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T_1 mapping performance and measurement repeatability: results from the multi-national T_1 mapping standardization phantom program (T1MES)

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Abstract

Background: The T_1 Mapping and Extracellular volume (ECV) Standardization (T1MES) program explored T_1 mapping quality assurance using a purpose-developed phantom with Food and Drug Administration (FDA) and Conformité Européenne (CE) regulatory clearance. We report T_1 measurement repeatability across centers describing sequence, magnet, and vendor performance.

Methods: Phantoms batch-manufactured in August 2015 underwent 2 years of structural imaging, B_0 and B_1 , and “reference” slow T_1 testing. Temperature dependency was evaluated by the United States National Institute of Standards and Technology and by the German Physikalisch-Technische Bundesanstalt. Center-specific T_1 mapping repeatability (maximum one scan per week to minimum one per quarter year) was assessed over mean 358 (maximum 1161) days on 34 1.5 T and 22 3 T magnets using multiple T_1 mapping sequences. Image and temperature data were analyzed semi-automatically. Repeatability of serial T_1 was evaluated in terms of coefficient of variation (CoV), and linear mixed models were constructed to study the interplay of some of the known sources of T_1 variation.

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Results: Over 2 years, phantom gel integrity remained intact (no rips/tears), B_0 and B_1 homogenous, and “reference” T_1 stable compared to baseline (% change at 1.5 T, $1.95 \pm 1.39\%$; 3 T, $2.22 \pm 1.44\%$). Per degrees Celsius, 1.5 T, T_1 (MOLLI 5s(3s)3s) increased by 11.4 ms in long native blood tubes and decreased by 1.2 ms in short post-contrast myocardium tubes. Agreement of estimated T_1 times with “reference” T_1 was similar across Siemens and Philips CMR systems at both field strengths (adjusted R^2 ranges for both field strengths, 0.99–1.00). Over 1 year, many 1.5 T and 3 T sequences/magnets were repeatable with mean CoVs < 1 and 2% respectively. Repeatability was narrower for 1.5 T over 3 T. Within T1MES repeatability for native T_1 was narrow for several sequences, for example, at 1.5 T, Siemens MOLLI 5s(3s)3s prototype number 448B (mean CoV = 0.27%) and Philips modified Look-Locker inversion recovery (MOLLI) 3s(3s)5s (CoV 0.54%), and at 3 T, Philips MOLLI 3b(3s)5b (CoV 0.33%) and Siemens shortened MOLLI (ShMOLLI) prototype 780C (CoV 0.69%). After adjusting for temperature and field strength, it was found that the T_1 mapping sequence and scanner software version (both $P < 0.001$ at 1.5 T and 3 T), and to a lesser extent the scanner model ($P = 0.011$, 1.5 T only), had the greatest influence on T_1 across multiple centers.

Conclusion: The T1MES CE/FDA approved phantom is a robust quality assurance device. In a multi-center setting, T_1 mapping had performance differences between field strengths, sequences, scanner software versions, and manufacturers. However, several specific combinations of field strength, sequence, and scanner are highly repeatable, and thus, have potential to provide standardized assessment of T_1 times for clinical use, although temperature correction is required for native T_1 tubes at least.

Keywords: T_1 mapping, Standardization, Calibration, Phantom, Repeatability, Extracellular volume

Introduction

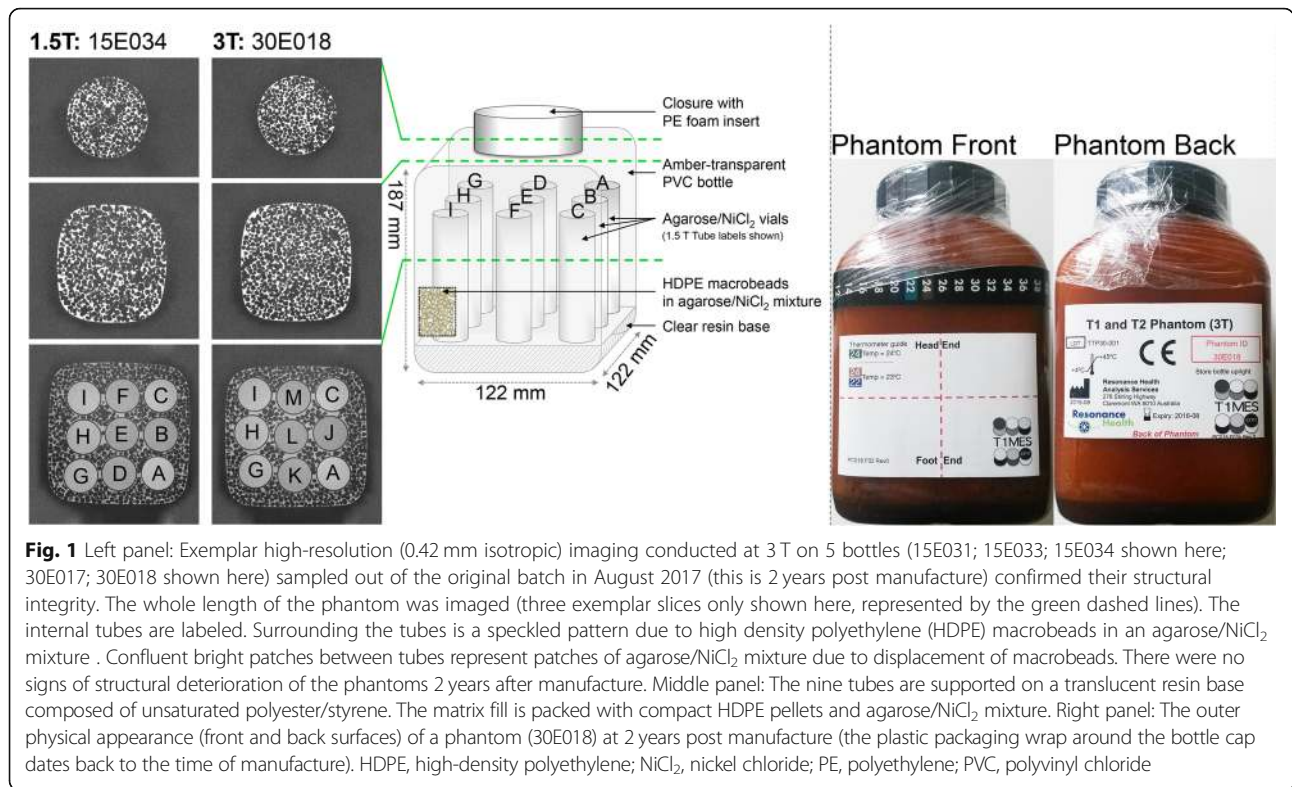
T_1 mapping aids clinicians in the assessment and diagnosis of myocardial disease. However, measurement needs to be stable over time with transferable values. Knowledge of normal reference ranges would benefit from not requiring local healthy subject scanning, and the pooling of multi-scanner datasets would have advantages such as increasing available sample sizes for the detection of subtle effects or subgroup analysis and increasing result robustness and generalizability, lowering the chance of unforeseen bias when compared to single-center data [1]. Combining results however introduces sequence, magnet, and field strength bias [2]. The field of T_1 mapping would therefore benefit from a “ T_1 standard” to enable cross-center T_1 mapping data pooling and delivery [3]—like the international normalized ratio (INR) which makes it possible to adjust the dosing of vitamin K antagonists regardless of which laboratory has performed the test [4].

The T_1 Mapping and Extracellular volume (ECV) Standardization (T1MES) phantom program was established to explore T_1 mapping quality assurance at 1.5 T and 3 T and understand the feasibility of delivering a “ T_1 standard” [5]. The first step toward the goal was development and mass-production of a phantom [5] and its European Union Conformité Européenne (CE) and United States Food and Drug Administration (FDA) regulatory clearance. In September 2015, cardiovascular magnetic resonance (CMR) centers worldwide joined the T1MES consortium and committed to submit a minimum of 12 months of center-specific T_1 mapping data. Data submitted have now been analyzed to explore phantom performance.

We report phantom data at 1 and 2 years using various T_1 mapping sequences, temperature sensitivity, and include platform performance, although we emphasize that comparison of different T_1 methods and systems was not the main aim, rather investigating long-term stability towards the “ T_1 standard”. This involved modeling some of the potential sources of the T_1 variation longitudinally and between T1MES centers to identify the most influential factors.

Methods

The development and description of the T1MES phantom (Fig. 1) has been previously reported [5]. Briefly, the T1MES phantom was designed to be field-strength specific (i.e., separate 1.5 T and 3 T models). Each phantom contains four tubes representing human native blood/myocardial T_1 and T_2 values (i.e., pre-gadolinium-based contrast agent [GBCA] values) and five tubes representing human post-GBCA blood/myocardial values. While the main aim of the present study was the collection and analysis of the multi-center data (see the “[Methods part 2—Multi-center phantom testing](#)” section), some other tests were applied to a small number of the phantoms during the 2 years to explore the utility of T1MES as a quality assurance device, and these tests are described here first (“[Methods part 1—Evaluation of the phantom](#)”). Imaging biomarker terms used follow the recommendations of the Quantitative Imaging Biomarkers Alliance (QIBA) of the Radiological Society of North America (RSNA) [6].



Methods part 1—Evaluation of the phantom

Structural integrity

Gel integrity and aging were checked at each submission time point for participating sites through the manual inspection of localizers that formed part of the minimum dataset requirement for participation. In addition, a high-resolution, isotropic, three-dimensional (3D) gradient echo sequence (0.42 mm³) was run on four phantoms (three 1.5 T phantoms; one 3 T phantom) at baseline (October 2015) and at 2 years post manufacturing in each case using a 3 T MAGNETOM Skyra (Siemens Healthineers, Erlangen, Germany; software syngo MR D13C). The sequence acquired two overlapping slabs (due to scanner software constraints), each with two directions of phase encoding, a slow repetition time (repetition time, TR = 17 ms), and narrow sampling bandwidth (250 Hz/pixel) for better signal-to-noise ratio (SNR). This sequence had weak T₁ and T₂ image contrast and was only for structural examination.

“Reference” rT₁ and rT₂ data

Baseline (October 2015) “reference” T₁ and T₂ values (rT₁, rT₂) were acquired at the Royal Brompton Hospital CMR Unit using basic single-slice TR = 10 s inversion recovery spin echo (IRSE, 8 inversion times [TI] from 25 to 3200 ms) and single-slice repetition time (TR) = 10 s SE (8 echo times [TE] from 10 to 640 ms) [5] respectively. These sequences were identically repeated at 2 years on the same

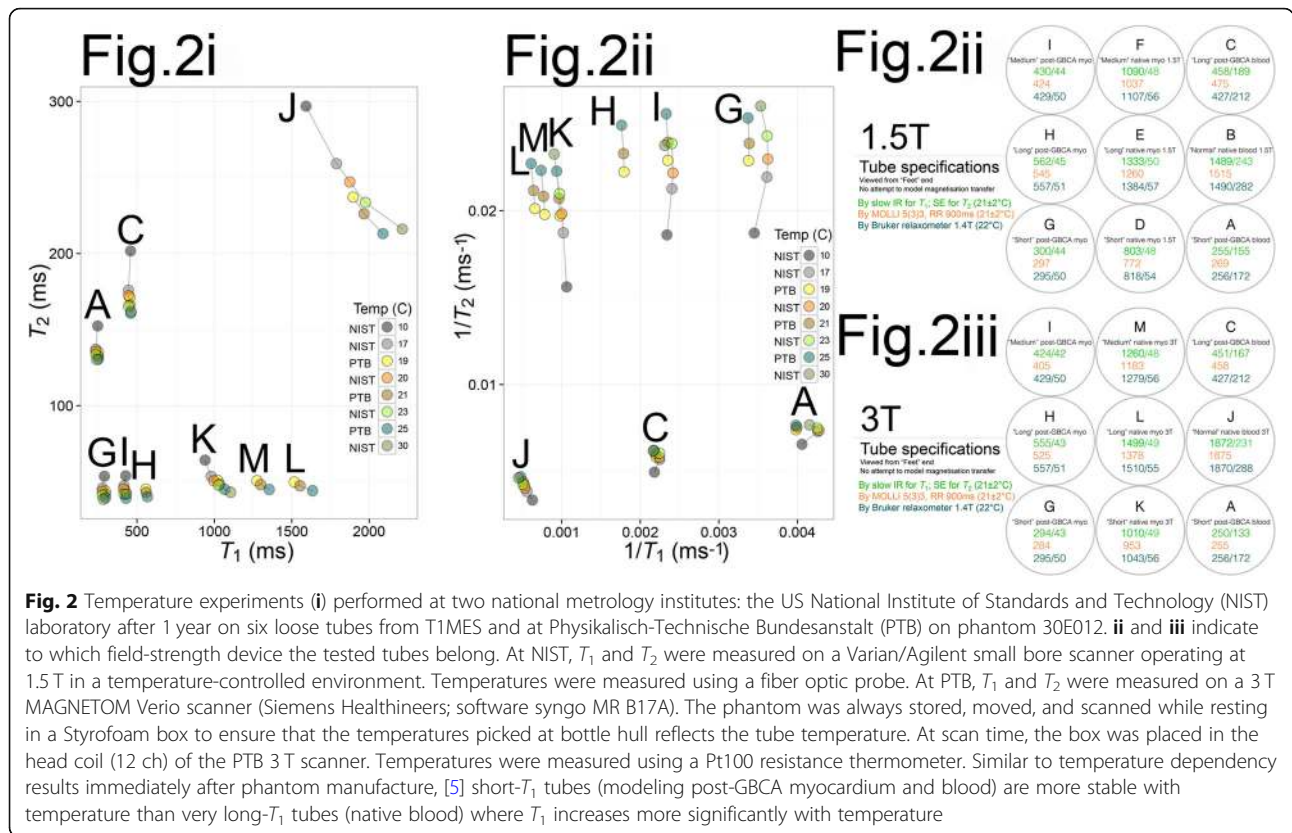
three 1.5 T phantoms and on the same three 3 T phantoms sampled from the production batch. The identifying serial numbers of the three 1.5 T phantoms were 15E031, 15E033, and 15E034, and these phantoms were scanned on a 1.5 T MAGNETOM Avanto [Siemens Healthineers; software syngo MR B17A]. The three 3 T phantoms were 30E001, 30E017, and 30E018, and they were scanned on a 3 T MAGNETOM Skyra [Siemens Healthineers; software syngo MR D13C].

Separate rT₁ and rT₂ data were acquired on the same 3 T phantom (30E021) at the German National Metrology Institute, Physikalisch-Technische Bundesanstalt (PTB) over a period of 1041 days (64 scans) commencing September 2015 (3 T MAGNETOM Verio (Siemens Healthineers; software syngo MR B17A). Sequences used for rT₁ and rT₂ were respectively basic single-slice TR = 8000 ms IRSE (IRSE, 7 TI from 25 to 4800 ms) and single-slice TR = 3000 ms SE (5 TE from 24 to 400 ms).

Temperature sensitivity

The following three methods were used:

First, controlled-temperature experiments over the range 10–30 °C were conducted at the United States National Institute of Standards and Technology (NIST) on six loose T1MES tubes at 1 year (Fig. 2i). T₁ and T₂ were measured at 10, 17, 20, 23, and 30 °C on an VnmrJ4



small-bore scanner operating at 1.5 T (Varian Medical Systems, Palo Alto, California, USA) in a temperature-controlled environment using a fiber optic temperature probe. T_1 was measured by IRSE (TR = 10 s, TI = 50–3000 ms) and T_2 by SE (TR = 10 s, TE = 15–960 ms).

Second, controlled-temperature experiments at 19, 21, and 25 °C were conducted at the PTB laboratory on T1MES phantom 30E012 at 1 year (also Fig. 2i). T_1 and T_2 were measured on a 3 T MAGNETOM Verio scanner (Siemens Healthineers; software syngo MR B17A) using a Pt100 resistance thermometer. T_1 was measured by IRSE (TR = 8000 ms, TI = 25–4800 ms) and T_2 by SE (TR = 3000 ms, TE = 24–400 ms).

Third, for each T1MES phantom scan at all centers, temperature was measured using liquid crystal thermometers adhered to every phantom. These measurements were pooled and analyzed to derive temperature-correction algorithms (see Statistical Analysis).

B_0 and B_1 uniformity

These uniformities and the fundamental distortion of B_1 by water dielectric permittivity especially at 3 T had been tested at baseline (October 2015, previously reported [5]). These uniformities were mapped later to check against “cracking” of the gel and subsequent impact of air gaps on B_0 in particular, while potential “clumping” of the plastic beads over time might in theory affect the B_1 [5]. We

therefore considered it prudent to check whether anything unexpected occurred over the long term.

B_0 uniformity was therefore mapped at 2 years in six phantoms, in the transverse slice, midway along the length of the tubes, using a multi-echo gradient echo sequence, based on the phase difference between known TEs [7]. A frequency range of ± 50 Hz across the phantom was considered acceptable, based on published T_1 mapping off-resonance sensitivity [8]. B_1 homogeneity was similarly evaluated using flip angle (FA) maps (double angle method using FA 60° and 120° [01, 2 × 01] with long TR [8 s], and 4 ms sinc [− 3π to + 3π] slice excitation profiles to minimize error due to FA variation through the slice).

Methods part 2—Multi-center phantom testing

Serial, multi-center T_1 mapping data

The T1MES user manual (https://doi.org/10.6084/m9.fig-share.c.3610175_D1.v1) defined strict scanning instructions (scanning and shim volume strictly at isocenter, use of same supporting materials, etc.). Each contributed T1MES dataset (localizers, sets of inversion recovery images, and inline scanner-generated T_1 maps, Fig. 3) underwent initial quality assurance, checking orientation, and isocenter (through visual inspection of localizers and maps and semi-automatically by inspecting metadata contained in Digital Imaging and Communications in Medicine [DICOM] headers “ImagePositionPatient” and “ImageOrientationPatient”) and to exclude

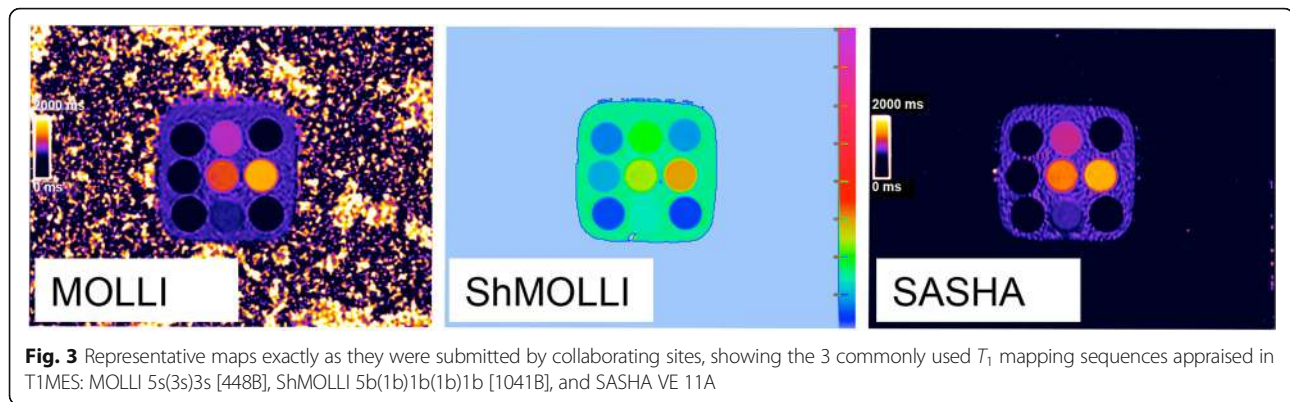


image artifacts. All Siemens sequences except MyoMaps product variant and all Philips (Philips Healthcare, Best, the Netherlands) sequences except CardiacQuant product variant were prototypes. Any tubes with artifacts detected by operator inspection of the submitted T_1 maps were excluded from the analysis. Software version changes were captured automatically from DICOM headers (“StationName” and “SoftwareVersion”).

The T_1 measurements from T1MES datasets (directly using only the parametric maps submitted, not by any T_1 fitting applied centrally to the submitted sets of T_1 recovery images) were carried out using a bespoke MATLAB pipeline (The MathWorks Inc., Natick, Massachusetts, USA, R2012b) assembled in collaboration with the US National Institutes of Health. From the data, T_1 for each of the nine tubes was measured in identically sized regions of interest (ROI) occupying the central 50% by area of each tube (accommodating ~ 40 independent pixels) and collated in a dedicated research electronic data capture instrument (REDCap [9, 10]).

Methods part 3—Statistical analysis

Analysis was performed using R (version 3.0.1, R Foundation for Statistical Computing, Vienna, Austria). Descriptive data are expressed as mean \pm standard deviation (SD) and standard error of the mean (SEM) as appropriate. Distribution of data were assessed on histograms and using the Shapiro-Wilk test.

Temperature sensitivity

Linear regression equations were used to relate temperature (predictor variable in degrees Celsius) and the response variable, phantom T_1 , by the formula: $T_1 = \text{Intercept} + (\beta * [\text{Temperature} - 21^\circ \text{C}])$, with β representing the temperature correction, and 21°C our arbitrarily chosen temperature for cross-center comparison.

Correlation with rT_1 times

Correlations between estimated and rT_1 times were derived using linear regression. Tests for significant inter-sequence and cross-vendor correlation differences

(setting null value to 0.001) were conducted with alpha 0.01 and confidence level 0.95 [11].

T_1 repeatability

After considering the normal values for native myocardial T_1 reported in the published literature (e.g., in [12–16] as mean \pm 1SD, though a 95% reference range is approximately \pm 2SD), where 1 SD of the mean native myocardial T_1 is generally ~ 20 – 30 ms at 1.5 T and ~ 50 ms at 3 T, we arbitrarily pre-defined as repeatable (and suitable for clinical/research use), T_1 mapping approaches where the estimated variance of serial T_1 data did not exceed $\frac{1}{2}$ of the above in vivo 1SD. For T_1 mapping at 1.5 T, this was ≤ 10 ms, i.e., CoV $\leq 1\%$; for T_1 mapping at 3 T ≤ 25 ms, i.e., CoV $\leq 2\%$.

The CoV between serial repeat T1MES scans was calculated as the ratio of the SD to the mean. We appraised CoV as a compound measure of all causes of change in the estimated T_1 of all nine tubes before and after temperature correction. We also appraised CoV after temperature correction separately for the four native and five post-GBCA tubes. Sequence-specific differences between the nine temperature-adjusted CoVs were calculated using paired t test with P value adjustment for multiple comparisons by the Bonferroni method (taking two-tailed $P < 0.01$ as significant).

Sources of T_1 variation

Using temperature-adjusted T_1 values of the “Medium” native myocardium tubes (tubes “F” and “M” respectively), we constructed linear mixed models to study the interplay of some known sources of T_1 variation in multi-center phantom data. We did this separately for 1.5 T and 3 T phantom data. Considering temperature-adjusted T_1 time as the response variable of interest, we examined the influence of phantom ID with and without the added effect of phantom age, as the combined fixed effect. With this, we then tested the following random effects:

- i) Main effects and interactions of scanner vendor/scanner model (Siemens, Philips or General Electric

- [GE; General Electric Healthcare, Waukesha, Wisconsin, USA]; e.g., for Siemens: MAGNETOM Aera vs. Avanto vs. Espree, etc.);
- ii) Main effects and interaction of sequence/scanner software version (considering all submitted variants of native modified Look-Locker inversion recovery [MOLLI] [17] sequences, shortened MOLLI [18] [ShMOLLI], native saturation-recovery single-shot acquisition [19] [SASHA], and saturation method using adaptive recovery times for cardiac T_1 mapping [20] [SMART]; e.g., for Philips: R4.1.3SP2 vs. R5.1.7SP2 vs. R5.2.0SP2, etc.).

The response variable T_1 fitted a normal probability distribution, so we estimated model parameters using maximum likelihood. ANOVA function using a type II Wald chi-square test evaluated the significance of fixed effects in the model. To compare models, Akaike and Bayesian information criteria (AIC, BIC) with the “smaller-is-better” criterion as well as chi-square values from inter-model ANOVA tests were used. The formulas used for model fitting and more definitions of the applied statistical tests are provided in Table 3.

Software upgrades

To explore whether software upgrades resulted in an abrupt “step” change in the temperature-adjusted T_1 reads, we performed piece-wise linear regression to check for any segmented relationship between the covariates “scan day” and “tube T_1 ” (considering tube “F” at 1.5 T and “M” at 3 T) [21]. For any broken-line relationship discovered, we defined slope parameters and break points where the linear relation/s changed and temporally correlated these with DICOM software metadata.

Results part 1—Evaluation of the phantom

Structural integrity

Individual inspection of all the localizers submitted by sites with each phantom dataset revealed no visible gel rips or tears down any of the tubes. At baseline and at 2 years following batch manufacturing (Supplementary Movie 1), phantoms were free of air bubbles and susceptibility artifacts at both field strengths. High-resolution imaging showed no evidence of gel rips or tears down any of the tubes, and the gels were intact in the mid slice—the piloted location for serial mapping (Fig. 1, left panel). T_1 maps collected through the midline of the phantom, using the specified TIMES scan setup, were free from off-resonance artifacts.

“Reference” rT_1 and rT_2 data

Phantom measurements (averaged across all tubes) collected at the Royal Brompton Hospital showed that temperature-

corrected rT_1 and rT_2 values at 2 years were stable compared to baseline. At 1.5 T, the 2-year temperature-adjusted $\%T_1$ change was $1.95 \pm 1.39\%$ SD, 0.37% SEM. At 3 T, the 2-year temperature-adjusted $\%T_1$ change was $2.22 \pm 1.44\%$ SD, 0.25% SEM. Three tesla measurements at PTB showed a 2-year temperature-adjusted $\%T_1$ change of $0.80 \pm 0.49\%$, SEM 0.16% (Supplementary Figure S1A). Although not the aim of this work, all reference T_2 parameters remained stable with relative <1% change at PTB (temperature-adjusted $\%T_2$ change over 2 years at 3 T = $1.65 \pm 0.26\%$ | 0.09%, Supplementary Figure S1B) and <2% change over 2 years at Royal Brompton Hospital.

Sequences from Siemens and Philips at 1.5 T collected at Royal Brompton Hospital showed strong correlation with rT_1 times with small offsets, as shown in Fig. 4 (left panel). There were no statistically significant differences between the rT_1 correlations for a given sequence when comparing Siemens to Philips platforms (Supplementary Table 1, panel C) or when individual sequences were compared on Philips (Supplementary Table S1, panel B). However, on Siemens, rT_1 correlations for SASHA were significantly stronger than for both MOLLI and ShMOLLI (Supplementary Table S1, panel A).

Sequences from Siemens and Philips at 3 T showed strong correlation with rT_1 times but with visibly imperfect absolute measurement of T_1 across tubes of longer T_1 times as shown in Fig. 4 (right panel). The correlations between absolute measurements of T_1 and rT_1 were similar for MOLLI and SASHA on Philips (Supplementary Table S2, panel B). On Siemens, the correlation between absolute measurements of T_1 and rT_1 was significantly stronger for SASHA when compared to MOLLI and ShMOLLI (Supplementary Table S2, panel A). The correlation between absolute measurements of T_1 and rT_1 for SASHA was significantly stronger on Siemens compared to Philips (Supplementary Table S2, panel C). Comparison of errors at 3 T relative to 1.5 T showed no significant difference (average SEM 5.49 vs. 4.04 respectively, $P = 0.428$).

Temperature sensitivity

Temperature experiments at both NIST and PTB using spin echo, long TR, and sequences (Fig. 2i) consistently showed that short- T_1 tubes (modeling post-GBCA myocardium and blood) were more stable with temperature than very long- T_1 tubes (native blood), where T_1 increased more significantly with temperature. T_1 increased by ~ 11 ms/ $^{\circ}$ C for a typical MOLLI 5s(3s)3s 1.5 T dataset in tube “B” (normal native blood). Conversely, T_1 decreased by ~ 1.2 ms/ $^{\circ}$ C in tube “G” (short post-GBCA myocardium). Temperature corrections for each of the nine tubes at both field strengths from the pooled multi-center analysis are presented separately (Supplementary Table S3) from temperature sensitivity

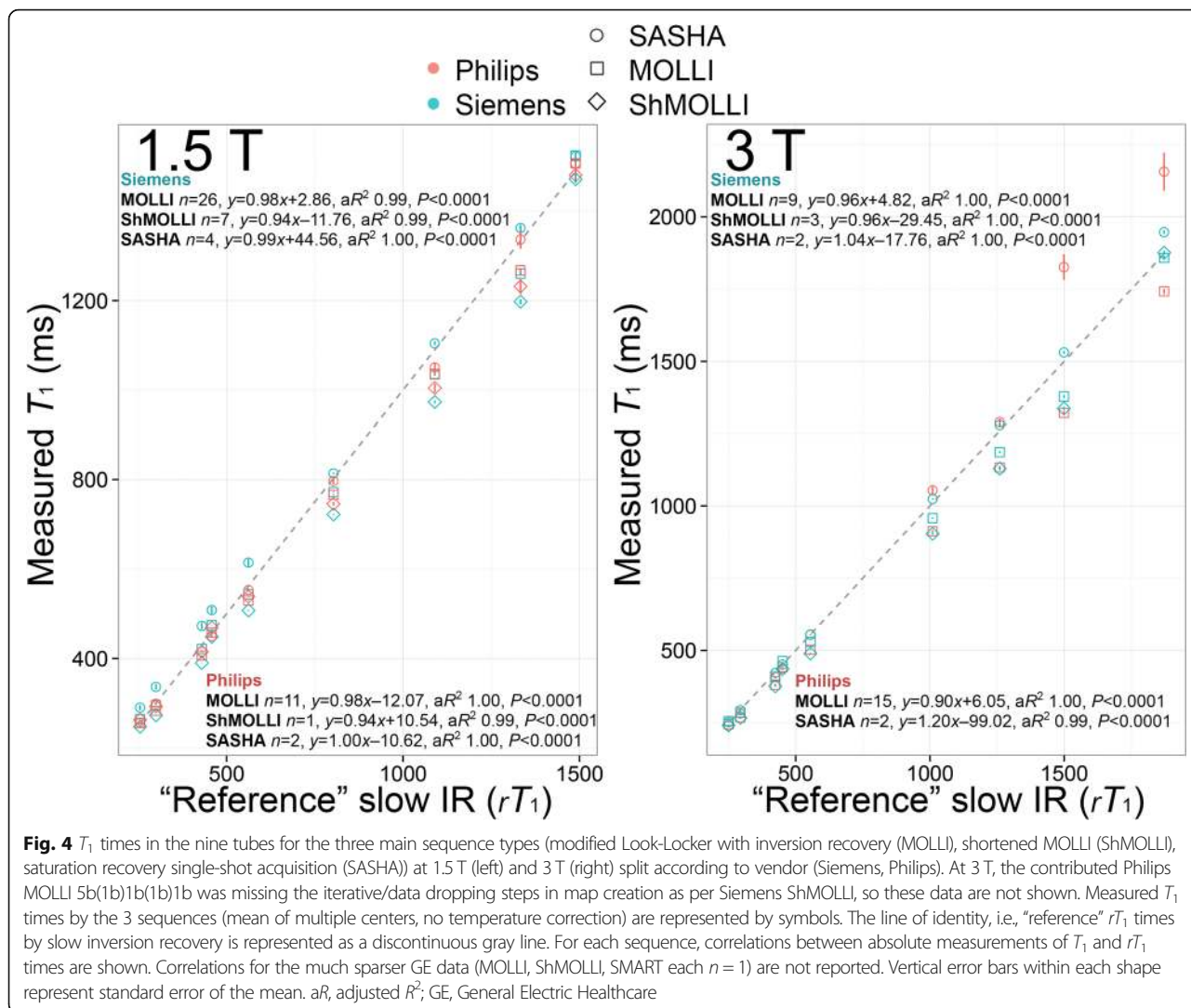


Fig. 4 T_1 times in the nine tubes for the three main sequence types (modified Look-Locker with inversion recovery (MOLLi), shortened MOLLi (ShMOLLi), saturation recovery single-shot acquisition (SASHA)) at 1.5 T (left) and 3 T (right) split according to vendor (Siemens, Philips). At 3 T, the contributed Philips MOLLi 5b(1b)1b(1b)1b was missing the iterative/data dropping steps in map creation as per Siemens ShMOLLi, so these data are not shown. Measured T_1 times by the 3 sequences (mean of multiple centers, no temperature correction) are represented by symbols. The line of identity, i.e., “reference” rT_1 times by slow inversion recovery is represented as a discontinuous gray line. For each sequence, correlations between absolute measurements of T_1 and rT_1 times are shown. Correlations for the much sparser GE data (MOLLi, ShMOLLi, SMART each $n = 1$) are not reported. Vertical error bars within each shape represent standard error of the mean. aR , adjusted R^2 ; GE, General Electric Healthcare

experiments of NIST and PTB due to the latter’s use of bespoke standard operating procedures.

B_0 and B_1 uniformity

B_0 uniformity at 2 years was delivered to within ± 9 Hz at 1.5 T and ± 10 Hz at 3 T (Fig. 5a–b). At 2 years after manufacture, the B_1 field distortion caused by the phantom, continued to be adequately flattened to within 10% of the B_1 at the center of the phantom, at both 1.5 T and 3 T (Fig. 5c–d).

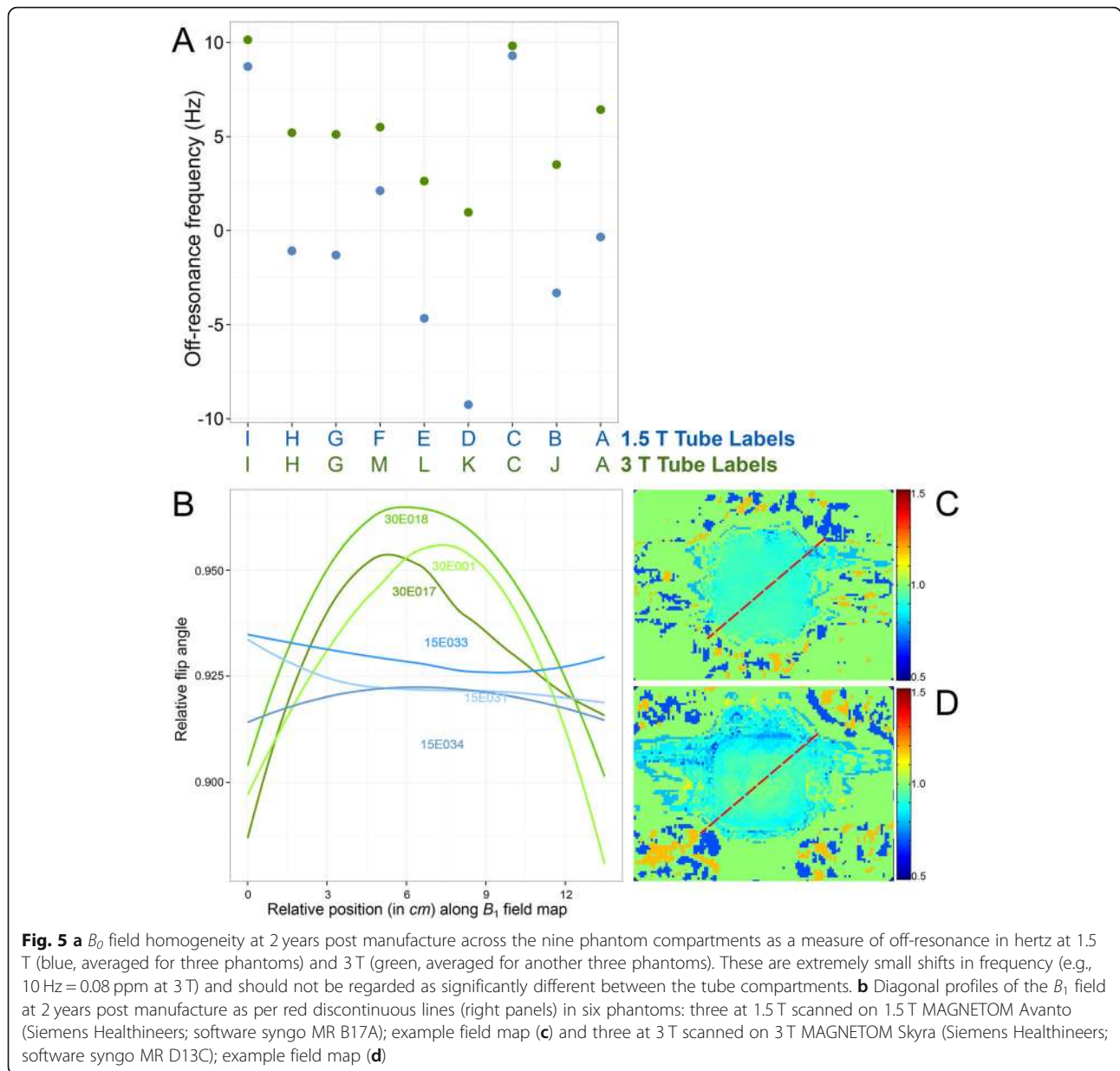
Results part 2—Multi-center phantom testing

Thirty-four of the magnet systems appraised were 1.5 T and 22 were 3 T (see [Supplementary Data File](#) and [Supplementary Table S4](#)) with manufacturers being 35 Siemens, 18 Philips, and 3 GE. Scan frequencies are reported in the [Supplementary Data File](#). The planned program was 1 year, but 24 centers voluntarily extended,

resulting in mean longitudinal data of 358 days, and the longest 1161 days.

Quality assurance of the contributed DICOM image files identified significant protocol deviations or sequence reconstruction failures in nine submissions, the majority of which took place at the start of the study (examples in [Supplementary Fig. S2](#)). These were excluded with subsequent advice sent to centers permitting later inclusion in most cases, except where a tube artifact affecting > 25% of a single center’s series led to exclusion of that tube from pooled analysis (see [Supplementary Data File](#)).

Fifty-two magnets non-exclusively contributed MOLLi datasets, 16 ShMOLLi, and 12 SASHA, and 2 magnets contributed SMART T_1 maps. Center-, session-, and sequence-specific T_1 mapping contributions are detailed in [Supplementary Data File](#). During the project, there were three software upgrades at centers: two Siemens and one GE (details in [Supplementary Fig. S4](#)).



T_1 repeatability

For serial multi-center data, temperature-unadjusted CoV per tube, per phantom, per sequence, and per magnet are detailed in the [Supplementary Data File](#). Temperature-adjusted CoV for the various native sequences together with inter-sequence and cross-platform differences, at 1.5 T and 3 T, are summarized in Tables 1 and 2 respectively and data for post-GBCA in Supplementary Tables S5 and S6. Results of native and post-GBCA sequence/scanner software version repeatability are provided in Tables 1 and 2 and in Supplementary Tables S5 and S6 respectively.

Based on these temperature-adjusted data, mean (\bar{x}) CoV were generally higher (implying poorer repeatability)

at 3 T than at 1.5 T and for the much sparser GE data over Siemens/Philips. Over 1 year, many 1.5 T and 3 T sequences/magnets were repeatable with \bar{x} CoV < 1% and < 2% respectively. For sequences optimized for native T_1 mapping applied to T1MES native tubes, we observed excellent repeatability for several sequences, for example, Siemens MOLLI 5s(3s)3s 448B (\bar{x} CoV = 0.27%) and Philips MOLLI 3s(3s)5s (\bar{x} CoV 0.54%) at 1.5 T (Table 1) and Philips MOLLI 3b(3s)5b (\bar{x} CoV 0.33%) and Siemens ShMOLLI (\bar{x} CoV 0.69%) at 3 T (Table 2).

Among sequences optimized for post-GBCA T_1 mapping applied to T1MES post-GBCA tubes, excellent repeatability was observed for several sequences, for example, Siemens ShMOLLI 1041B (\bar{x} CoV 0.21%) and

Table 1 Temperature-adjusted (normalized to 21 °C) native T_1 and CoV (%) at 1.5 T summarized by vendor and sequence

Summary table of \bar{x} CoV of T_1 (%) across 4 tubes			Order of tube IDs follows their orientation in the scanned bottle			
Siemens	MOLLI 5s(3s)3s [448B]	0.27				
Philips	MOLLI 3s(3s)5s	0.54				
Siemens	SASHA	0.56				
Siemens	ShMOLLI 5b(1b)1b(1b)1b [1041B]	0.64				
Philips	SASHA	0.92				
Philips	ShMOLLI 5b(1b)1b(1b)1b ^c	1.04				
General Electric	MOLLI 5b(1b)1b(1b)1b ^d	1.28				
General Electric	SMART	3.00				
Platform	Sequence [prototype, #]		\bar{x} CoV of T_1 (%), global $\bar{x} \pm SD$ of T_1 according to native tube ID [tube rT_1 by slow IR]			
			F [1090 ms]	E [1333 ms]	D [803 ms]	B [1489 ms]
Siemens	MOLLI MyoMaps product 5s(3s)3s [5]		1.07, 1041 ± 23	1.38, 1273 ± 35	0.85, 771 ± 6	1.45, 1538 ± 46
	MOLLI 3b(3s)3b(3s)5b [448, 3]		0.87, 1019 ± 11	1.08, 1225 ± 17	0.65, 765 ± 6	1.19, 1462 ± 21
	MOLLI 5b(3s)3b [448, 3]		0.18, 1005 ± 13	0.53, 1217 ± 24	0.26, 762 ± 6	0.74, 1464 ± 23
	MOLLI 5b(3s)3b [448B, 2]		0.65, 1052 ± 7	0.82, 1289 ± 10	0.61, 784 ± 4	1.00, 1555 ± 16
	MOLLI 5b(3s)3b [780, 2]		0.70, 1048 ± 14	1.18, 1276 ± 20	0.61, 776 ± 6 sic.	1.15, 1536 ± 24
	MOLLI 5b(3s)3b [780B, 4]		0.76, 1043 ± 9	0.99, 1279 ± 17	0.67, 776 ± 6	1.45, 1550 ± 37
	MOLLI 5s(3s)3s [448, 2]		0.89, 1035 ± 10	1.51, 1261 ± 25	1.04, 770 ± 9	1.44, 1538 ± 24
	MOLLI 5s(3s)3s [448B, 1] ^a		0.14, 1051 ± 1	0.38, 1282 ± 5	0.17, 783 ± 1	0.38, 1553 ± 6
	MOLLI 5s(3s)3s [780B, 2]		0.81, 1043 ± 15	0.69, 1277 ± 15	0.42, 771 ± 6	0.90, 1546 ± 25
	MOLLI 5s(3s)3s [1041, 1]		1.11, 1045 ± 12	0.69, 1285 ± 9	0.81, 772 ± 6	0.96, 1560 ± 15
	MOLLI 5s(3s)3s [1041B, 1]		1.27, 1033 ± 13	1.80, 1232 ± 22	0.66, 771 ± 5	1.47, 1494 ± 22
	ShMOLLI 5b(1b)1b(1b)1b [448, 1]		1.01, 989 ± 10	1.39, 1223 ± 17	0.60, 729 ± 4	1.50, 1510 ± 23
	ShMOLLI 5b(1b)1b(1b)1b [448C, 1]		0.61, 983 ± 6	0.64, 1218 ± 8	0.63, 727 ± 5	1.57, 1507 ± 22
	ShMOLLI 5b(1b)1b(1b)1b [780B, 3]		2.16, 945 ± 34	3.29, 1147 ± 62	0.98, 711 ± 11	3.70, 1392 ± 85
	ShMOLLI 5b(1b)1b(1b)1b [1048, 1]		0.70, 972 ± 7	0.71, 1208 ± 9	0.68, 715 ± 5	1.01, 1489 ± 15
	ShMOLLI 5b(1b)1b(1b)1b [1041B, 1] ^a		0.59, 978 ± 6	0.67, 1201 ± 8	0.43, 721 ± 3	0.86, 1501 ± 13
SASHA [4]		0.39, 1104 ± 17	0.65, 1362 ± 27	0.47, 814 ± 8	0.73, 1522 ± 30	
Philips	MOLLI CardiacQuant product 5s(3s)3s [2]		0.53, 1025 ± 13	0.86, 1265 ± 19	0.50, 760 ± 7	1.04, 1516 ± 28
	MOLLI 3b(3s)3b(3s)5b [2]		0.47, 1006 ± 17	0.94, 1247 ± 39	0.46, 751 ± 6	1.25, 1503 ± 46
	MOLLI 3s(3s)3s(3s)5s [1]		0.94, 1026 ± 10	1.66, 1251 ± 21	0.69, 765 ± 5	2.15, 1450 ± 31
	MOLLI 3s(3s)5s [1] ^a		0.20, 936 ± 2	1.11, 1100 ± 12	0.27, 722 ± 2	0.59, 1323 ± 8
	MOLLI 5b(3s)3b(3s)2b [1]		6.78, 1168 ± 79	6.32, 1374 ± 87	3.04, 816 ± 25	6.53, 1665 ± 109
	MOLLI 5b(3s)3b [1]		1.18, 1027 ± 12	1.48, 1265 ± 19	0.67, 758 ± 5	1.16, 1522 ± 18
	MOLLI 5s(3s)3s [3]		0.89, 1012 ± 17	1.35, 1233 ± 31	1.20, 756 ± 9	1.76, 1456 ± 58
	ShMOLLI 5b(1b)1b(1b)1b ^c [1]		0.89, 1027 ± 9	1.37, 1270 ± 17	0.50, 753 ± 4	1.41, 1522 ± 21
	SASHA [2]		1.02, 1057 ± 44	0.89, 1336 ± 84	0.87, 798 ± 30	0.89, 1505 ± 33
	General Electric	MOLLI 5b(3s)5b [1]		11.18, 653 ± 73	7.69, 853 ± 66	8.45, 591 ± 50
MOLLI 5b(1b)1b(1b)1b ^d [1] ^a			1.40, 575 ± 8	1.76, 642 ± 11	0.78, 543 ± 4	1.19, 1002 ± 12
SMART [1]			2.05, 1063 ± 22	4.76, 1273 ± 61	0.95, 798 ± 8	4.22, 1430 ± 60

The term sequence refers to either MOLLI, ShMOLLI, SASHA, or SMART
MOLLI/ShMOLLI protocol nosology has the number of inversions per experiment as the total count of numbers outside brackets, image cycles are outside brackets, pause cycles are within brackets, and cycle lengths defined in terms of either heart beats (b) or seconds (s)
CoV coefficient of variation, ID identity code, GBCA gadolinium-based contrast agent, GE General Electric, MOLLI modified Look-Locker inversion recovery, rT_1 "reference" slow inversion recovery T_1 , SASHA saturation-recovery single-shot acquisition, SD standard deviation, ShMOLLI shortened MOLLI, SMART saturation method using adaptive recovery times for cardiac T_1 mapping, T Tesla
^aDenotes the T_1 mapping sequence/software combination with lowest overall CoV% for a given vendor where multiples exist. P values for differences in CoV between sequences/vendors are reported for this highly repeatable sequence where multiples exist. Less favorable CoVs (> 1%, see the " T_1 repeatability" section) are in italics. Post-GBCA tubes are not shown here as their data are reported separately in relation to post-GBCA sequences in Supplementary Table 5
^bDenotes the number of different magnets submitting that particular sequence from which the average CoVs were derived
^cUsing iterative/data dropping steps in map creation as per ShMOLLI
^dIn the absence of iterative/data dropping steps in map creation as per ShMOLLI
 \bar{x} = average CoV across the 4 native tubes for a given sequence
 \bar{x} = where more than one sequence type was submitted, individual CoVs were then averaged to derive \bar{x} CoV; while for single sequence submissions \bar{x} CoV is from the global mean $T_1 \pm SD$ for that one sequence

Table 2 Temperature-adjusted (normalized to 21 °C) native T_1 and CoV (%) at 3 T summarized by vendor and sequence

Summary table of \bar{x} CoV of T_1 (%) across 4 tubes			Order of tube IDs follows their orientation in the scanned bottle			
Philips	MOLLI 3b(3s)5b	0.33				
Siemens	ShMOLLI 5b(1b)1b(1b)1b [780C]	0.69				
Siemens	MOLLI 5b(3s)3b [448B]	0.95				
Siemens	SASHA	1.29				
Philips	SASHA	3.76				
GE	SMART	3.76 sic				
Platform	Sequence [prototype, #]	\bar{x} CoV of T_1 (%), global $\bar{x} \pm$ SD of T_1 according to native tube ID [tube rT_1 by slow IR]				
		M [1260 ms]	L [1499 ms]	K [1010 ms]	J [1872 ms]	
Siemens	MOLLI MyoMaps product 5s(3s)3s [4]	1.28, 1182 \pm 20	1.61, 1367 \pm 29	1.02, 955 \pm 13	2.42, 1831 \pm 62	
	MOLLI 3b(3s)3b(3s)5b [448, 1]	1.16, 1092 \pm 13	1.92, 1252 \pm 24	0.56, 889 \pm 5	1.58, 1724 \pm 27	
	MOLLI 5b(3s)3b [448B, 1] ^a	0.36, 1233 \pm 4	0.78, 1449 \pm 11	0.76, 986 \pm 7	0.86, 2012 \pm 17	
	MOLLI 5b(3s)3b [780B, 2]	0.99, 1204 \pm 14	1.13, 1410 \pm 18	0.74, 970 \pm 9	1.50, 1912 \pm 28	
	MOLLI 5s(3s)3s [780B, 1]	1.46, 1222 \pm 18	2.36, 1435 \pm 34	1.76, 980 \pm 17	3.04, 1916 \pm 58	
	ShMOLLI 5b(1b)1b(1b)1b [780B, 2]	1.17, 1155 \pm 27	1.62, 1361 \pm 35	0.80, 922 \pm 17	2.45, 1882 \pm 41	
	ShMOLLI 5b(1b)1b(1b)1b [780C, 1] ^a	0.78, 1109 \pm 9	0.89, 1322 \pm 12	0.64, 888 \pm 6	1.51, 1892 \pm 29	
	SASHA [2]	2.04, 1284 \pm 30	1.15, 1534 \pm 27	0.70, 1025 \pm 13	1.27, 1949 \pm 35	
	Philips	MOLLI 3b(3s)3b(3s)5b [4]	1.64, 1147 \pm 52	2.42, 1343 \pm 80	1.05, 923 \pm 29	3.90, 1785 \pm 115
MOLLI 3b(3s)5b [1] ^a		0.01, 1055 \pm 1	0.49, 1217 \pm 6	0.05, 862 \pm 1	0.78, 1710 \pm 13	
MOLLI 3s(3s)5s [1]		5.41, 1175 \pm 64	3.72, 1392 \pm 52	7.13, 937 \pm 67	3.56, 1833 \pm 65	
MOLLI 5b(3s)3b [3]		4.18, 1098 \pm 48	5.44, 1286 \pm 73	4.37, 868 \pm 42	6.33, 1784 \pm 106	
MOLLI 5s(3s)3s [6]		2.40, 1131 \pm 70	2.41, 1320 \pm 108	2.60, 906 \pm 34	2.71, 1727 \pm 131	
MOLLI 5b(1b)1b(1b)1b ^c [2]		2.98, 948 \pm 97	4.53, 1082 \pm 137	0.96, 794 \pm 60	6.29, 1286 \pm 282	
SASHA [2]		3.11, 1290 \pm 46	3.40, 1826 \pm 171	2.11, 1054 \pm 38	6.41, 2157 \pm 267	
General Electric	SMART [1]	2.85, 1150 \pm 33	4.13, 1227 \pm 51	2.04, 973 \pm 20	6.03, 1419 \pm 86	

Less favorable CoVs (> 2%, see the “ T_1 repeatability” section) are in italics. Post-GBCA tubes are not shown here as their data are reported separately in relation to post-GBCA sequences in Supplementary Table 6

sic = text is quoted exactly as it stands in the original, i.e., this is not a typo. Abbreviations as in Table 1

^aDenotes the T_1 mapping sequence/software combination with lowest overall CoV% for a given vendor where multiples exist

^bDenotes the number of different magnets submitting that particular sequence from which the average CoVs were derived

^cIn the absence of iterative/data dropping steps in map creation as per ShMOLLI

\bar{x} = average CoV across the 4 native tubes for a given sequence

\bar{x} = where more than one sequence type was submitted, individual CoVs were then averaged to derive \bar{x} CoV; while for single sequence submissions \bar{x} CoV is from the global mean $T_1 \pm$ SD for that one sequence

Siemens MOLLI 4s(1s)3s(1s)2s 448B (x CoV 0.26%) at 1.5 T (Supplementary Table S5) and Siemens MOLLI 4b(1s)3b(1s)2b 448B (x CoV 0.10%) and Siemens ShMOLLI (x CoV 0.28%) at 3 T (Supplementary Table S5).

Sources of T_1 variation

Linear mixed models which excluded centers that had experienced a software change during the time of data collection (Table 3 and Supplementary Tables S7 and S8) indicated that temperature-adjusted T_1 differs significantly between sequences and software versions at both field strengths ($P < 0.001$ all) and between magnet models at 1.5 T ($P = 0.011$). Notably, phantom age had no significant effect on T_1 (model A2).

Software upgrades

The two software upgrades on Siemens both occurred towards the end of longitudinal data submissions (Supplementary Fig. S4A, 15E031; 4B, 30E017), so the paucity of data points post-upgrade events precluded statistical testing for significant T_1 shifts. For the software upgrade on GE, however, differences between pre- and post-linear regression slopes (Supplementary Fig. S4C, 30E012) indicate a marginally significant T_1 shift event ($P = 0.024$).

Discussion

To our knowledge, this is the first multi-center study using a wide range of T_1 mapping methods to study the interplay of some of the known sources of measured T_1 variation. The basis of this work is the TIMES phantom, and a major aim of this work was to evaluate how this

Table 3 Linear mixed models for 1.5 T and 3 T multi-center temperature-adjusted T_1 mapping data (normalized to 21 °C, considering the “Medium” native myocardium tubes “F” and “M” respectively). The best model is “A5” at 1.5 T and “A3” at 3 T

	Model fitting formulas	AIC	BIC	Log likelihood	χ^2	P value	Best model
1.5 T model	A1: $T_1 \sim$ sequence + (1 ID)	5850.3	5920.8	- 2909.1	“ref”	“ref”	“ref”
	A2: $T_1 \sim$ sequence + (1 ID/age) [worse fit with age]	5852.3	5927.2	- 2909.1	0.0	1.000	-
	A3: $T_1 \sim$ sequence \times software + (1 ID)	5725.4	5883.9	- 2826.7	164.9	< 0.0001	-
	A4: $T_1 \sim$ sequence \times software + vendor + (1 ID) [vendor non-contributory to fit]	5725.4	5883.9	- 2826.7	0.0	1.000	-
	A5: $T_1 \sim$ sequence \times software + model + (1 ID)	5723.4	5890.8	- 2823.7	5.9	0.051	A5
	A6: $T_1 \sim$ sequence \times software + vendor \times model + (1 ID) [vendor non-contributory to fit]	5723.4	5890.8	- 2823.7	0.0	1.000	-
Final model (A5) parameters		Variance	SD	χ^2	P value		
Random effects	ID	2547.2	50.47	“ref”	“ref”		
Fixed effects	Sequence (β range - 32.3 to 455.6 ^a)	/	/	1101.0	< 0.0001		
	Software (β range 385.1 to 522.3 ^a)	/	/	63.8	< 0.0001		
	Model (β range - 161.8 to 16.0 ^a)	/	/	11.2	0.011		
	Sequence: Software (β range - 161.8 to - 6.9 ^a)	/	/	149.6	< 0.0001		
3 T model	A1: $T_1 \sim$ sequence + (1 ID)	4390.4	4447.0	- 2181.2	“ref”	“ref”	“ref”
	A2: $T_1 \sim$ sequence + (1 ID/age) [worse fit with age]	4392.4	4453.1	- 2181.2	0.0	1.000	-
	A3: $T_1 \sim$ sequence \times software + (1 ID)	4238.3	4339.4	- 2094.2	174.1	< 0.0001	A3
	A4: $T_1 \sim$ sequence \times software + vendor + (1 ID) [vendor non-contributory to fit]	4238.3	4339.4	- 2094.2	0.0	1.000	-
	A5: $T_1 \sim$ sequence \times software + model + (1 ID)	4239.2	4348.4	- 2092.6	3.1	0.212	-
	A6: $T_1 \sim$ sequence \times software + vendor \times model + (1 ID) [vendor non-contributory to fit]	4239.2	4348.4	- 2092.6	0.0	1.000	-
Final model (A3) parameters		Variance	SD	χ^2	P-value		
Random effects	ID	372.4	19.3	“ref”	“ref”		
Fixed effects	Sequence (β range - 114.5 to 38.5 ^a)	/	/	732.8	< 0.0001		
	Software (β range - 203.0 to 32.7 ^a)	/	/	46.0	< 0.0001		
	Sequence: Software (β range - 48.8 to 267.3 ^a)	/	/	189.0	< 0.0001		

The symbol (1||ID) in the model formulas refers to the random effect of individual phantoms (by identity number). At 1.5 T models, A5 and A6 have equal AIC/BIC but given the lack of statistically significant χ^2 improvement from A5 to A6 ($P = 1.0$); A5 is considered the most parsimonious model. Age refers to phantom age at scanning since date of manufacture, AIC Akaike information criterion, BIC Bayesian information criterion, χ^2 chi-square, ref reference, SD standard deviation

^aComplete list of β coefficients for model A5 at 1.5 T and A3 at 3 T are provided in Supplementary Tables 7 and 8 respectively

performed in use at multiple centers. The phantom appears sufficiently robust for T_1 mapping quality assurance purposes. Based on our data, estimated phantom T_1 values may show substantial variation between different T_1 mapping sequences, scanner software versions, and potentially also scanner models, and temperature correction, at least for native T_1 tubes, is necessary to achieve the desired repeatability. Inspection of the Supplementary Table also suggests that there are occasional performance variations producing outliers even within a given T_1 mapping sequence prototype/software combination across different magnets. In spite of this variation, however, several specific combinations of field strength, sequence, and scanner generally exhibit excellent repeatability. Given the choices, the CMR community may prefer to standardize and use combinations with high

repeatability for future clinical and research use, e.g., CoV < 1% at 1.5 T and < 2% at 3 T, while seeking to optimize less repeatable combinations. Less data were available for GE in T1MES, so the observed variability requires verification in a larger more representative sample.

Agreement with “reference” slow scanning, rT_1 times was slightly greater for SASHA compared to MOLLI or ShMOLLI, which both slightly underestimated T_1 , mainly from the known T_2 -related underestimation which is larger when measuring the myocardium [13–15], (Fig. 4, Supplementary Tables S1 and S2) and not due to magnetization transfer, which is negligible in these agar phantoms [22].

The phantom data presented here suggest that the “ T_1 standard” framework remains possible, but the wide

variety of different sequence options, vendors, and field strengths distributed the data over too many categories for reliable modeling, unlike the “locked-down” approach mentioned below which does have that strong advantage. Further work will explore the transferability of clinical measurement based on in vitro phantom calibration.

TIMES is but one of a number of phantom objects that have been used or proposed to support T_1 mapping quality assurance (elaborated in Table 4). Some alternatives that target native and post-GBCA myocardial and blood relaxation times in support of T_1 mapping work are (1) Brompton phantom by Vassiliou et al. [23], (2) Hypertrophic Cardiomyopathy Registry [24] phantom (HCMR) by Piechnik et al., and (3) International Society for Magnetic Resonance in Medicine (ISMRM)/NIST MR imaging phantom [25].

At 1 year, CoV for T_1 tubes of the Vassiliou [26] phantom on a single Siemens scanner ranged from 1.0 to 3.6% using only the native MOLLI 5s(3s)3s sequence, compared to 0.27 to 3.0% in TIMES (for 1.5 T MOLLI 5s(3s)3s [448B] on Siemens to SMART on GE respectively). This within sequence precision heterogeneity, as

already alluded to by Kellman et al. [27], linked to protocol modifications within MOLLI, may partly explain some of the higher CoV for specific MOLLI sequences compared to ShMOLLI where centers obligatorily scanned using a fixed 5b(1b)1b(1b)1b sequence with conditional fitting. As mentioned above for the dispersion of TIMES results, this “real-world” heterogeneity of T_1 mapping sequences highlights that while T_1 mapping research and innovation calls for multiple flexible prototypes, the resultant diversity poses a nontrivial challenge to multi-center standardization. Conversely, the more “locked-down” approach like that adopted for ShMOLLI, albeit less editable by external researchers, potentially facilitates standardization.

The design of phantoms for this purpose (although often seen as trivial) is challenging: to have a large sufficient ROI in each sample tube and a good number of sample tubes in the main jar of the phantom, the overall size of the phantom increases to the point that B_1 non-uniformity at 3 T due to the dielectric permittivity of any water-based phantom becomes problematic. While of course B_1 is non-uniform in vivo, one purpose of phantom tests is to eliminate uncertainty or at least enable

Table 4 Comparison of recently reported phantoms for cardiac T_1 mapping quality assurance

	Dedicated T_1 device		Combined T_1/T_2 device	
Phantom	HCMR (cardiac specific)	Brompton (cardiac specific)	TIMES (cardiac specific)	ISMRM/NIST system phantom (not cardiac specific)
Field strength specificity				
Tube ingredients	NiCl ₂ -doped agar + carrageenan	NiCl ₂ -doped agar	NiCl ₂ -doped agar	NiCl ₂ -doped water, MnCl ₂ -doped agar
Structure	9 T_1/T_2 tubes in an amber PVC sealed jar that is ~25% smaller in total volume compared to TIMES.	Air-filled box containing 4 duplicate T_1/T_2 glued glass tubes (total 8) requires cylindrical MRI test bottles on either side for B_1 field/reference frequency calibration.	9 plastic T_1/T_2 tubes in an amber PVC sealed jar with inter-tube gaps packed with a carefully specified agarose/ HDPE plastic macrobead/ NiCl ₂ -doped fill that flattened the B_1 field and provided sufficient B_0 homogeneity to obviate the need for side phantoms.	A layer of 14 T_1 spheres and a separate layer of 14 T_2 spheres so no unified T_1/T_2 compartment representing the relaxation parameters of the human heart.
Cardiac T_1/T_2 coverage	Health and disease, 9 biologies but limited T_2 coverage (57 and 75 ms).	Health only, 4 biologies: (1) native myocardium, (2) native blood, (3) post-GBCA myocardium, and (4) post-GBCA blood.	Health and disease, 9 biologies and broad T_2 coverage (1.5 T, 44, 48, 50, 155, 189, and 243 ms). Also see Supplementary Table 3.	Health and disease, but of the 14 T_1 spheres (ranging from 21 to 2038 ms), half are not useful to cardiac T_1 mapping as T_1 times are too short for either native or post-GBCA myocardium/blood. Half the T_2 spheres (ranging from 11 to 581 ms) are also anti-physiological for cardiac mapping. Also, the T_1/T_2 ratio is not representative of cardiac tissue.
MT coverage	–	–	–	–
Regulatory clearance	–	–	FDA, CE-mark	–
Developers	Piechnik et al.	Vassiliou et al.	Captur et al.	NIST/ISMRM

CE Conformité-Europeen, FDA Food and Drug Administration, HDPE high-density polyethylene, MT magnetization transfer, PVC poly vinyl chloride

controlled testing of factors such as B_1 rather than introduce their own errors. Conversely, if the phantom is made very small, then the truly acquired pixel size of in vivo T_1 mapping methods (which obviously must not be modified or adapted [28] for the phantom scans) becomes important compared to the sample tube inner diameters leading to questions over SNR [29] and the impact of Gibbs artifact, for example.

Another obstacle for relaxation time phantoms (on top of basic water T_1 and T_2 temperature and pressure variations, gel instabilities [30, 31], small leakages, dehydration, or ion migration effects over time) is the differing temperature dependence of each species of paramagnetic ion's relaxivity (r_1 , r_2) and furthermore frequency dispersion in all of these physical properties between 60 MHz and 120 MHz [32–36]. Efforts to use alternative chemistry have so far not provided the range of T_1 and T_2 needed for this application.

The results confirmed, as is widely known in the field, that the T_1 measured by even nominally identical mapping sequences can differ significantly between software revisions. Scanner software upgrades are not uncommon in centers, and we encountered three such events among contributing partners. There was a measurable shift on at least one system, with potentially important implications for the field as alluded to in the 2017 consensus statement on multi-parametric mapping by the Society for Cardiovascular Magnetic Resonance (SCMR) [37]. Future work using phantoms at scale, across vendors and over time, would ideally continue monitoring center-specific data before and after potential shift events to more fully understand their impact on T_1 allowing the community to set up mitigation approaches.

Clearly, sources of T_1 variance and drift over time and between centers are numerous. Some T_1 variance is related to differences between sequences and magnet platforms: to the scanner itself (equipment drift or equipment shifts from updates of software, replacement of parts, routine service recalibrations, etc.), to the environment (temperature, pressure, humidity), and to the operator (phantom positioning inside scanner bore). In a phantom study, some T_1 variance may additionally have been due to intra-phantom differences (liquid/agar status over time collectively contributing to phantom aging) and inter-phantom manufacturing variations. In T1MES, we mitigated potential inter-phantom manufacturing variations by exercising extreme caution in design and prototyping and by the strict batch manufacturing processes to medical device grade standards. Drifts in the system setup (long-term stability) would tend to be “adjusted” out by the shim and center frequency (CF) and transmitter B_1 reference (TxREF) adjustments that could otherwise bias estimated T_1 . Any receiver coil or gain changes would most likely cancel out of T_1 maps as

constant across the series of input images to T_1 derivation (appearing in A and B maps instead). It is considered unlikely that gradient performance would drift significantly, and any significant sudden changes in that are also unlikely even when the eddy-current compensation is adjusted on routine service visits. However, we have shown how unforeseen software changes could potentially cause “step-wise” changes in the estimated T_1 records. To directly model the interplay between all these potential sources of T_1 variation in longitudinal phantom studies would be a daunting task, and we considered only a subset of these sources in our analyses.

Though B_0 and B_1 homogeneity tests in T1MES (Fig. 5) were stable, we did measure slightly greater field inhomogeneity at the phantom edge and corners compared to the core (phantom corners and edges are the zones which our T1MES data flagged up as areas of least B_0/B_1 homogeneity). The tubes with the highest T_1 times are the tubes most susceptible to such field inhomogeneities, and this suggests that a future iteration of T1MES could be further optimized by rearrangement to position such tubes in the most uniform regions.

The results of this study are relevant to clinicians or researchers choosing sequences for T_1 mapping, and we highlight 3 take home messages as follows:

- i. Sequence/software combinations in Tables 1 and 2, with a CoV $\leq 1\%$ and 2% respectively, are sufficiently repeatable for clinical/research use (see rationale in the “Methods part 3—Statistical analysis” and “ T_1 repeatability” sections). For the native myocardium T1MES tube at 1.5 T (reference $T_1 \sim 1037$ ms by MOLLI), a $\leq 1\%$ CoV is well within ± 1 SD of normal values for T_1 , meaning that small-magnitude biological changes (e.g., diffuse myocardial fibrosis), will be resolvable with high precision [37]. By contrast, CoVs of 3 or 6% at 1.5 T (variance of 31 and 62 ms respectively) will undermine a center's ability to resolve small-magnitude changes, and they may also impact the resolution of large-magnitude biological changes [37] (like amyloid, iron overload, Fabry disease, acute myocardial injury). Our data indicate that many of the investigated sequences for Siemens and Philips at 1.5 T exhibited CoVs $\leq 1\%$. At 3 T, MOLLI (Siemens and Philips), ShMOLLI (Siemens), and SASHA (Siemens) exhibited CoVs $\leq 2\%$. The absence of GE from this list is due to limited participation.
- ii. For developers, the highlighting of sequence performance may encourage work to refine sequences in some cases.
- iii. The prospect of using phantom calibration to remove the need for local reference ranges [37] is

potentially becoming more tangible than before, yet more work is needed to deliver such calibration.

These data may facilitate the use of T_1 mapping as a useful clinical biomarker. T_1 standardization will also be important to facilitate clinical research. At the present time, there are 25 registered clinical research studies using T_1 mapping (clinicaltrials.gov accessed January 2019).

Limitations

Presented results are not intended to compare the SNR obtained in the phantom setup in different machines but are aimed at studying long-term stability. In spite of our outreach to advertise enrollment at study start, GE centers are under-represented. The scarcity of multi-center data for many of the specific T_1 mapping sequence, software, and field-strength combinations has limited our ability to deliver a “ T_1 standard”. Eight centers enrolled, received a phantom, but then could not deliver the serial phantom scans, and four centers did not meet the 1 year minimum data submission requirement. Each center provided legitimate justifications for the missing data, and furthermore, we highlight that contributions to T1MES were entirely voluntary, with no center receiving remuneration for any staff or scanner time invested in the program.

In total, 3 magnets deviated from the prescribed nominal intervals in some of their initially submitted datasets. Phantom repeatability studies represent an in vitro experiment. Some sequences and scanners that performed well in vitro may exhibit greater variability in the face of biological tissue (magnetization transfer, flow, motion), patient characteristics (fast/slow heart rates, heart rate variability, breath-hold length, implanted metal devices), and scanning process (non-isocentering, scanning in multiple planes). The T1MES phantom does not capture magnetization transfer. The biomarker performance that was measured here accounts for T_1 mapping in vitro. While this is obviously related to diagnostic performance in vivo, cardiac motion, etc., introduce yet another degree of freedom, and different hardware/software combinations might deal with this differently. In addition, CoV as a single metric may not be the best way to pick a pulse sequence for repeatability when the data is only derived from a static phantom. A very precise (reproducible) T_1 sequence that requires long breatholds or is heart rate-variability-sensitive might perform very well with the T1MES phantom setup but perform poorly in patients with varying biology, e.g., that cannot hold their breath well or who are in atrial fibrillation. The phantom has no intracellular/extracellular component, so any researcher seeking to model “ECV” using pre- and post-GBCA T_1 mapping values

derived from T1MES (see example data in Supplementary Table S9) must bear this in mind.

The phantom design, ROI size, and coil setup instructions all aimed to make the fundamental image noise SNR an irrelevant factor in the CoV results. The artificially high and “clean” SNR of a phantom setup (if without phantom-related distortions, as we have shown in the “ B_0 and B_1 uniformity” section) is clearly not directly an indicator of satisfactory in vivo performance. It contains many sequence-related factors that may have different consequences in vivo [29], and such SNR comparisons are *not* the aim of this work. Ideally, participating sites would have proved this by sending multiple T_1 maps per session. However, contributions were already voluntary, and further demands on sites could not be supported. For the same reason, it was not feasible to demand long reference T_1/T_2 data for each participating site, and this was never part of the original enrollment commitment. As some indication of the typical short-term thermal noise random jitter, the raw (temperature-unadjusted) T_1 values from some sites that did anyway submit several repeated T_1 maps each session are plotted in Supplementary Fig. S3.

Conclusion

The T1MES phantom developed to CE/FDA manufacturing standards for T_1 mapping is a robust quality assurance device. T_1 mapping can now be quality assured on a multi-center scale, fulfilling a key requirement for the use of T_1 mapping in clinical decision-making or as a surrogate endpoint in drug trials. A good number of T_1 measurement sequence/scanner model/vendor/field strength combinations are remarkably repeatable, but some combinations less so, with coefficients of variation exceeding 1–2% over 1 to 2 years. Given the alternatives, we recommend that combinations with poor repeatability are deprioritized for clinical and research use. In spite of the use of a large number of different sequence prototypes and products in this study, a number of agreements and encouraging results are reported. Phantom calibration of T_1 mapping which obviates the need for local reference ranges, enabling the establishment of a “ T_1 standard” to facilitate multi-center T_1 studies, comes closer, with further work required to address this.

Supplementary information

Supplementary information accompanies this paper at <https://doi.org/10.1186/s12968-020-00613-3>.

Additional file 1: Supplementary Movie 1. High-resolution imaging of phantom 30E017 at baseline (October 2015, **Left panel**) and at two years post manufacturing (**Right panel**). (MP4 69,750 kb)

Additional file 2: Supplementary file containing list of Supplementary Tables and Figures referenced in the main text.

Additional file 3. Supplementary data file reporting all the center-, session- and sequence-specific T_1 mapping contributions to the T1MES program.

Abbreviations

3D: Three-dimensional; AIC: Akaike information criteria; BIC: Bayesian information criteria; CE: Conformité Européenne; CF: Center frequency; CMR: Cardiovascular magnetic resonance; CoV: Coefficient of variation; DICOM: Digital Imaging and COmmunications in Medicine; ECV: Extracellular volume; FA: Flip angle; FDA: Food and Drug Administration; GBCA: Gadolinium-based contrast agent; GE: General Electric Healthcare; HCMR: Hypertrophic Cardiomyopathy Registry; INR: International normalized ratio; IRSE: Inversion recovery spin echo; ISMRM: International Society for Magnetic Resonance in Medicine; MOLL: Modified Look-Locker inversion recovery; NIST: National Institute of Standards and Technology; PTB: Physikalisch-Technische Bundesanstalt; QIBA: Quantitative Imaging Biomarkers Alliance; REDCap: Research electronic data capture; ROI: Region of interest; RSNA: Radiological Society of North America; rT_1 : "Reference" T_1 ; rT_2 : "Reference" T_2 ; SASHA: Saturation recovery single-shot acquisition; SCMR: Society for Cardiovascular Magnetic Resonance; SD: Standard deviation; ShMOLL: Shortened MOLL; SMART: Saturation method using adaptive recovery times for cardiac T_1 mapping; SNR: Signal-to-noise ratio; T: Tesla; T1MES: T_1 mapping and extracellular volume standardization; TE: Echo time; TR: Repetition time; TxREF: Transmitter B_1 reference

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Authors' contributions

JCM together with PG and GC conceived the project. PG provided continuous expert physics support for the project, oversaw the experimental testing of phantoms from prototype out to 2 years, scanned the phantom, expertly reviewed the manuscript and analysis, and assisted with all aspects of the project. GC and PG designed the prototypes and final phantom. GC curated the phantom data, managed the consortium, developed the analytical pipeline (with help from PK and RB), organized and analyzed the data, scanned the phantom including experimental tests, performed the statistical analysis, wrote the manuscript, and built the figures. JCM expertly reviewed the manuscript and assisted with all aspects of the project. AB, YY, RJE, GB, CT, LR, and NF analyzed the data. RB, BI, and KK performed the PTB and NIST experiments respectively. RB and PK provided analytical support and scanned the phantom. DMH, AG, DAB, RZ, TD, and JJ provided expert appraisal of the manuscript and analytical approaches. JPG, LEMP, CBL, CBD, HJL, RS, SWL, CB, SV, AF, CdL, FDC, MV, PC, DD, AJT, FPM, SP, DM, MAM, SMG, ML, JJ, and RN scanned the phantom. MS and NC provided expert advice. All authors reviewed the manuscript. The authors read and approved the final manuscript.

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Availability of data and materials

Datasets collected for the current study are available from the corresponding author upon reasonable request.

Ethics approval and consent to participate

Not applicable

Consent for publication

Not applicable

Competing interests

An industrial group (Resonance Health) had been funded to mass-produce the described T1MES phantoms to CE mark/FDA standards to permit international distribution in the T1MES academic program. WP is an employee of Resonance Health. AG is an employee of Siemens Healthineers. PH is an employee of Philips Healthcare. No financial or non-financial competing interests exist for any of the other authors.

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