a strong dependency over PDFF (R = 0.945, *P*<0.001), while T1<sub>w</sub> itself was independent to PDFF (R = -0.029, *P* = 0.904). This is a solid demonstration of the ability of T1<sub>w</sub> to avoid errors introduced by fat.

A technical limitation of the current MT-ME method is the dependency of T1 and T1<sub>w</sub> on B<sub>1</sub><sup>+</sup>. It can cause estimation bias and increased intra-subject variation, as analyzed in Supporting Information Section D and Supporting Information Figure S4. To reduce the T1 dependency on B<sub>1</sub><sup>+</sup>, a small flip angle, which is 5°, and a long TR, which is 9.3 ms were chosen in this work<sup>61</sup>. Supporting information Figure S5 displays the measured intra-subject

standard deviation of T1 and T1<sub>w</sub> for all subjects. The root-mean-squares of the intra-subject standard deviation are 71.6 ms and 59.4 ms for T1 and T1<sub>w</sub> respectively, larger than the values reported in Thompson et al.<sup>9</sup>, which corrected B<sub>1</sub><sup>+</sup> inhomogeneity. Nevertheless, T1<sub>w</sub> measurements from MT-ME in this work are still sufficient to differentiate normal liver from medium- to high-stage fibrosis or steatosis of liver. According to literature, the differences of T1 is larger than 150 ms between normal liver versus fibrotic liver with fibrosis stage >=3, or between normal liver versus steatosis liver<sup>12,62,63</sup>, more than two times of the root-mean-squares of the intra-subject standard deviation for both T1 and T1<sub>w</sub>. In the future, technical improvement to resolve B<sub>1</sub><sup>+</sup>-related variation of T1 and T1<sub>w</sub> measurements will be investigated. One possible approach is to acquire a separate low-resolution B<sub>1</sub><sup>+</sup> map. Another possible approach is to incorporate dual-flip angle acquisition to map B<sub>1</sub><sup>+</sup> and T1 together<sup>64–66</sup>.

There are several limitations of the study. First, the clinically healthy group comprised subjects without clinical diagnosis of liver diseases. However, given the high prevalence of liver diseases, it is possible that these subjects also possessed liver abnormalities to some extent, which may contribute to the large variation of the T1, PDFF, and R2\* values within the group. For the patient study, the sample size is small. The four recruited patients had varied types and stages of diseases (2 patients with NAFLD and 2 with NASH). The overall PDFF values in patient group are relatively low (the median is 9.8% and the maximum is 15.9%), which was unable to show the feasibility of the multiparametric quantification from MT-ME under high-PDFF condition. A possible reason is that the patients had different durations of disease history and a variety of treatments which may have changed the pathological characteristics of liver to a different extent. Furthermore, the histology biomarkers including the fibrosis, fat concentration, and iron concentration were not available for this pilot study. The cross-correlation analysis between quantitative parameters from MT-ME and pathological markers were not accessible. Besides, the intrasession repeatability of MT-ME was not available in the patient group due to the additional time constraints associated with clinical scans.

In the future, studies will be performed on larger patient cohorts with histological validation. The inter-session repeatability and inter-vendor reproducibility of MT-ME will be assessed. Iron is another confounder of T1 measurement in liver. Increased iron concentration can directly reduce T1<sup>67</sup>. Co-registered, simultaneous measurement of T1<sub>w</sub> and R2\* using MT-ME provides the opportunity to develop and incorporate an R2\* based iron-compensated algorithm<sup>68</sup> into the MT-ME pipeline for the estimation of fat-independent, iron-compensated T1, which may serve as a better indicator of liver fibrosis scores. In addition, advanced shimming techniques to improve B0 field homogeneity can be potentially integrated into the application of MT-ME to further enhance the image quality and precision of the multiparametric quantification<sup>69,70</sup>.

#### 5. Conclusion

A novel 3D MT-ME technique was developed, enabling simultaneous whole-liver quantification of joint T1, water-specific T1 (T1<sub>w</sub>), PDFF, and R2\* within a 5-minute free-breathing acquisition. The T1, T1<sub>w</sub>, PDFF, and R2\* estimated by MT-ME showed great

intra-session repeatability and substantial agreement with reference values in a phantom and in vivo subjects. The proposed MT-ME technique is clinically promising for the comprehensive characterization of liver tissue properties.

#### **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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#### Figure 1:

(A) Pulse sequence diagram for the proposed MT-ME technique. Non-selective inversion recovery(IR) preparation pulse is applied periodically followed by 3D SPGR readouts. Six-echo bipolar gradients were used for data collection. The center k-space readout was set on kz (superior-inferior) direction. Pre-phaser and spoiler of readout was not included in the simplified figure. (B): Simplified illustration of k-space sampling strategy. A stack-of-star sampling pattern with golden-angle in-plane and Gaussian-density randomized reordering in kz direction was implemented. The training data formed with center k-space lines was acquired every 8 readouts in kz (superior-inferior) direction for better capture of respiratory motion. Rest of the data forms the imaging data.



#### Figure 2:

Flow chart of the reconstruction of MT-ME images and the quantification of T1, T1<sub>w</sub>, PDFF, and R2\*.

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#### Figure 3:

Phantom Measurements. (A) T1, T1<sub>w</sub>, PDFF, and R2\* maps from reference sequences and MT-ME. The Inversion-recovery spin-echo (IR-SE) sequence without and with water excitation were used as reference for joint and water-specific T1. q-DIXON provided the reference for PDFF and R2\*. The quantitative maps from MT-ME showed great agreement with references. (B) Bland-Altman plots shows good repeatability of T1, T1<sub>w</sub>, PDFF, and R2\* estimated with MT-ME. (C) Good correlation was demonstrated of T1, T1<sub>w</sub>, PDFF, and R2\* from MT-ME and reference with high Pearson correlation coefficient *R* and ICC, as labeled on each plot.



#### Figure 4:

Demonstration of MT-ME images with three spatial dimensions and three temporal dimensions (IR dimension, multi-echo dimension, and respiratory dimension). The images of end-inspiration and end-expiration were displayed in coronal view at the last TI of echo1, echo 2, and echo 6. The red solid reference line on images of echo 1 is the position of liver dome at end-expiration, while the dashed reference line is the position at end-inspiration. Images of different TI times (266.8 ms and 1766.4 ms) were displayed in axial view. The liver and vessels were well delineated.



#### Figure 5:

Representative anatomical water/fat images and T1, T1<sub>w</sub>, PDFF, and R2\* maps from a clinically healthy 29-year-old volunteer. (A) Water and fat images from q-DIXON, VIBE-DIXON, and MT-ME. Vessels can be well defined on water images of MT-ME and VIBE-DIXON. (B) Good consistency was showed between MT-ME maps and references. The in vivo T1 reference was 2D MOLLI with a 11-second breath-hold, while the PDFF and R2\* reference was q-DIXON with 12-second breath hold. The T1<sub>w</sub> reference was not available.



#### Figure 6:

Representative water/fat images and T1, T1<sub>w</sub>, PDFF, and R2\* maps from a 64-year-old patient with NASH. (A) Water and fat images from q-DIXON, VIBE-DIXON, and MT-ME. (B) T1, T1<sub>w</sub>, PDFF and R2\* maps. The mean T1, PDFF, and R2\* of reference were 779 ms, 15.5%, 89 s<sup>-1</sup>, respectively, while the mean T1, T1<sub>w</sub>, PDFF, and R2\* measured from MT-ME were 785 ms, 643 ms, 15.5%, and 85 s<sup>-1</sup>.

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#### Figure 7:

The significance of T1 correction in the estimation of PDFF. Two in vivo examples are display in (A) and (B). Without T1 correction (middle panel of A and B), the PDFF estimated from MT-ME is higher than q-DIXON reference. With correction (right panel of A and B), the PDFF estimated from MT-ME matches well with q-DIXON results. (C) the regression analysis between uncorrected MT-ME PDFF and q-DIXON shows R = 0.979, ICC = 0.935; the regression between corrected MT-ME PDFF and q-DIXON shows improved correlation R = 0.993 and increased ICC = 0.990.

(A) Intra-session repeatability of T1, T1<sub>w</sub>, PDFF, and R2\*



(A) Bland-Altman plots demonstrated a good intra-session repeatability of the multiparametric mapping from MT-ME. The overall intra-session differences for T1 and T1<sub>w</sub> were less than 3%, while the differences for PDFF and R2\* were less than 10%. (B) The regression analysis of T1, PDFF, and R2\* measured with MT-ME and references showed good agreement in the clinically healthy subjects. The Pearson correlation coefficient *R* was 0.990, 0.976, and 0.953, respectively. (C) The regression analysis of T1, PDFF, and R2\* measured with MT-ME analysis of T1, PDFF, and R2\* measured with MT-ME and references showed good agreement in patients. The Pearson correlation coefficient *R* was 0.998, >0.999, and >0.999, respectively.



#### Figure 9:

(A) Dot-line diagram of the relation between T1 and T1<sub>w</sub> estimated on all subjects using MT-ME. The T1<sub>w</sub> appears lower than joint T1 in all subjects. (B) regression of the differences of T1 and T1<sub>w</sub> against PDFF. A significantly strong correlation with R = 0.960, P < 0.001 is demonstrated. (C) No correlation was found between T1<sub>w</sub> against PDFF, with R = 0.016, P = 0.781.

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#### Table 1:

Mean and standard deviation measurements of T1,  $T1_w$ , PDFF, and R2\* from MT-ME and reference, the absolute relative differences between MT-ME reference, and P value of paired t-test are provided for clinically healthy and patient groups.

		T1 [ms]	T1 <sub>w</sub> [ms]	PDFF [%]	R2* [s-1]
Clinically healthy	MT-ME	851.7±80.2	797.6±59.3	7.1±6.2	52.0±10.7
	References	MOLLI	N/A	q-DIXON	q-DIXON
		814.7±78.1		7.2±6.6	51.6±11.5
	Absolute relative differences [%]	4.6%	N/A	1.4%	0.8%
	Р	< 0.001 *	N/A	0.853	0.337
Patient	MT-ME	911.2±83.2	829.8±119.5	10.0±5.6	55.9±20.0
	References	MOLLI	N/A	q-DIXON	q-DIXON
		869.3±68.0		9.9±5.7	57.8±21.2
	Absolute relative differences [%]	4.8%	N/A	1.0%	3.3%
	Р	0.028*	N/A	0.252	0.075

indicates statistical significance.