

**Document Version**

Final published version

**Licence**

CC BY

**Citation (APA)**

Starekova, J., Weingärtner, S., Rutkowski, D. R., Fullerton, G. C., Bae, W. C., Do, H. P., Madhuranthakam, A. J., Malis, V., Hernando, D., & More Authors (2026). Cross-Vendor Validation of Proton Density Fat Fraction and T<sub>1</sub> Mapping Using a Combined Proton Density Fat Fraction—T1 Phantom. *Journal of Magnetic Resonance Imaging*, 64(2), 482-492. <https://doi.org/10.1002/jmri.70344>

**Important note**

To cite this publication, please use the final published version (if applicable).  
Please check the document version above.

**Copyright**

In case the licence states “Dutch Copyright Act (Article 25fa)”, this publication was made available Green Open Access via the TU Delft Institutional Repository pursuant to Dutch Copyright Act (Article 25fa, the Taverne amendment). This provision does not affect copyright ownership.  
Unless copyright is transferred by contract or statute, it remains with the copyright holder.

**Sharing and reuse**

Other than for strictly personal use, it is not permitted to download, forward or distribute the text or part of it, without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license such as Creative Commons.

**Takedown policy**

Please contact us and provide details if you believe this document breaches copyrights.  
We will remove access to the work immediately and investigate your claim.

## RESEARCH ARTICLE OPEN ACCESS

# Cross-Vendor Validation of Proton Density Fat Fraction and $T_1$ Mapping Using a Combined Proton Density Fat Fraction— $T_1$ Phantom

Jitka Starekova<sup>1</sup>  | Sebastian Weingärtner<sup>2</sup>  | David R. Rutkowski<sup>3</sup>  | Garrett C. Fullerton<sup>1,4</sup>  | Won C. Bae<sup>5</sup> | Hung P. Do<sup>6</sup>  | Ananth J. Madhuranthakam<sup>7,8</sup>  | Vadim Malis<sup>5</sup> | Sheng Qing Lin<sup>7</sup>  | Suraj D. Serai<sup>9</sup>  | Takeshi Yokoo<sup>7</sup>  | Scott B. Reeder<sup>1,4,10,11,12</sup>  | Jean H. Brittain<sup>3</sup>  | Diego Hernando<sup>1,4</sup> 

<sup>1</sup>Department of Radiology, University of Wisconsin-Madison, Madison, Wisconsin, USA | <sup>2</sup>Department of Imaging Physics, Delft University of Technology, Delft, the Netherlands | <sup>3</sup>Calimetrix, Madison, Wisconsin, USA | <sup>4</sup>Department of Medical Physics, University of Wisconsin-Madison, Madison, Wisconsin, USA | <sup>5</sup>Department of Radiology, University of California San Diego, San Diego, California, USA | <sup>6</sup>Canon Medical Systems USA, Tustin, California, USA | <sup>7</sup>Department of Radiology, University of Texas, Southwestern Medical Center, Dallas, Texas, USA | <sup>8</sup>Department of Radiology, Mayo Clinic in Rochester, Rochester, Minnesota, USA | <sup>9</sup>Department of Radiology, Children's Hospital of Philadelphia, University of Pennsylvania School of Medicine, Philadelphia, Pennsylvania, USA | <sup>10</sup>Department of Biomedical Engineering, University of Wisconsin-Madison, Madison, Wisconsin, USA | <sup>11</sup>Department of Medicine, University of Wisconsin-Madison, Madison, Wisconsin, USA | <sup>12</sup>Department of Emergency Medicine, University of Wisconsin-Madison, Madison, Wisconsin, USA

**Correspondence:** Diego Hernando ([dhernando@wisc.edu](mailto:dhernando@wisc.edu))

**Received:** 19 January 2026 | **Revised:** 10 April 2026 | **Accepted:** 14 April 2026

**Keywords:** CSE-MRI | PDFF | phantom | quantification | reproducibility |  $T_1$

## ABSTRACT

**Background:** In chronic liver disease, fat and fibroinflammatory changes often coexist. However, their biomarkers, proton density fat fraction (PDFF) and  $T_1$ , are typically assessed separately. Their reproducibility under mutual confounding remains unclear.

**Purpose:** To assess multicenter, multi-vendor reproducibility of confounder-corrected chemical shift-encoded (CSE)-MRI-based PDFF mapping and MOLLI-based  $T_1$  mapping using a combined PDFF- $T_1$  phantom.

**Study Type:** Prospective phantom study.

**Phantom:** Commercial PDFF- $T_1$  Phantom (Model 725) with varying PDFF (0%–30%) and  $T_1$  (200–1400 ms) values.

**Field Strength/Sequence:** 1.5 T and 3 T multi-echo, three-dimensional spoiled-gradient-echo (SGRE) sequence for PDFF mapping, and MOLLI sequence (5(3)3 acquisition scheme) using two-dimensional SGRE readouts for  $T_1$  mapping across four centers and vendors.

**Assessment:** PDFF and  $T_1$  maps were acquired using standardized protocols. PDFF maps were reconstructed locally, while  $T_1$  maps were generated using a centralized algorithm. All maps were quantitatively analyzed by a single radiologist using standardized region-of-interest placement. Phantom temporal stability was assessed at one center across five sessions over 9 months (baseline, retest, 1 week, 6 and 9 months).

**Statistical Tests:** Intraclass correlation coefficients (ICC), reproducibility coefficients (RDC), and linear regression analysis were used. A  $p$  value < 0.05 was considered statistically significant.

**Results:** PDFF showed overall excellent reproducibility (ICC = 0.987, RDC = 3.7%), with increased variability at higher  $T_1$  values (RDC up to 7.9% at  $T_1$  = 1400 ms).  $T_1$  mapping showed good reproducibility in the absence of fat (RDC 16–161 ms at PDFF = 0%),

This is an open access article under the terms of the [Creative Commons Attribution](https://creativecommons.org/licenses/by/4.0/) License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2026 The Author(s). *Journal of Magnetic Resonance Imaging* published by Wiley Periodicals LLC on behalf of International Society for Magnetic Resonance in Medicine.

but moderate to poor reproducibility in the presence of fat, with RDC increasing up to 1553 ms at PDFF 30%. Temporal stability was excellent  $ICC \geq 0.998$  for both PDFF and  $T_1$ , and RDC of 1.1%–1.3% for PDFF and 52–57 ms for  $T_1$ .

**Data Conclusion:** This phantom study demonstrated high reproducibility of PDFF, whereas  $T_1$  reproducibility deteriorated at higher fat and  $T_1$  levels, underscoring the need for fat-corrected  $T_1$  mapping for reliable assessment of fibroinflammatory changes.

**Evidence Level:** N/A.

**Technical Efficacy:** Stage 1.

## 1 | Introduction

Metabolic dysfunction-associated steatotic liver disease (MASLD) is the most prevalent chronic disease of the liver, affecting approximately 38% of the global adult population [1]. The earliest and hallmark feature of MASLD is excessive fat accumulation in the liver (steatosis). The presence of fat can progress to metabolic dysfunction-associated steatohepatitis (MASH), a more severe form involving inflammation, hepatocellular injury, and fibrosis [1, 2]. If left untreated, the condition may advance to cirrhosis, liver failure, and hepatocellular carcinoma. MASLD is also linked to a range of extrahepatic complications [1, 3].

Given its high prevalence and progressive nature, there is an unmet need for reliable noninvasive biomarkers to assess liver fat and fibroinflammatory changes in both clinical practice and research settings [4, 5]. Confounder-corrected chemical shift-encoded MRI (CSE-MRI) can measure proton density fat fraction (PDFF), a well-validated quantitative biomarker for liver steatosis [3, 6].  $T_1$  mapping has shown promise as a noninvasive biomarker for liver fibroinflammatory changes (via native  $T_1$ ) and function (via post-contrast  $T_1$  using hepatocyte-specific contrast agents) [7, 8].

However, both PDFF and  $T_1$  measurements can be affected by multiple confounding factors. The presence of fat leads to bias in  $T_1$  estimates when using conventional (fat-uncorrected) methods. Changes in  $T_1$  values, such as those caused by fibroinflammatory changes, can lead to bias (overestimation) of PDFF estimates if no appropriate mitigation method, such as low flip angle or  $T_1$ -correction, is employed [3, 9–12]. Reliable quantification of both steatosis and fibroinflammatory activity is important, as they have distinct implications for MASLD diagnosis, staging, and treatment strategies in the clinic.

Chemical shift encoded MRI (CSE-MRI)-based PDFF quantification is considered the leading noninvasive assessment method for quantifying liver steatosis [3]. Among existing  $T_1$  mapping techniques, modified Look-Locker inversion recovery (MOLLI) methods, originally developed for cardiac imaging, are widely available and have been used for liver imaging [13, 14]. However, MOLLI is susceptible to bias in the presence of fat [13]. Given the high prevalence of steatosis, the reproducibility of MOLLI-based  $T_1$  mapping in the clinical setting remains uncertain.

Multicenter studies enable highly controlled validation of quantitative MRI methods across centers, vendors, and platforms. PDFF and  $T_1$  phantoms have been used in several multicenter studies [15–20]. However, there is limited data on the reproducibility of PDFF and  $T_1$  mapping in the simultaneous presence of varying combinations of PDFF and  $T_1$  values.

Therefore, this study evaluated the multicenter, multi-vendor reproducibility of CSE-MRI-based PDFF and MOLLI-based  $T_1$  quantification at 1.5T and 3.0T using a combined quantitative PDFF- $T_1$  phantom.

## 2 | Materials and Methods

This prospective phantom study was conducted across four participating centers (University of Wisconsin–Madison, University of Texas Southwestern Medical Center, Children's Hospital of Philadelphia, and Canon Medical Systems USA's Training Academy, Irvine, California) from February 2022 to December 2022.

### 2.1 | PDFF- $T_1$ Phantom

A single commercial phantom (Model 725 Special Order PDFF- $T_1$  Phantom, Calimetrix, Madison, WI) was used in this study. The phantom includes 16 cylindrical vials (volume: 20 mL, length: 60 mm, outer diameter: 27 mm) arranged in an asymmetrical grid. Each vial was filled with a gel formulation designed to mimic a specific combination of nominal PDFF values (0%–30%) and nominal water-specific  $T_1$  (200–1400 ms), as specified by the phantom manufacturer (Figure 1). These values were selected to span the physiologically relevant range of PDFF and water-specific  $T_1$  values observed in the human liver, including both native and post-contrast conditions [6]. The phantom gels achieve water-specific  $T_1$  values that remain approximately the same at 1.5T and 3.0T. To improve magnetic field homogeneity, coil loading, and minimize artifacts, the phantom's spherical housing was filled with a doped water solution. Water solution doping shortens the  $T_1$  and  $T_2$  relaxation times to better match the background contrast of human tissue. Additionally, gel-based vials were used to reduce motion-related and susceptibility artifacts caused by air bubbles.

Reference PDFF values for each vial were defined by phantom construction and verified through measurements using a temperature-adjusted, confounder-corrected reconstruction of CSE-MRI data [21].  $T_1$  values in the fat-free vials were verified using a conventional inversion-recovery spin-echo sequence (flip angle = 90°, TE = 9 ms, TR = 7000 ms, TI = 50, 400, 900, 1800 ms).  $T_1$  estimates were obtained using magnitude-based exponential fitting across inversion times. Verification imaging at 3.0T yielded values of  $195 \pm 1.7$  ms,  $619 \pm 3.8$  ms,  $1025 \pm 16.4$  ms, and  $1435 \pm 19.6$  ms. For vials with fat content, water-specific  $T_1$  was assumed to be the same as in the absence of fat (i.e., the nominal water-specific  $T_1$  values were assumed to be constant

### Plain Language Summary

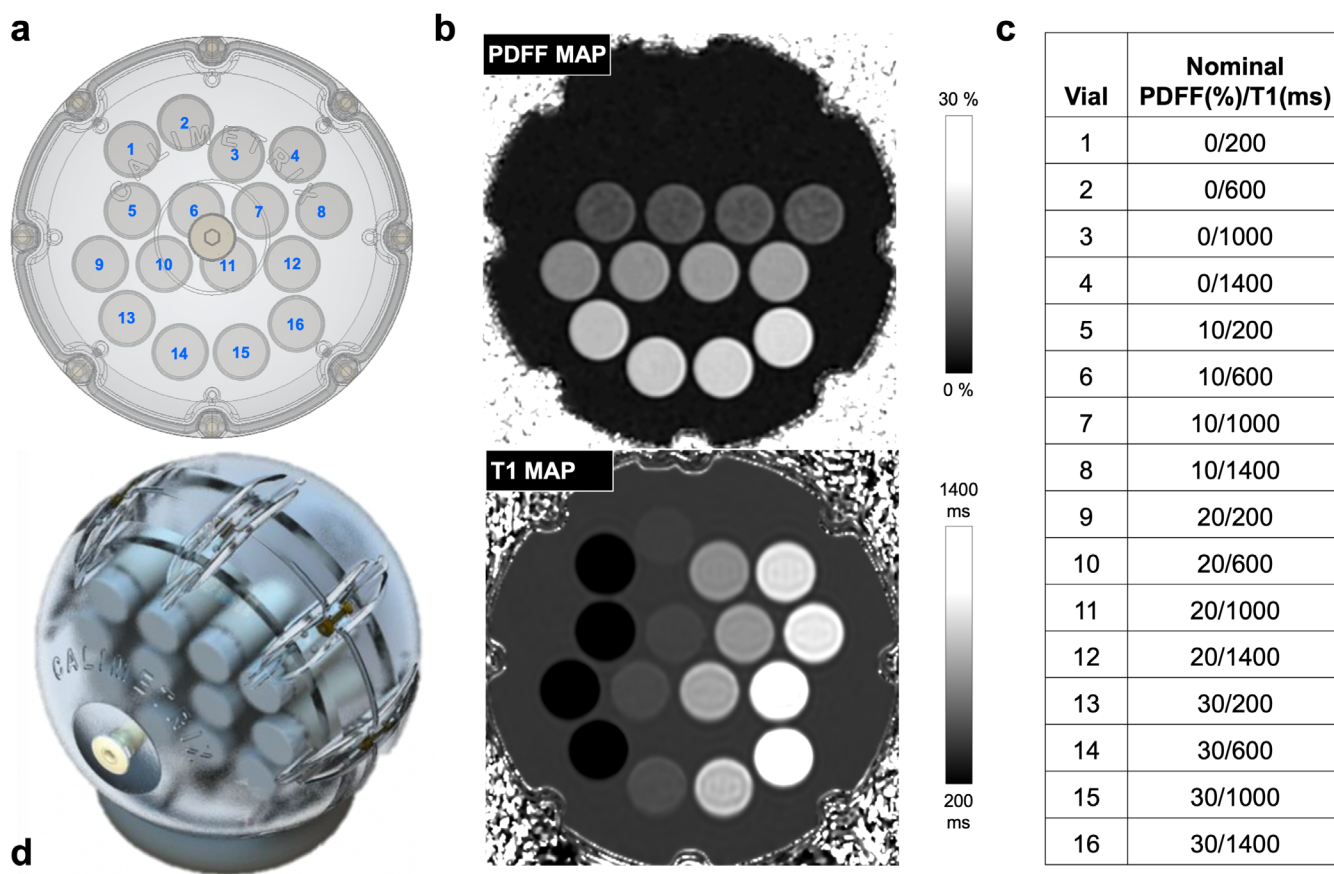
Magnetic resonance imaging (MRI) is used to measure liver fat and T1, a marker of tissue damage and inflammation. However, fat can interfere with T1 measurements. To test the reliability of these scans, we sent a specialized test object with known fat and T1 values to four hospitals. We found that while fat measurements were consistent across different MRI machines, T1 measurements varied significantly, especially when high levels of fat were present. This shows that standard MRI methods need improvement to ensure patients receive reliable liver disease assessments, regardless of which hospital or machine they visit.

across PDFF values) [10]. Vial verification was performed in accordance with standard Calimetrix Quality Management System (QMS) procedures.

## 2.2 | Imaging Experiments

The study involved four centers, each equipped with MRI systems from a different vendor (Figure 1). Specifically, the following vendors and systems were included: GE HealthCare 1.5T Signa Artist and 3.0T Signa Premier (Waukesha, WI, USA) at University of Wisconsin–Madison (center 1); Philips Healthcare 1.5T and 3.0T Ingenia systems (Amsterdam, Netherlands) at University of Texas Southwestern Medical Center (center 2); Siemens Healthineers 1.5T Magnetom Avanto and 2.89T Magnetom Skyra (Erlangen, Germany) at Children's Hospital of Philadelphia (center 3); and Canon Medical Systems 1.5T Vantage Orian and 2.89T Vantage Galan (Ottawa, Tochigi, Japan) at Canon Medical Systems USA's Training Academy, Irvine, California (center 4).

Two quantitative techniques were employed [1]: a 3D multi-echo, spoiled-gradient-echo (SGRE) sequence for PDFF mapping, and [2] a MOLLI sequence with 5(3)3 acquisition scheme using 2D



| Field strength   Vendor | GE HealthCare | Philips Healthcare | Siemens Healthineers | Canon Medical Systems |
|-------------------------|---------------|--------------------|----------------------|-----------------------|
| 1.5T                    | Signa Artist  | Ingenia            | Magnetom Avanto      | Vantage Orian         |
| 3T                      | Signa Premier | Ingenia            | Magnetom Skyra*      | Vantage Galan*        |

**FIGURE 1** | Illustration and photograph of the PDFF-T<sub>1</sub> phantom (a) with corresponding PDFF and T<sub>1</sub> map (b). The phantom was designed as an array of 16 vials immersed in a doped fill solution, contained within a spherical housing. Each vial contained a distinct combination of PDFF and water-specific T<sub>1</sub> values (nominal values) as shown in the table (c). The phantom was imaged at four different centers with different MR vendors (d). Abbreviation: PDFF, proton density fat fraction. \*2.89T.

SGRE readouts for  $T_1$  mapping. A low flip angle SGRE readout was used with MOLLI to minimize fat- and  $B_0$  inhomogeneity-related effects on  $T_1$  estimates [22]. Attempts were made to match acquisition parameters across centers and vendors; however, exact alignment was constrained by hardware and pulse sequence variations among vendors. Of note,  $T_1$  mapping could not be performed on the Philips 1.5T system at Center 2 as the SGRE-based MOLLI sequence was unavailable on that specific platform.

For PDFF quantification, centers were instructed to match the following acquisition parameters: number of slices 14, slice thickness of 4 mm,  $TE_1$  of 1.0–1.2 ms (for both 1.5 T and 3 T MRI systems),  $\Delta TE$  of 1.8–2.1 ms for 1.5 T and 0.8–1.0 ms for 3 T, six

echoes acquired in a single echo train for 1.5 T and two interleaved echo trains for 3 T, and a flip angle of 3° for both 1.5 T and 3 T (Table 1).

For  $T_1$  quantification, centers were instructed to use the following acquisition parameters: number of slices 1 through the middle of the vials, slice thickness of 4 mm,  $TI_1$  of 97–137 ms for 1.5 T and 97–115 ms for 3 T MRI systems,  $\Delta TI$  of 80 ms for both 1.5 T and 3 T, TR of 2.8–3.7 ms for both 1.5 T and 3 T, flip angle of 12° for 1.5 T and 10° for 3 T, 5(3)3 acquisition scheme (Table 2).

The phantom was shipped to participating centers via an overnight courier service in a protective, foam-padded case to reduce the risk of damage during transit (Figure S1). All centers

**TABLE 1** | Imaging parameters for proton density fat fraction (PDFF) mapping acquisition.

| Parameter                     | Recommended          | Center 1<br>GE HealthCare | Center 2<br>Philips<br>Healthcare | Center 3<br>Siemens<br>Healthineers | Center 4<br>Canon Medical<br>Systems |
|-------------------------------|----------------------|---------------------------|-----------------------------------|-------------------------------------|--------------------------------------|
| 1.5 T                         |                      |                           |                                   |                                     |                                      |
| Coil/channels (n)             | Head or head-neck    | Head-neck/21              | Head/15                           | Head/8                              | Head-neck/16                         |
| Matrix size                   | 148 × 148            | 148 × 148                 | 144 × 144                         | 128 × 128                           | 144 × 144                            |
| FOV (cm <sup>2</sup> )        | 26 × 26              | 26 × 26                   | 26 × 26                           | 26 × 26                             | 26 × 26                              |
| Slice thickness (mm)          | 4                    | 4                         | 4                                 | 4                                   | 4                                    |
| First TE (ms)                 | 1.0 to 1.2           | 1.1                       | 1.2                               | 1.11                                | 1.2                                  |
| TE spacing (ms)               | 1.8 to 2.1           | 1.9                       | 1.3                               | 2.0                                 | 1.2                                  |
| TR (ms)                       | Min <sup>a</sup>     | 23.0                      | 8.9                               | 12.5                                | 8.8                                  |
| No. of echo trains/<br>echoes | 1/6                  | 1/6                       | 1/6                               | 1/6                                 | 1/6                                  |
| FA (degrees)                  | 3                    | 3                         | 5                                 | 3                                   | 3                                    |
| Pixel BW (Hz)                 | 1351                 | 1351                      | 1578                              | 1085                                | 1302                                 |
| 3 T <sup>c</sup>              |                      |                           |                                   |                                     |                                      |
| Coil/channels (n)             | Head or head-neck    | Head-neck/19              | Head/32                           | Head/8                              | Head-neck/16                         |
| Matrix size                   | 140–160 ×<br>140–160 | 160 × 160                 | 164 × 163                         | 128 × 128                           | 144 × 144                            |
| FOV (cm <sup>2</sup> )        | 26 × 26              | 26 × 26                   | 26 × 26                           | 26 × 26                             | 26 × 26                              |
| Slice thickness (mm)          | 4                    | 4                         | 4                                 | 4                                   | 4                                    |
| First TE (ms)                 | 1.0–1.2              | 1.1                       | 1.2                               | 1.11                                | 1.2                                  |
| TE spacing (ms)               | 0.8 to 1.0           | 0.9                       | 1.0                               | 2.0                                 | 1.0                                  |
| TR (ms)                       | Min <sup>a</sup>     | 7.3                       | 7.5                               | 12.5                                | 7.6                                  |
| No. of echo trains/<br>echoes | 2/6                  | 2/6                       | 1/6                               | 1/6 <sup>b</sup>                    | 1/6                                  |
| FA (degrees)                  | 3                    | 3                         | 3                                 | 3                                   | 3                                    |
| Pixel BW (Hz)                 | 1136 to 1429         | 1136                      | 1494                              | 1090                                | 1302                                 |

Abbreviations: BW, bandwidth; FA, flip angle; FOV, field of view; No, number; TE, echo time; TR, repetition time.

<sup>a</sup>Minimum achievable with the provided echo times and other system constraints.

<sup>b</sup>Bipolar readout.

<sup>c</sup>2.89 T for Siemens and Canon.

**TABLE 2** | Imaging parameters for T<sub>1</sub> mapping acquisition.

| Parameter              | Recommended       | Center 1<br>GE HealthCare | Center 2<br>Philips<br>Healthcare | Center 3<br>Siemens<br>Healthineers | Center 4<br>Canon Medical<br>Systems |
|------------------------|-------------------|---------------------------|-----------------------------------|-------------------------------------|--------------------------------------|
| <b>1.5 T</b>           |                   |                           |                                   |                                     |                                      |
| Coil/channels (n)      | Head or head-neck | Head-neck/21              | n/a                               | Head/8                              | Head-neck/16                         |
| FOV (cm <sup>2</sup> ) | 26 to 36          | 36 × 27                   |                                   | 26 × 21                             | 36 × 27                              |
| Matrix size            | 128 × 128         | 128 × 128                 |                                   | 128 × 128                           | 128 × 128                            |
| Slice thickness (mm)   | 6 to 8            | 8                         |                                   | 8                                   | 6                                    |
| TR (ms)                | 2.8 to 3.7        | 3.1                       |                                   | 3.6                                 | 3.6                                  |
| Initial TI (ms)        | 97 to 137         | 98                        |                                   | 100                                 | 97                                   |
| ΔTI (ms)               | 80                | 80                        |                                   | 100                                 | 80                                   |
| Pixel BW (Hz)          | 480–980           | 488                       |                                   | 1090                                | 977                                  |
| FA (degrees)           | 12                | 12                        |                                   | 12                                  | 12                                   |
| Acquisition scheme     | 5(3)3             | 5(3)3                     |                                   | 5(3)3                               | 5(3)3                                |
| <b>3 T<sup>a</sup></b> |                   |                           |                                   |                                     |                                      |
| Coil/channels (n)      | Head or head-neck | Head-neck/21              | Head/15                           | Head/8                              | Head-neck/16                         |
| FOV (cm <sup>2</sup> ) | 26 to 36          | 36 × 27                   | 30 × 24                           | 26 × 21                             | 36 × 27                              |
| Matrix size            | 128 × 128         | 128 × 128                 | 132 × 126                         | 128 × 128                           | 128 × 128                            |
| Slice thickness (mm)   | 6 to 8            | 8                         | 8                                 | 8                                   | 8                                    |
| TR (ms)                | 2.8 to 3.7        | 3.1                       | 3.0                               | 3.7                                 | 3.3                                  |
| Initial TI (ms)        | 97 to 115         | 103                       | 85                                | 100                                 | 97                                   |
| ΔTI (ms)               | 80                | 80                        | 80                                | 100                                 | 80                                   |
| Pixel BW (Hz/pixel)    | 480–980           | 488                       | 553                               | 781                                 | 977                                  |
| FA (degrees)           | 10                | 10                        | 10                                | 10                                  | 10                                   |
| Acquisition scheme     | 5(3)3             | 5(3)3                     | 5(3)3                             | 5(3)3                               | 5(3)3                                |

Abbreviations: BW, bandwidth; FA, flip angle; FOV, field of view; T<sub>1</sub>, inversion time; TR, repetition time.<sup>a</sup>2.89 T for Siemens and Canon.

were instructed to store the phantom at room temperature when not in use and to allow it to acclimate to the MRI room temperature for at least 30 min before data acquisition, to ensure optimal image quality and consistency. The phantom was leveled within the head or head-neck coil using foam pads and a manufacturer-provided rubber base (where compatible) to accommodate varying coil geometries and secure the phantom to prevent movement during data acquisition. Reproducible positioning was ensured by aligning the system's laser crosshairs with the phantom's integrated orientation markings (cross) to place the phantom at the magnet isocenter.

The temperature of the phantom was measured just before imaging using a self-adhesive temperature-indicating strip (Cole-Parmer, Vernon Hills, IL, USA). Upon return to Center 1 after multicenter imaging, the phantom housing and vials were visually inspected by the study coordinator to ensure no damage had occurred during transit prior to stability testing.

To evaluate the stability of the phantom's PDFF and T<sub>1</sub> values over time, multiple repeated acquisitions were performed at Center 1

using 1.5 T and 3.0 T systems above, following the same imaging protocols described above using the same hardware and software. Two exams were conducted during the initial session on the same day: a baseline test and a retest exam performed following repositioning of the phantom and reconnection of the coil. Follow-up exams were conducted at 1 week, 6 months, and 9 months (Figure S1).

## 2.3 | Image Processing and Quantitative Analysis

PDFF maps were automatically reconstructed online at the respective centers using the vendor-provided reconstruction software and subsequently transferred to Center 1 for analysis. Because vendor-specific T<sub>1</sub> map reconstruction was not available for some systems, all T<sub>1</sub> maps from the participating centers were processed centrally at Center 1 using a standardized MOLLI fitting algorithm implemented in Matlab (MathWorks, Natick, MA) [23, 24].

An experienced radiologist (16 years of MRI experience, JS) used OsiriX DICOM viewer (v14.1.2 Pixmeo, Bernex, Geneva, Switzerland) to place circular regions of interest (ROIs), 1.9 cm in

diameter, on each of the 16 vials per slice. For the PDFF maps, ROIs were placed on three consecutive central slices, and voxel values were averaged across them. For the  $T_1$  map, ROIs were placed on a single central slice, and the mean voxel value was recorded.

## 2.4 | Validation of Phantom Homogeneity

To verify the spatial homogeneity of the phantom and provide an independent reference for water-specific  $T_1$  in the presence of fat, an additional validation was performed at Center 1 (3.0T). A saturation-recovery chemical shift-encoded (SR-CSE) method was used to achieve simultaneous fat-water separation and  $T_1$  mapping of the water compartment (water-specific  $T_1$ ) [25]. Acquisition parameters included: FOV  $28 \times 28 \text{ cm}^2$ ,  $128 \times 96$  matrix,  $2 \times$  auto-calibrated parallel imaging in the phase-encoding direction, 8.59 ms TR, 5 monopolar echoes per TR ( $TE_1 = 1.24 \text{ ms}$ ,  $\Delta TE = 1.41 \text{ ms}$ ), 3 saturation-recovery preparations ( $TS = [200 \text{ ms}, 715 \text{ ms}, \text{ and } 1231 \text{ ms}]$ ) and one acquisition without saturation preparation, 4 mm slice thickness, 6 mm gap between adjacent slices, averaged across 5 NEX. To assess phantom homogeneity, standardized ROIs were drawn in three spatially separated slices of water-specific  $T_1$  maps along the length of each vial. Mean and standard deviation of water-specific  $T_1$  measurements were computed for each vial across the three slices.

## 2.5 | Statistical Analysis

Statistical analysis was performed using R (v4.1.0., tidyverse v1.3.1, ggplot2 v3.3.6, irr v0.84.1, rstatix v0.7.0). For the multicenter, multi-vendor validation of PDFF and  $T_1$  mapping, the intraclass correlation coefficients (ICC) were calculated to assess overall agreement across scanners and vendors. ICC values were interpreted as  $< 0.50 = \text{poor}$ ,  $0.50\text{--}0.75 = \text{moderate}$ ,  $0.75\text{--}0.90 = \text{good}$ , and  $> 0.90 = \text{excellent}$  [26]. Reproducibility coefficients (RDC) were calculated to quantify measurement precision. Overall RDC values were calculated by pooling data across all vials within specific nominal ranges (PDFF: 0%–30%;  $T_1$ : 200–1400 ms) to provide a global estimate of measurement variation. Linear regression was performed using measured PDFF/ $T_1$  values as the dependent variables and corresponding nominal phantom values as the independent variables to quantify linear association. To assess measurement consistency over time (i.e., phantom stability), the ICC and RDC were similarly calculated. For the validation of phantom homogeneity, the mean, standard deviation, coefficient of variation (CV), and ICC were calculated.

Outlier detection was performed using the interquartile range (IQR) method. Data points were considered outliers if they fell below the first quartile or above the third quartile by more than 1.5 times the IQR and classified as extreme outliers if they exceeded three times the IQR.

For all tests,  $p < 0.05$  was considered statistically significant.

## 3 | Results

PDFF and  $T_1$  mapping data were successfully acquired at all centers except for  $T_1$  mapping at Center 2 (1.5T Philips). The

measured phantom temperature prior to imaging ranged from  $18^\circ\text{C}$  to  $23^\circ\text{C}$  across all centers and systems.

### 3.1 | PDFF Reproducibility

Overall, excellent agreement was observed for PDFF measurements across centers and vendors (ICC 0.987 [95% CI: 0.975–0.995], RDC 3.7%, Table 3). Linear regression analysis demonstrated a strong linear association between measurements ( $R^2$  range: 0.95–1.00, Figure 2).

Some variability was observed at low  $T_1$  values (RDC 0.7%–3.5% for  $T_1 \leq 600 \text{ ms}$ ), whereas higher variability occurred at longer  $T_1$  (RDC up to 7.9% for  $T_1 = 1400 \text{ ms}$  and PDFF 30%) (Table 3, Figure 2, and Figure S2).

### 3.2 | $T_1$ Reproducibility

In fat-free vials (PDFF 0%), excellent agreement was observed for MOLLI-based  $T_1$  measurements across centers and vendors (ICC 0.996 [95% CI 0.984–1.0]). Across all vials, including those with fat, overall agreement was good (ICC 0.885 [95% CI 0.793–0.951], RDC 499 ms, Figure 3). Linear regression analysis showed moderate linear association between measurements ( $R^2$  range: 0.76–1; Figure 3).

Variability increased with higher PDFF and longer  $T_1$ , with RDC values ranging from 16 ms (Vial 1, 0% PDFF) to 1553 ms (Vial 16, 30% PDFF) (Table 3, Figure 3).

### 3.3 | Phantom Temporal Stability

Upon return to Center 1, inspection of the phantom housing and vials revealed no visible signs of damage. Excellent agreement was observed between initial and follow-up acquisition measurements at Center 1 for PDFF, with an overall ICC of 0.999 (1.1%) at 1.5T, and ICC of 0.999 (RDC 1.28%) at 3.0T (Figure 4).

Similarly, excellent agreement was observed between initial and follow-up acquisition measurements for  $T_1$  values at Center 1 (Figure 5), with an overall ICC of 0.999 (RDC 52 ms) at 3.0T, and an ICC of 0.998 (RDC 57 ms) at 1.5T. While a slight trend of increasing  $T_1$  values over time was observed at 3.0T (e.g., a maximal increase of 80 ms [5.8%] in vial 4 over 9 months), this trend was not apparent on the 1.5T data.

The surface temperature of the phantom, as indicated by the temperature-indicating strip before each imaging session, ranged between  $19^\circ\text{C}$ – $20^\circ\text{C}$  at 3.0T and  $18^\circ\text{C}$ – $20^\circ\text{C}$  at 1.5T.

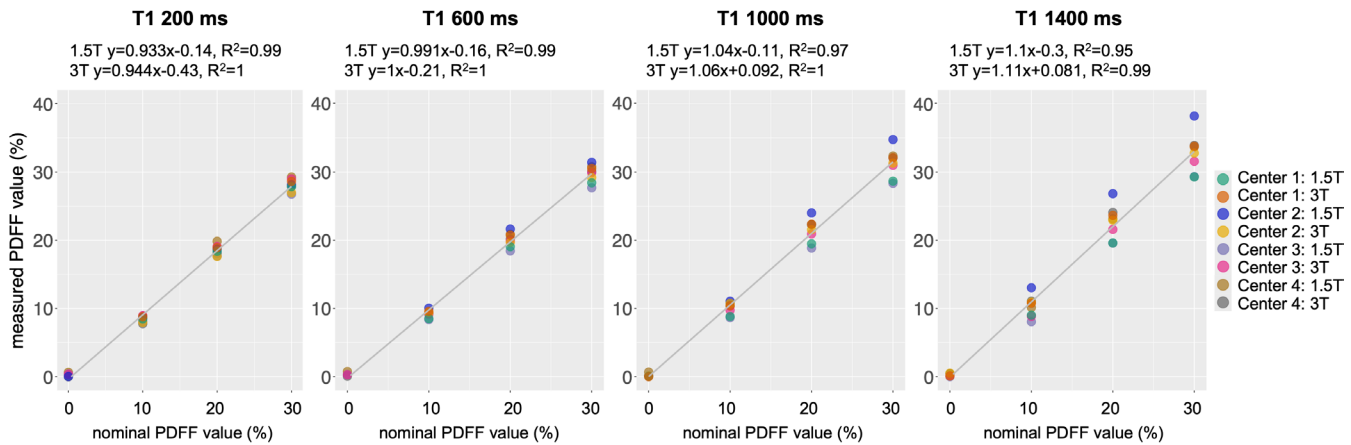
### 3.4 | Validation of Phantom Homogeneity

The SR-CSE validation confirmed high spatial homogeneity across all phantom vials. The CV for  $T_1$  measurements across spatially separated slices ranged from 0.02% to 2.43%, with an ICC of 1.0 [95% CI: 0.999–1.0], indicating negligible through-plane variation (detailed results provided in Table S2).

**TABLE 3** | Reproducibility coefficient for chemical shift-encoded proton density fat fraction (PDFFF) and low flip angle spoiled-gradient-echo-based MOLLI T<sub>1</sub> mapping.

| PDFFF Overall RDC: 3.7% |           |                       |      |         | T <sub>1</sub> Overall RDC: 499 ms |           |                       |       |          |
|-------------------------|-----------|-----------------------|------|---------|------------------------------------|-----------|-----------------------|-------|----------|
| Vial                    | PDFFF (%) | T <sub>1</sub> w (ms) | SD   | RDC (%) | Vial                               | PDFFF (%) | T <sub>1</sub> w (ms) | SD    | RDC (ms) |
| 1                       | 0         | 200                   | 0.25 | 0.71    | 1                                  | 0         | 200                   | 5.6   | 15.6     |
| 2                       | 0         | 600                   | 0.28 | 0.77    | 2                                  | 0         | 600                   | 13.1  | 36.3     |
| 3                       | 0         | 1000                  | 0.23 | 0.63    | 3                                  | 0         | 1000                  | 28.5  | 79.1     |
| 4                       | 0         | 1400                  | 0.23 | 0.63    | 4                                  | 0         | 1400                  | 58.2  | 161.4    |
| 5                       | 10        | 200                   | 0.45 | 1.25    | 5                                  | 10        | 200                   | 9.0   | 24.9     |
| 6                       | 10        | 600                   | 0.52 | 1.44    | 6                                  | 10        | 600                   | 22.9  | 63.6     |
| 7                       | 10        | 1000                  | 0.88 | 2.43    | 7                                  | 10        | 1000                  | 47.0  | 130.4    |
| 8                       | 10        | 1400                  | 1.57 | 4.34    | 8                                  | 10        | 1400                  | 86.3  | 239.1    |
| 9                       | 20        | 200                   | 0.76 | 2.09    | 9                                  | 20        | 200                   | 7.8   | 21.7     |
| 10                      | 20        | 600                   | 1.02 | 2.84    | 10                                 | 20        | 600                   | 44.2  | 122.4    |
| 11                      | 20        | 1000                  | 1.69 | 4.68    | 11                                 | 20        | 1000                  | 131.8 | 365.4    |
| 12                      | 20        | 1400                  | 2.41 | 6.68    | 12                                 | 20        | 1400                  | 298.3 | 826.7    |
| 13                      | 30        | 200                   | 0.89 | 2.48    | 13                                 | 30        | 200                   | 8.7   | 24.2     |
| 14                      | 30        | 600                   | 1.25 | 3.46    | 14                                 | 30        | 600                   | 90.4  | 250.6    |
| 15                      | 30        | 1000                  | 2.09 | 5.78    | 15                                 | 30        | 1000                  | 269.9 | 748.1    |
| 16                      | 30        | 1400                  | 2.86 | 7.93    | 16                                 | 30        | 1400                  | 560.1 | 1552.5   |

Abbreviation: RDC, Reproducibility coefficient.



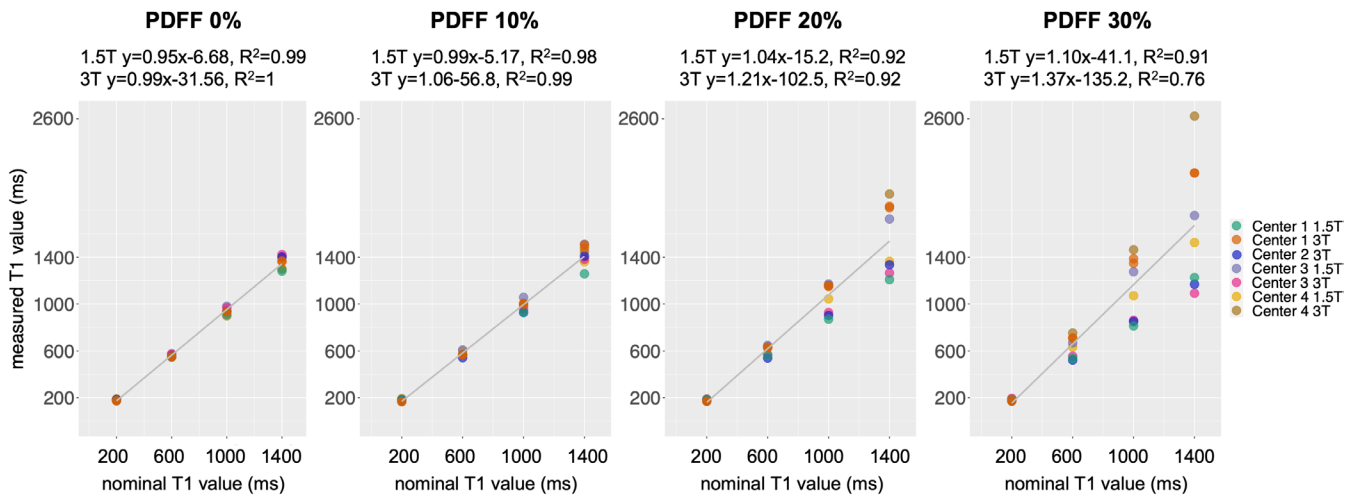
**FIGURE 2** | Linear regression shows good agreement in PDFFF values across centers and platforms, with higher variability in the presence of high fat and T<sub>1</sub>. Depicted are reference (nominal) and measured CSE-MRI PDFFF values across 4 centers and 8 different systems. The columns represent vials with nominal water-specific T<sub>1</sub> of 200 ms, 660 ms, 1000 ms and 1400 ms, respectively. The regression equations predict measured PDFFF (y-axis) based on nominal PDFFF (x-axis). Vendors: Center 1, GE HealthCare; Center 2, Philips Healthcare; Center 3, Siemens Healthineers; Center 4, Canon Medical Systems. Abbreviations: PDFFF, proton density fat fraction; CSE-MRI, chemical shift-encoded MRI.

#### 4 | Discussion

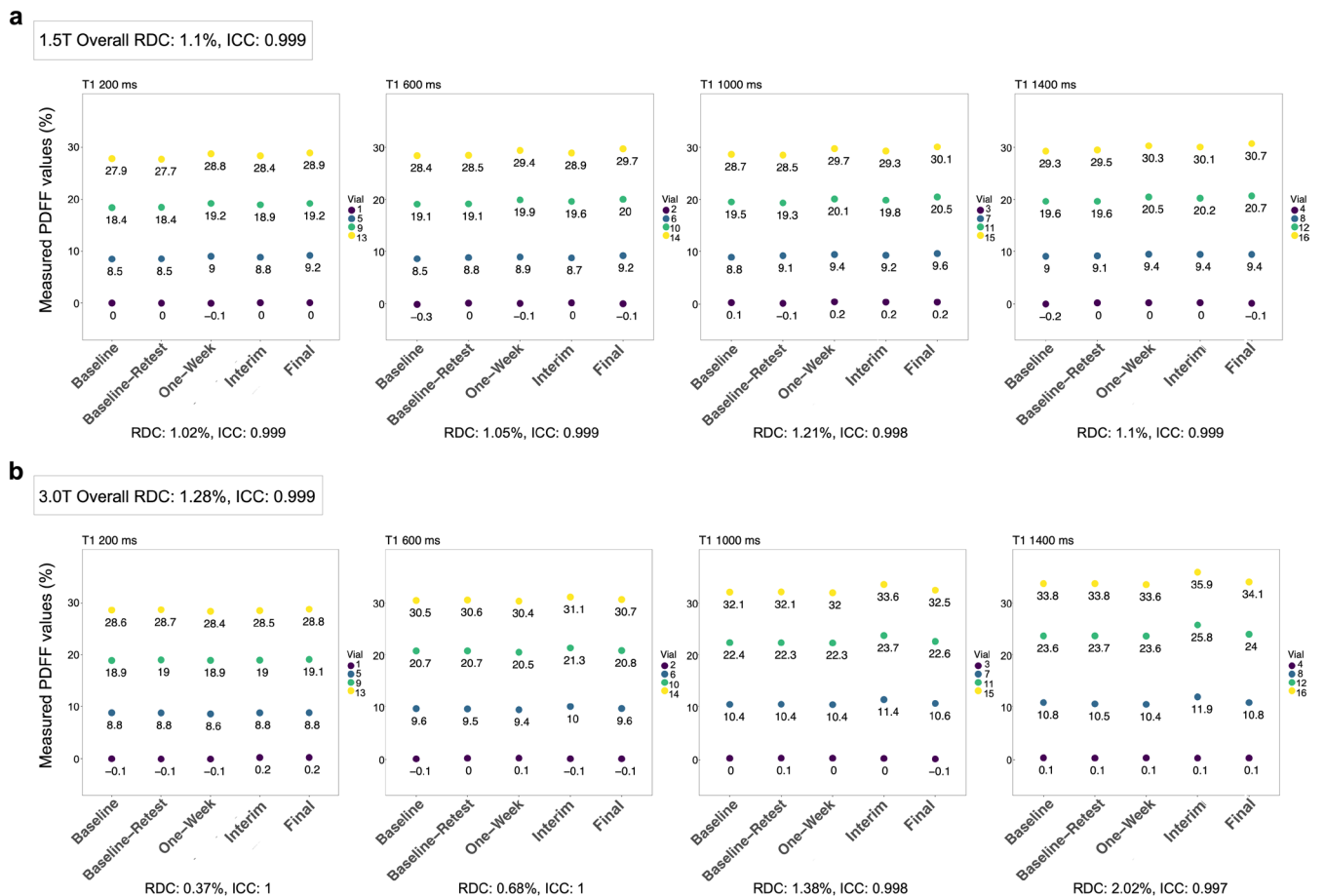
In this work, we conducted a phantom study across four centers with four vendors and multiple platforms at both 1.5T and 3T, using CSE-MRI and MOLLI-based protocols to evaluate the reproducibility of PDFFF and T<sub>1</sub> quantification, respectively. Since

abnormal fat accumulation and fibroinflammatory changes often coexist in liver diseases such as MASLD, joint evaluation of both biomarkers has high clinical relevance.

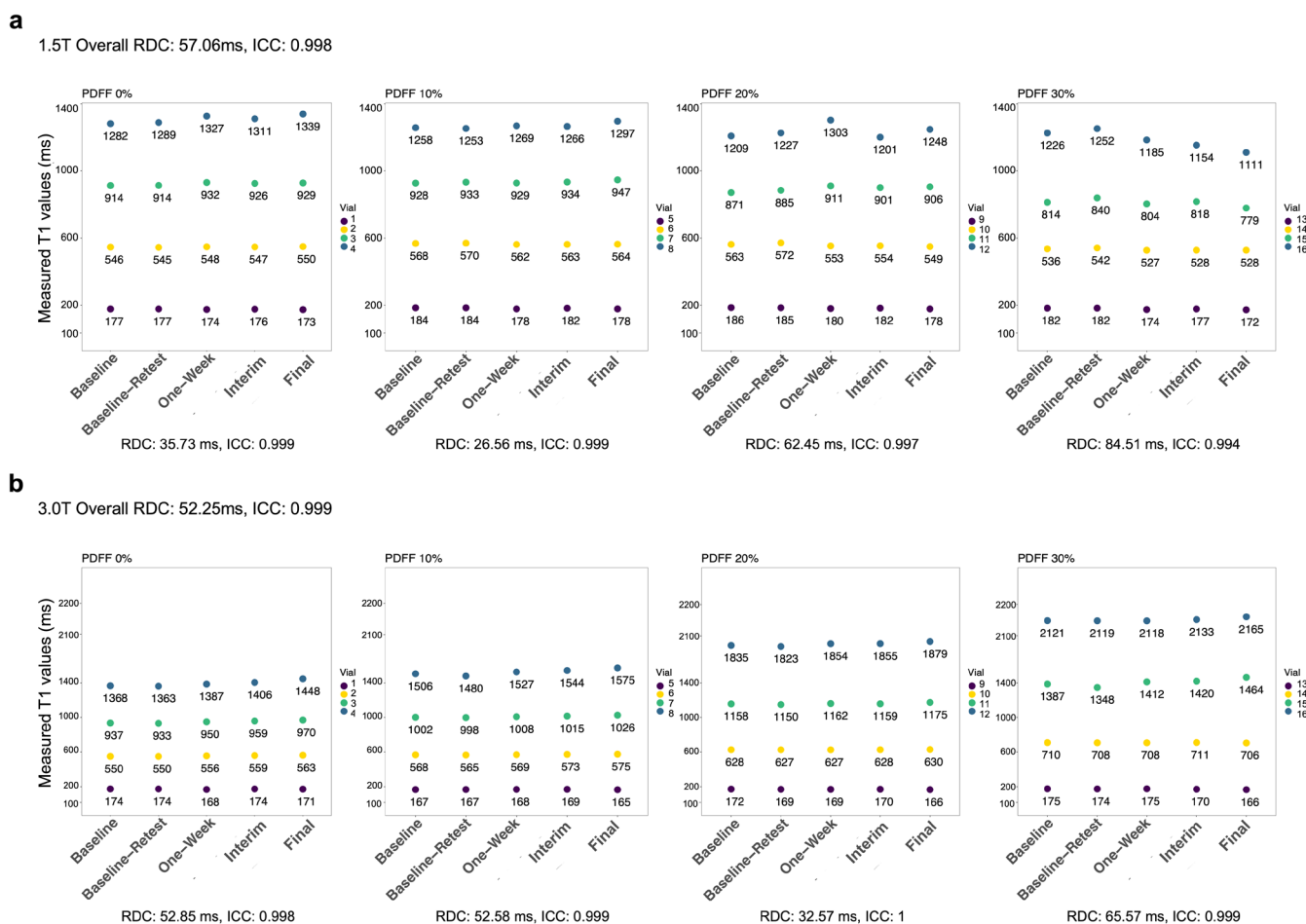
Using a commercially available PDFFF-T<sub>1</sub> phantom with concomitantly modulated PDFFF and T<sub>1</sub> values, we demonstrated



**FIGURE 3** | Linear regression shows observed agreement in  $T_1$  values across the centers and platforms, with low variability in the absence of fat, and higher variability in the presence of high fat and  $T_1$ . Depicted are nominal water-specific  $T_1$  values and measured  $T_1$  values from low flip angle SGRE  $T_1$  mapping across 4 centers and 7 different scanners. Unlike in vivo, where  $T_1$  values vary depending on the scanner's field strength,  $T_1$  values in the phantom are nearly the same at 1.5T and 3T. The regression equations predict measured  $T_1$  (y-axis) based on nominal  $T_1$  (x-axis). Vendors: Center 1, GE HealthCare; Center 2, Philips Healthcare; Center 3, Siemens Healthineers; Center 4, Canon Medical Systems. Abbreviation: SGRE, spoiled-gradient-echo.



**FIGURE 4** | The phantom was imaged at Center 1 several times to test its stability, before and after it was returned to Center 1. Excellent agreement was observed for PDFF between these acquisitions with an overall ICC = 0.999 at 3.0T and 1.5T. Two exams were performed during the initial session (same day): A baseline exam and a retest after repositioning the phantom and reconnecting of the coil. Follow-up exams were conducted at 1 week, 6 months (interim exam), and 9 months later (final exam). Abbreviation: RDC, reproducibility coefficient.



**FIGURE 5** | The phantom was imaged at Center 1 several times to test its stability, before and after it was returned to Center 1. Excellent agreement was observed (ICC=0.999 at 3.0T, and ICC=0.998 at 1.5T). A slight upward drift in  $T_1$  values was noted at 3.0T but not at 1.5T. Two exams were performed during the initial session (same day): A baseline exam and a retest after repositioning the phantom and reconnecting of the coil. Follow-up exams were conducted at 1 week, 6 months (interim exam) and 9 months later (final exam). Abbreviation: RDC, reproducibility coefficient.

overall high multicenter, multi-vendor reproducibility of CSE-MRI based fat quantification at both 1.5T and 3T. Agreement was excellent across most tested PDFF- $T_1$  combinations; however, reproducibility deteriorated at higher  $T_1$  values and higher PDFF levels, with increased variability and systematic overestimation of PDFF. These findings indicate that while PDFF measurements are robust across a broad physiological range, measurement precision is reduced under conditions of prolonged  $T_1$  and high fat content, which are clinically relevant in advanced liver disease.

This behavior is consistent with the known limitations of conventional PDFF mapping methods, which may exhibit residual  $T_1$ -bias depending on the  $T_1$ -weighting parameters (TR, flip angle) and the differences between the  $T_1$  value of water and the  $T_1$  value of fat [11, 27]. Such bias becomes increasingly relevant in tissues with prolonged water  $T_1$ , including those affected by fibroinflammatory changes, and may therefore impact PDFF quantification in progressive stages of MASLD [27]. Further mitigation of  $T_1$  bias is likely achievable through post-processing approaches (given the known acquired flip angles and assumed or measured  $T_1$  values), or by employing nonsteady state  $T_1$ -independent acquisitions [28–30].

Our findings highlight the need for the development of confounder-corrected techniques to ensure accurate and reliable  $T_1$  quantification across sites and patient populations. While MOLLI-based  $T_1$  mapping is reproducible at low PDFF, it is confounded at higher PDFF values due to fat-related signal interference, resulting in substantial bias and variability in  $T_1$  measurements [13]. Standardization strategies, such as system calibration and consistent acquisition protocol, may partially mitigate these limitations. Alternatively, several confounder-corrected liver  $T_1$  mapping techniques have been proposed, correcting for the presence of fat [9, 10, 13, 25, 31, 32]. Once such advanced  $T_1$  mapping methods are implemented across all major vendors, multicenter phantom studies will be needed to validate and benchmark performance. Confounder-corrected methods may enable increased reproducibility of  $T_1$  mapping of the liver, even in the presence of elevated PDFF.

Excellent longitudinal stability of  $T_1$  values was observed at 1.5T. A slight upward trend in  $T_1$  values was noted at 3.0T. This upward trend at 3.0T may be due to confounding factors that impact MOLLI-based  $T_1$  quantification, including  $B_0$  drift, shimming, or RF instability. Drift caused by temperature fluctuation

appears less likely given the lack of monotonic changes in temperature across longitudinal scans.

## 5 | Limitations

Although the phantom was temperature-stabilized prior to scanning at each site, maintaining uniform temperature across sites and systems in the MRI suite was not possible and may have introduced variability between sites. Temperature fluctuations during imaging due to system heating may partially explain discrepancies between the estimated and reference PDFF and  $T_1$  values. However, our acquisition protocol represents a realistic scenario for real-world deployment of phantom-based quality assurance. Therefore, this slight variability is likely representative of the subsequent use of phantom-based quality assurance in the clinic or in clinical trials. Furthermore, any temperature dependence may vary across vendors for PDFF due to differences in reconstruction algorithms. In contrast,  $T_1$  maps were reconstructed offline using a single method (as some systems lacked online reconstructions for the acquisition parameters used in this project), minimizing such variability [16].

Another aspect to consider is the use of a head or head-neck coil rather than a body array coil, which would typically be used in clinical abdominal imaging. While this choice was made to facilitate consistent and reproducible positioning across centers, it may affect image quality parameters such as signal-to-noise ratio and coverage. While B1 inhomogeneity is a known confounder for MOLLI, its impact was minimized by the small size of the phantom and position at isocenter. While important, this effect was considered secondary to the more pronounced confounding effects of fat observed in this study.

Our study was based exclusively on phantoms rather than ex vivo or in vivo tissue. While this limits the direct clinical applicability of our findings, it provides greater control over imaging conditions. Conducting a study involving patient or specimen imaging across multiple centers would be logistically challenging [33]. Finally, although we included four major MRI vendors to enhance generalizability, expanding to additional platforms and clinical environments would further strengthen the applicability of our results.

## 6 | Conclusion

This multicenter, multi-vendor, multi-platform study demonstrated high reproducibility of CSE-based PDFF measurements, using a phantom with simultaneously modulated PDFF and  $T_1$  values. Although PDFF measurements were generally robust, reproducibility deteriorated under conditions of prolonged  $T_1$  and high fat content, which are common in advanced liver disease. In contrast, MOLLI-based  $T_1$  mapping showed markedly reduced reproducibility in the presence of increasing fat content, with substantial variability at higher PDFF levels. These findings support the established high reproducibility of PDFF as a quantitative imaging biomarker and emphasize the importance of further development and validation of  $T_1$  mapping techniques, particularly for use in patients with coexisting steatosis and fibroinflammatory changes. Our results support the development

and standardization of liver MRI biomarkers across clinical trials and clinical practice.

## Funding

J.H.B. and D.H. report grant support from: NIH R41 EB025729, NIH R44 EB025729. S.B.R. and D.H. report grant support from: NIH R01 EB031886. A.J.M. reports grant support from: NIH R01 CA283663. S.W. acknowledges funding from: European Research Council ERC Grant no. 101078711; Nederlandse Hartstichting, Grant no. 03-004-2022-0079.

## Disclosure

S.B.R. has ownership interests in Calimetrix, Madison, WI, USA. Unrelated to this work, he reports ownership interests in Reveal Pharmaceuticals, VistaAI, RevOps, Elucent Medical, and he provides consulting services for Amgen. He receives research support from the John H. Juhl Professorship. The University of Wisconsin-Madison received research support from GE HealthCare and Bracco Diagnostics.

## References

1. Z. M. Younossi, M. Kalligeros, and L. Henry, "Epidemiology of Metabolic Dysfunction-Associated Steatotic Liver Disease," *Clinical and Molecular Hepatology* 31 (2024): S32–S50.
2. B. J. Perumpail, M. A. Khan, E. R. Yoo, G. Cholankeril, D. Kim, and A. Ahmed, "Clinical Epidemiology and Disease Burden of Nonalcoholic Fatty Liver Disease," *World Journal of Gastroenterology* 23 (2017): 8263–8276.
3. J. Starekova, D. Hernando, P. J. Pickhardt, and S. B. Reeder, "Quantification of Liver Fat Content With CT and MRI: State of the Art," *Radiology* 301 (2021): 250–262.
4. S. B. Reeder and J. Starekova, "MRI Proton Density Fat Fraction for Liver Disease Risk Assessment: A Call for Clinical Implementation," *Radiology* 309 (2023): e232552, <https://doi.org/10.1148/radiol.232552>.
5. J. Patel, R. Bettencourt, J. Cui, et al., "Association of Noninvasive Quantitative Decline in Liver Fat Content on MRI With Histologic Response in Nonalcoholic Steatohepatitis," *Therapeutic Advances in Gastroenterology* 9 (2016): 692–701.
6. T. Yokoo, S. D. Serai, A. Pirasteh, et al., "Linearity, Bias, and Precision of Hepatic Proton Density Fat Fraction Measurements by Using MR Imaging: A Meta-Analysis," *Radiology* 286 (2018): 486–498.
7. F. V. De Oliveira Terzi, D. B. Parente, G. C. Camargo, et al., "MRI-Derived Extracellular Volume to Assess Liver Fibrosis in Patients With Metabolic-Associated Steatotic Liver Disease," *Abdom Radiol* 64, no. 2 (2026): 493–501.
8. M. Haimerl, N. Verloh, F. Zeman, et al., "Gd-EOB-DTPA-Enhanced MRI for Evaluation of Liver Function: Comparison Between Signal-Intensity-Based Indices and  $T_1$  Relaxometry," *Scientific Reports* 7 (2017): 43347.
9. R. B. Thompson, K. Chow, D. Mager, J. J. Pagano, and J. Grenier, "Simultaneous Proton Density Fat-Fraction and Imaging With Water-Specific  $T_1$  Mapping (PROFIT<sub>1</sub>): Application in Liver," *Magnetic Resonance in Medicine* 85 (2021): 223–238.
10. Y. Muslu, D. Tamada, N. T. Roberts, et al., "Free-Breathing, Fat-Corrected  $T_1$  Mapping of the Liver With Stack-Of-Stars MRI, and Joint Estimation of  $T_1$ , PDFF,  $R_2^*$ , and B1+," *Magnetic Resonance in Medicine* 92 (2024): 1913–1932.
11. C.-Y. Liu, C. A. McKenzie, H. Yu, J. H. Brittain, and S. B. Reeder, "Fat Quantification With IDEAL Gradient Echo Imaging: Correction of Bias from  $T_1$  and Noise," *Magnetic Resonance in Medicine* 58 (2007): 354–364.
12. X. Wang, T. J. Colgan, L. A. Hinshaw, et al., " $T_1$ -Corrected Quantitative Chemical Shift-Encoded MRI," *Magnetic Resonance in Medicine* 83 (2020): 2051–2063.

13. R. B. Thompson, R. Sherrington, C. Beaulieu, et al., "Reference Values for Water-Specific  $T_1$  of the Liver at 3T:  $T_2^*$ -Compensation and the Confounding Effects of Fat," *Journal of Magnetic Resonance Imaging* 60 (2024): 2063–2075.
14. L. A. Gilligan, J. R. Dillman, J. A. Tkach, S. A. Xanthakos, J. K. Gill, and A. T. Trout, "Magnetic Resonance Imaging  $T_1$  Relaxation Times for the Liver, Pancreas and Spleen in Healthy Children at 1.5 and 3T," *Pediatric Radiology* 49 (2019): 1018–1024.
15. D. Hernando, S. D. Sharma, M. Aliyari Ghasabeh, et al., "Multisite, Multivendor Validation of the Accuracy and Reproducibility of Proton-Density Fat-Fraction Quantification at 1.5T and 3T Using a Fat-Water Phantom: Proton-Density Fat-Fraction Quantification at 1.5T and 3T," *Magnetic Resonance in Medicine* 77 (2017): 1516–1524.
16. H. H. Hu, T. Yokoo, M. R. Bashir, et al., "Linearity and Bias of Proton Density Fat Fraction as a Quantitative Imaging Biomarker: A Multicenter, Multiplatform, Multivendor Phantom Study," *Radiology* 298 (2021): 640–651.
17. J. K. Jang, S. S. Lee, B. Kim, et al., "Agreement and Reproducibility of Proton Density Fat Fraction Measurements Using Commercial MR Sequences Across Different Platforms: A Multivendor, Multi-Institutional Phantom Experiment," *Investigative Radiology* 54 (2019): 517–523.
18. J. Starekova, D. Rutkowski, W. C. Bae, et al., "Multi-Center, Multi-Vendor Validation of Simultaneous MRI -Based Proton Density Fat Fraction and  $R_2^*$  Mapping Using a Combined Proton Density Fat Fraction-  $R_2^*$  Phantom," *Journal of Magnetic Resonance Imaging* 62 (2025): 800–811.
19. S. Roujol, S. Weingärtner, M. Foppa, et al., "Accuracy, Precision, and Reproducibility of Four  $T_1$  Mapping Sequences: A Head-To-Head Comparison of MOLLI, ShMOLLI, SASHA, and SAPHIRE," *Radiology* 272 (2014): 683–689.
20. G. Captur, A. Bhandari, R. Brühl, et al., " $T_1$  Mapping Performance and Measurement Repeatability: Results From the Multi-National  $T_1$  Mapping Standardization Phantom Program (TIMES)," *Journal of Cardiovascular Magnetic Resonance* 22 (2020): 31.
21. D. Hernando, S. D. Sharma, H. Kramer, and S. B. Reeder, "On the Confounding Effect of Temperature on Chemical Shift-Encoded Fat Quantification: Effect of Temperature on CSE Fat Quantification," *Magnetic Resonance in Medicine* 72 (2014): 464–470.
22. C.-Y. Liu, C. Noda, B. Ambale-Venkatesh, Y. Kassai, D. Bluemke, and J. A. C. Lima, "Evaluation of Liver  $T_1$  Using MOLLI Gradient Echo Readout Under the Influence of Fat," *Magnetic Resonance Imaging* 85 (2022): 57–63.
23. D. R. Messroghli, A. Radjenovic, S. Kozierke, D. M. Higgins, M. U. Sivananthan, and J. P. Ridgway, "Modified Look-Locker Inversion Recovery (MOLLI) for High-Resolution  $T_1$  Mapping of the Heart," *Magnetic Resonance in Medicine* 52 (2004): 141–146.
24. S. K. Piechnik, V. M. Ferreira, E. Dall'Armellina, et al., "Shortened Modified Look-Locker Inversion Recovery (ShMOLLI) for Clinical Myocardial  $T_1$ -Mapping at 1.5 and 3T Within a 9 Heartbeat Breathhold," *Journal of Cardiovascular Magnetic Resonance* 12 (2010): 69.
25. G. C. Fullerton, D. Tamada, Y. Muslu, D. Hernando, and S. B. Reeder, "Saturation Recovery Chemical Shift-Encoded  $T_1$  Mapping in the Liver. Oral Scientific Session," *International Society of Magnetic Resonance in Medicine (ISMRM) Annual Meeting* (2025). Abstract number 0816.
26. T. K. Koo and M. Y. Li, "A Guideline of Selecting and Reporting Intraclass Correlation Coefficients for Reliability Research," *Journal of Chiropractic Medicine* 15 (2016): 155–163.
27. J. Kühn, C. Jahn, D. Hernando, et al., " $T_1$  Bias in Chemical Shift-Encoded Liver Fat-Fraction: Role of the Flip Angle," *Journal of Magnetic Resonance Imaging* 40 (2014): 875–883.
28. J. Tang, D. Tamada, R. do Vale Souza, et al., "Optimized Motion-Insensitive PDFF Mapping of the Liver," *Magnetic Resonance in Medicine* 95 (2026): 249–267.
29. C.-Y. Liu, C. A. McKenzie, H. Yu, J. H. Brittain, and S. B. Reeder, "Fat Quantification With IDEAL Gradient Echo Imaging: Correction of Bias From  $T_1$  and Noise," *Magnetic Resonance in Medicine* 58 (2007): 354–364.
30. C. D. G. Hines, A. Frydrychowicz, G. Hamilton, et al., " $T_1$  Independent,  $T_2^*$  Corrected Chemical Shift-Based Fat-Water Separation With Multi-Peak Fat Spectral Modeling Is an Accurate and Precise Measure of Hepatic Steatosis," *Journal of Magnetic Resonance Imaging* 33 (2011): 873–881.
31. N. Wang, T. Cao, F. Han, et al., "Free-Breathing Multitasking Multi-Echo MRI for Whole-Liver Water-Specific  $T_1$ , Proton Density Fat Fraction, and Quantification," *Magnetic Resonance in Medicine* 87 (2022): 120–137.
32. N. T. Roberts, D. Tamada, Y. Muslu, D. Hernando, and S. B. Reeder, "Confounder-Corrected  $T_1$  Mapping in the Liver Through Simultaneous Estimation of  $T_1$ , PDFF,  $R_2^*$ , and  $B_1+$  in a Single Breath-Hold Acquisition," *Magnetic Resonance in Medicine* 89 (2023): 2186–2203.
33. T. He, P. Kirk, D. N. Firmin, et al., "Multi-Center Transferability of a Breath-Hold  $T_2$  Technique for Myocardial Iron Assessment," *Journal of Cardiovascular Magnetic Resonance* 10 (2008): 11.

### Supporting Information

Additional supporting information can be found online in the Supporting Information section. **Figure S1:** Phantom shipping workflow and imaging schedule across centers imaging at Center 1 was performed at multiple points throughout the study to evaluate phantom integrity and monitor potential drift in proton density fat fraction (PDFF) and  $T_1$  values. Two exams were conducted during the initial session on the same day (a baseline exam and followed by a retest after repositioning the phantom and reconnecting the coil). Follow-up exams took place at 1 week, 6 months (interim exam; after the phantom was shipped from Center 3 to Center 1) and 9 months (final exam). The phantom was transported between participating centers using an overnight courier service within a protective, foam-padded case. **Figure S2:** Good reproducibility with low bias was observed for PDFF mapping across PDFF values (0%–30%) and  $T_1$  values (200–1400 ms). A moderate overestimation and increased variability were observed in measurements with long  $T_1$ , particularly at higher PDFF values. This effect is likely due to residual  $T_1$  weighting in conventional PDFF mapping methods. These methods use a low flip angle between 3°C–5°C, which still leads to residual  $T_1$  bias when the  $T_1$  of water is much longer than the  $T_1$  of fat (approximately 350 ms). Note: PDFF, proton density fat fraction. **Table S1:** Proton density fat fraction (PDFF) and  $T_1$  reproducibility coefficient (RDC) and percentage RDC (% RDC). **Table S2:** Validation of phantom homogeneity via saturation-recovery chemical shift-encoded (SR-CSE)  $T_1$  mapping.