GUIDELINES

A standard system phantom for magnetic resonance imaging

Karl F. Stupic1 | Maureen Ainslie2 | Michael A. Boss3 | Cecil Charles2 |
Andrew M. Dienstfrey1 | Jeffrey L. Evelhoch4 | Paul Finn5 | Zydrunas Gimbutas1 |
Jeffrey L. Gunter6 | Derek L. G. Hill7 | Clifford R. Jack6 | Edward F. Jackson8 |
Todor Karaulanov9 | Kathryn E. Keenan1 | Guoying Liu10 | Michele N. Martin1 |
Pottumarthi V. Prasad11 | Nikki S. Rentz1 | Chun Yuan12 | Stephen E. Russek1 |

1Physical Measurement Laboratory, National Institute of Standards and Technology, Boulder, Colorado, USA
2Department of Radiology, Duke University, Durham, North Carolina, USA
3American College of Radiology, Philadelphia, Pennsylvania, USA
4Merck Research Laboratories, West Point, Pennsylvania, USA
5University of California, Los Angeles, California, USA
6Department of Radiology, Mayo Clinic, Rochester, Minnesota, USA
7Centre for Medical Image Computing, University College London, London, United Kingdom
8Medical Physics, University of Wisconsin, Madison, Wisconsin, USA
9CaliberMRI, Inc., Boulder, Colorado, USA
10National Institute of Biomedical Imaging and Bioengineering, National Institutes of Health, Bethesda, Maryland, USA
11Radiology/CAI, NorthShore University HealthSystem, Evanston, Illinois, USA
12Radiology, University of Washington, Seattle, Washington, USA

Correspondence
Karl F. Stupic, Physical Measurement Laboratory, National Institute of Standards and Technology, 325 Broadway, Boulder, CO 80305, USA.
Email: karl.stupic@nist.gov

Purpose: A standard MRI system phantom has been designed and fabricated to assess scanner performance, stability, comparability and assess the accuracy of quantitative relaxation time imaging. The phantom is unique in having traceability to the International System of Units, a high level of precision, and monitoring by a national metrology institute. Here, we describe the phantom design, construction, imaging protocols, and measurement of geometric distortion, resolution, slice profile, signal-to-noise ratio (SNR), proton-spin relaxation times, image uniformity and proton density.

Methods: The system phantom, designed by the International Society of Magnetic Resonance in Medicine ad hoc committee on Standards for Quantitative MR, is a 200 mm spherical structure that contains a 57-element fiducial array; two relaxation time arrays; a proton density/SNR array; resolution and slice-profile insets. Standard imaging protocols are presented, which provide rapid assessment of geometric distortion, image uniformity, $T_1$ and $T_2$ mapping, image resolution, slice profile, and SNR.
Results: Fiducial array analysis gives assessment of intrinsic geometric distortions, which can vary considerably between scanners and correction techniques. This analysis also measures scanner/coil image uniformity, spatial calibration accuracy, and local volume distortion. An advanced resolution analysis gives both scanner and protocol contributions. SNR analysis gives both temporal and spatial contributions.

Conclusions: A standard system phantom is useful for characterization of scanner performance, monitoring a scanner over time, and to compare different scanners. This type of calibration structure is useful for quality assurance, benchmarking quantitative MRI protocols, and to transition MRI from a qualitative imaging technique to a precise metrology with documented accuracy and uncertainty.

KEYWORDS
MRI standards, phantom, quality assurance, quantitative MRI

1 | INTRODUCTION

With the rise of quantitative MRI to complement qualitative MRI, it is imperative to assess system stability and understand the comparability of measurements across MRI scanners. One solution is to use a standard imaging phantom, an inanimate object used to characterize or calibrate an imaging system. A standard phantom would be one that is commonly used and accepted by the imaging community; has stable, precisely defined properties to allow monitoring of scanner performance and accuracy of image-based measurements; is fully documented; has long-term monitoring and maintenance; and is traceably connected to the International System of Measurements. The need for a standard MRI system phantom was voiced at a National Institute of Standards and Technology (NIST) workshop entitled “Imaging as a Biomarker: Standards for Change Measurements in Therapy,” that was held on September 14-15, 2006.1 The workshop action items included: (1) Design phantoms that may better characterize the time-related physical performance of imaging systems, and the performance of specific functional and molecular-based measurements; (2) Define the physical performance of different imaging platforms required to measure change analysis; and (3) Develop and share open source tools to analyze phantom or simulated data.

The Ad Hoc Committee for Standards in Quantitative Magnetic Resonance (SQMR) was formed, under the auspices of the International Society of Magnetic Resonance in Medicine (ISMRM), to address these action items in the areas that pertained to MRI. The committee developed recommendations for a system phantom2 that could be used to determine the accuracy, stability, and comparability of MRI scanners. The desired measurements include:

1. Radio frequency (RF) field, $B_1$, non-uniformity
2. Static main magnetic field, $B_0$, non-uniformity
3. Geometric linearity
4. Gradient amplitude
5. Slice position and profile
6. Image uniformity
7. Resolution (high-contrast detectability)
8. Signal-to-noise ratio (SNR) (low-contrast detectability)
9. Accuracy and precision of measurement of proton spin relaxation times: $T_1$ (longitudinal) and $T_2$ (transverse) and proton density
10. System constancy

Such a system phantom can be used to: (1) track scanner performance at a particular site over time, as well as compare performance with other scanners; (2) determine the accuracy of certain quantitative measurements, such as $T_1$ mapping, and assist in the development of appropriate imaging protocols to obtain desired accuracy; (3) validate scanner and protocols for participation in clinical trials; and (4) determine the best phantom fabrication techniques for future phantoms, for example, application-specific phantoms.

Previous successful efforts to create standard phantoms include the American College of Radiology (ACR) MRI phantom3 and the Alzheimer’s Disease Neuroimaging Initiative (ADNI) MRI phantom.4 The ACR magnetic resonance accreditation phantom is a water-filled cylindrical phantom with relaxation times and conductivities in the biological range of interest. The phantom contains a resolution inset, slice profile wedges, a grid to determine geometric distortion, and a contrast array. The phantom was primarily designed as a method to assist site accreditation performed by the ACR and as a mechanism to improve quality control by repeated imaging on a weekly basis. The spherical ADNI phantom
is used for measurements of SNR, contrast-to-noise ratio (CNR), and geometric distortion in scan protocols intended to monitor changes in the brain morphology over time. From 2004 to 2009, the ADNI study acquired a phantom image with each participant study.4 The key results from this phantom plus human-participant study were: (1) the drift in scanner gradient calibration over months to years was less than the magnitude of step-wise changes in the gradient calibration introduced by field service calibration; (2) using phantom derived linear scaling corrections was not better than affine registration at correcting within-subject longitudinal-imaging-series gradient calibration changes or gradient drift. For ADNI-2 (2010-2017), phantom images were acquired for site certification and after system upgrades. These images were used to ensure that off-line gradient non-linearity corrections based on scanner model could be performed correctly.5 In ADNI-3 (2018-present), all scanners in the study have full three-dimensional (3D) on-board corrections for gradient non-linearity, and sites are requested to use ADNI phantoms for certification if possible. The standard system phantom described here builds off the ACR and ADNI phantoms, addressing the need for a more general and precise calibration object that has properties traceable to primary standards connected to the International System of Units (SI).2

This paper describes the design, construction, accuracy requirements, and measurement protocols of prototype system phantoms designed by the ISMRM SQMR ad hoc committee. The system phantom is a precise imaging artifact for quantitatively measuring image distortions and validating many types of quantitative mapping. The system phantom has subsequently been commercialized with over 100 units sold.6 The commercial system phantom has some minor changes in its design to accommodate more efficient manufacture, reduce cost, and incorporate some improvements that have been requested by customers. These changes are documented in Supporting Information Sec. 9. Here, we note that the spin-relaxation times and geometric tolerances of the commercial phantoms vary slightly from the prototypes and calibration data for the particular phantom, as determined from the serial number, should be used. The 3D mechanical design files, analysis software, analysis instructions, and imaging data can be found online, as described in the Data Availability Statement.

A single phantom cannot provide all the calibration measurements required for assessing the accuracy of MRI scanners and quantitative imaging protocols. However, this system phantom can address many of the important measurement and quality control issues. A universally used phantom is essential to allow sites to measure system stability and readily compare scanner performance with other scanners. There are several elements of the MRI system phantom described here that make it unique:

1. SI traceable parameters including geometric dimensions (mm), nuclear magnetic resonance (NMR) parameters such as proton spin relaxation times (s), and material compositions that dictate contrast properties.
2. Dimensionally stable construction using a rigid precision-joined framework and materials that are both low thermal-expansion and low water-uptake.
3. All components including design, 3D models, imaging protocols, detailed description of construction materials, and imaging analysis routines are in the public domain.
4. Long-term maintenance/monitoring of stability and accuracy by a national metrology institute.

2 | METHODS

This section gives an overview of the design and construction of prototype MRI system phantoms and the process to provide SI-traceable calibration of phantom properties. Two prototype system phantoms were built for this study. A more comprehensive description of the construction, solutions, and materials can be found in the Supporting Information. The design and construction specifications are meant as a common reference for commercial versions, not as specific requirements. The prototype phantoms are modular so that all components may be disassembled and reassembled in a variety of configurations for testing alternative designs.

2.1 | System phantom design

The prototype MRI system phantom, shown in Figure 1, consists of a water-filled spherical polycarbonate shell with a 200 mm inner diameter (ID). The spherical diameter was chosen to mimic a human head and was designed to fit in most head coil assemblies. A spherical design also allows for easy rotation of the phantom. Inside the spherical shell is a framework consisting of five 8.0 mm-thick polyphenylene sulfide (PPS) plates rigidly connected with PPS rods and kinematic mounts. PPS was chosen since it is a high-performance plastic with low water absorption, low thermal expansion, and good machinability. The PPS plates are annealed for stress relief and then machined to an inspected flatness of less than 0.04 mm. The plates support 57 fiducial spheres, a 14 element NiCl2 array, a 14 element MnCl2 array, a 14 element proton density array, two resolution insets, and wedges for slice profiles. The plates, as shown in Figure 1, contain the following structures: Plate 1, 5 fiducial spheres on bottom; Plate 2, 13 fiducial spheres on top and slice profile wedges on bottom; Plate 3, 21 fiducial spheres on top, proton density array on bottom; Plate 4, coarse resolution inset in the plate, 13 fiducial spheres on top, MnCl2 array on bottom; Plate 5,
The phantom coordinate system was chosen to match typical scanner/patient coordinates, with +Z along the scanner bore axis (superior), +X toward the left, and +Y down (posterior), when mounting the phantom in the default orientation. A set of eye decals is included on the phantom shell to allow easy recognition and placement of the phantom within the scanner, as seen in Figure 1B. The phantom is placed in the same fashion as a patient laying supine on the imaging bed. The origin of the phantom is defined as the center of the central fiducial sphere as shown in Figure 1C. The phantom is designed so that the plates are in the XZ (coronal) plane with left (L), right (R), superior (S), and inferior (I) marked on each plate, along with the plate identifier. Phantom identification information, along with the serial number, are etched into plate 5, which are easily readable in the prescribed 3D scan (Figure 1A). In the default orientation, the MR parameter arrays (NiCl₂, MnCl₂, proton density arrays) are imaged with a coronal scan. Recommended protocols for measuring the arrays and insets on clinical MRI systems, for the main MRI vendors, are given in the Supporting Information, Tables S4 through S10.

The inner frame of the prototype phantoms was designed to be very rigid and allow the fiducial-sphere centers to be located within a ±0.1 mm accuracy. Details of the plate locating mechanism are shown in Supporting Information Figure S1. A hemispherical machined surface on the post mates with a set of inclined plates to precisely locate the plates and eliminate torsional bending. The precision construction of the prototypes comes at an increased cost due to the need for computer-controlled machining. It was deemed necessary for the prototypes to have such precision to allow the study of scanner geometric distortion.

2.2 | Fiducial array

The system phantom contains a 3D fiducial array that can be used to assess geometric distortion, image uniformity and $B_1$ homogeneity. The fiducial array was modeled on the ADNI phantom,⁴ which has successfully been used to correct geometric distortions and image nonuniformity. The fiducial...
array spheres all have the same fill solutions with the same $T_1$ and $T_2$ properties ($T_1 = 407 \text{ ms} \pm 6 \text{ ms}, T_2 = 347 \text{ ms} \pm 6 \text{ ms} \text{ at } 20.0^\circ\text{C}, 3.0 \text{ T}$), which means they can be used to assess image homogeneity and calculate $B_1$ variation, both transmit, $B_1^+$, and receive, $B_1^-$, fields.

The fiducial array consists of a set of 57 precision-machined polyvinyl chloride (PVC) 10.0 mm ± 0.1 mm inner-diameter spheres on a 3D cubic lattice, with a lattice spacing of 40 mm ± 0.1 mm. Figure 2 shows a coronal MRI with a subset of the fiducial spheres along with 10 mm diameter circular regions of interest (ROIs) at their prescribed positions, before automated sphere location. Figure 2 shows standard output from the Python-based analysis package, after sphere location, plotting image uniformity and geometric distortion along the $x$, $y$, $z$ axes, respectively. The fiducial spheres are numbered 1 through 57, starting at plate 1, increasing from right to left, then inferior to superior. For data presentation purposes, the 57 fiducial array elements are divided into seven groups: 27 internal spheres on central $3 \times 3 \times 3$ grid, 5 spheres at each of the six outward faces of the phantom. Each set of fiducials is represented by a different color in the plots in Figure 2.

The spheres are filled with a CuSO$_4$ solution made by mixing 0.802 g CuSO$_4$ $\cdot$ 5H$_2$O per liter of deionized water (Supporting Information Sec. 4). The Cu$^{+2}$ ion concentration was measured, using an inductively coupled plasma-optical emission spectroscopy and a NIST traceable Cu standard (such as NIST standard reference material 3114) to be $0.18901 \pm 0.00055 \text{ mg/g}$ or $2.969 \pm 0.008 \text{ mM}$. A small amount of blue dye (erioflavine dihydrochloride, 0.12 µg/ml)

---

**Figure 2**

A. Coronal 3D gradient echo image of the fiducial array along with 10.0 mm circular ROIs located on prescribed 40 mm grid, which allows for visual identification of geometric distortion. S, I, L, R refer to superior, inferior, left, right directions, respectively. The scan is at 3T using a body transmit coil and a head receive coil with 0.97 mm isotropic voxels, TR = 6.3 ms, TE = 1.89 ms. The nonuniform gradient corrections are turned off. B. Normalized integrated intensity of all 57 spheres obtained after the sphere location procedure. C,D,E. Geometric distortion along the $x$, $y$, $z$ directions, respectively. Geometric distortion, $\delta\vec{R} = \vec{R}_a - \vec{R}_p$, is defined as the distance between the apparent position $\vec{R}_a$ and the prescribed position $\vec{R}_p$ of the sphere centers after an affine transformation has accounted for rotation, translation, and overall scale correction.
was added to assist in the visual recognition of the fiducial elements. The fiducial spheres are machined in two parts from PVC stock, glued together and attached to the plates using an M8-1.25 threaded stud, as shown in Supporting Information Figure S1D. PVC was chosen because it is inexpensive, can be easily glued, and has a low permeability to water. The spheres are sealed with a tapered polyvinylidene fluoride (PVDF) plug. The fiducial solution was chosen to have short $T_1$ times to give, for most scans in the protocol, high signal and contrast with respect to the surrounding fill composed of long-$T_1$ deionized water. High contrast is beneficial for automated location of the fiducial sphere positions.

### 2.3 MR parameter arrays (NiCl$_2$, MnCl$_2$, and proton density arrays)

The MR parameter array spheres are 20 mm outer diameter (OD), 15 mm internal diameter (ID) polypropylene spheres that are filled, heat sealed and then, epoxy sealed to an M8-1.25 threaded mounting stud (Supporting Information Figure S1D), similar to those used for the fiducial spheres. Since the spheres are epoxied onto a hemispherical indent, as opposed to being integrally machined with the threaded stud, there is more positional uncertainty in the MR parameter arrays than in the fiducial array. Polypropylene spheres were chosen since they are inexpensive, commercially available, and have one of the lowest water/air permeation rates among all of the plastics tested. However, the OD, ID, and sphericity of the polypropylene spheres are not well controlled: see Figure S2D inset. The polypropylene spheres are made by thermally welding together two molded hemispheres, which results in a thicker wall in the weld plane.

The $T_1$ and $T_2$ values were modified by doping high-purity water with NiCl$_2$ and MnCl$_2$ in their respective arrays (Supporting Information Sec. 4). The proton density values were modified by making solutions of 5 % to 100 % high-purity water with the balance D$_2$O. Proton density is defined as the concentration of MR-visible protons in a sample/tissue, relative to that in the same volume of water at the same temperature. All solutions use American Chemical Society (ACS) reagent grade, or higher quality, water. A small quantity of NiCl$_2$ (0.26 mg Ni$^{2+}$/mL) was added to the proton density solutions to reduce the relaxation times. Green dye (28 µg/mL erioglaucine disodium salt, 28 µg/mL tartrazine) was added to the NiCl$_2$-doped array solutions, red dye (49.6 µg/mL allure red AC) was added to the MnCl$_2$-doped array solutions, and yellow dye (53 µg/mL tartrazine) was added to the proton density solutions to make the arrays easily identifiable by the user. Each array consists of 14 spheres, 10 equally spaced on a 50 mm radius, and 4 spheres internal to the outside 10 spheres on a 40 mm grid. The sphere positions, NiCl$_2$, MnCl$_2$, D$_2$O concentrations, and measured $T_1$, $T_2$ values (at 20°C, 1.5 T and 3.0 T) for the NiCl$_2$, MnCl$_2$, and proton density arrays are listed in the Supporting Information in Tables S1-S3, respectively.

The NiCl$_2$ and MnCl$_2$ arrays were chosen to have a wide range of values to mimic both endogenous tissues and tissue containing standard commercial contrast agents. $T_1$ and $T_2$ relaxation times for the arrays, along with typical values for tissue, are shown in Figure 3. The MnCl$_2$-doped array spans a range of relaxation time values much closer to that of tissue; however, it has considerably more temperature and field dependence than the NiCl$_2$-doped array (Supporting Information Sec. 7). $T_1$ and $T_2$ can be measured on both the NiCl$_2$-doped and MnCl$_2$-doped arrays; although historically, the NiCl$_2$ array was designed to cover a suitable range of $T_1$ values, while the MnCl$_2$ array was chosen to cover a suitable range of $T_2$ values. Also shown in Figure 3 are model fits to the data, assuming that the relaxation rates are linearly proportional to the paramagnetic salt concentration. The deviation of the measured data from the model and the relation to the measurement uncertainties are discussed in the Supporting Information Sec 2.

The NiCl$_2$ doping concentrations were chosen to give a $\sqrt{2}$ progression of $T_1$ values from approximately 20 ms to 2000 ms at a $B_0$ field of 1.5 T. NiCl$_2$ was chosen because the relaxivity of paramagnetic spin-1 Ni$^{2+}$ ions on water proton spins has been extensively studied. $T_1$ values are relatively insensitive to the magnetic field strength and to temperature over the range of application for this phantom. However, the paramagnetic relaxation enhancement due to...
Ni\textsuperscript{2+} has a complex behavior in the region of application with non-monotonic variation of the relaxivities with field and temperature.\textsuperscript{12} $T_1$ and $T_2$ for the higher concentration NiCl\textsubscript{2} solutions, Ni-5 to Ni-14, have a minimum near 22°C at 3 T (Supporting Information Sec 7), close to typical MRI bore temperatures. The presence of a minimum eliminates the large linear variation of $T_1$ and $T_2$ with temperature, giving a maximum variation of 4% over the range of expected bore temperatures (16°C to 26°C).

The MnCl\textsubscript{2} concentrations are varied from 0.013 mM to 1.704 mM and, as with the NiCl\textsubscript{2} array, the concentrations were spaced by factors of $\sqrt{2}$. The variation of $T_1$ and $T_2$ is a linear function of temperature over the expected range of bore temperatures (16°C to 26°C), with a variation at 3 T of 2.7%/°C and 1.6%/°C for $T_1$ and $T_2$, respectively.

### 2.4 Resolution and slice profile insets

The resolution insets are modeled on the ACR guidance,\textsuperscript{13} and the slice profile wedges are designed in accordance with National Electrical Manufacturers Association (NEMA) MS 5-2018 standard.\textsuperscript{14} Photographs and MR images of resolution and slice profile insets are shown in Figure 4.

The phantom contains two resolution insets consisting of $4 \times 4$ arrays of holes. A coarse resolution inset is drilled directly into plate 4 with hole diameters of 0.6 mm, 0.7 mm, 0.8 mm, 0.9 mm, and 1.0 mm ± 0.1 mm diameters. The fine resolution inset is contained within a rectangular prism filled with CuSO\textsubscript{4} solution to give high contrast and has hole diameters of 0.4 mm, 0.5 mm, 0.6 mm, 0.7 mm, and 0.8 mm ± 0.1 mm diameters. Each hole array is duplicated at a 10° angle, sharing one common hole, to assess resolution along non-principal axes. In total, there are 155 holes in each of the resolution insets (5 hole sizes, 31 per size). The advantage of this resolution inset is that it is easy to interpret visually, compact, easy to manufacture, and there is considerable community experience with it.

The slice profile inset consists of two 10° angle wedges contained within a rectangular prism filled with CuSO\textsubscript{4} solution. The wedges are oriented so that one has a positive and one has a negative inclination. The presence of a pair of wedges allows for the slice profile to be corrected for the misalignment of the slice plane relative to the base plane of the wedges.

### 2.5 Safety

For most of the imaging studies, the shell was filled with 4 L of deionized water. This was done for two reasons: (1) safety, if the phantom spilled, the outer fill has no toxicity or

---

**FIGURE 4** A,B, Photo and MRI of resolution inset. C,D, Photo and MRI of slice profile inset.
disposal issues; and (2) practicality, the phantom could be easily refilled at any site. The main toxicity issue is due to the NiCl₂ solutions. A total of 52 mg Ni²⁺ is included in the phantom. The NiCl₂ is contained in the thick-walled polypropylene contrast spheres that are double sealed with both a weld and epoxy. These spheres are very robust and can take large impacts without breakage or leakage.

2.6 | SI traceability

A calibration object requires its properties to be precisely measured and traceable to a common set of definitions and units. The measured values must include a well-defined uncertainty. A protocol for traceable spin relaxation measurements for MRI phantoms can be found in NIST SP250-9715 and measurement detail can be found in Supporting Information Sec. 6. Spin relaxation time measurements are SI traceable via a metrology NMR system with a calibrated time base. The NMR time base, a 10 MHz oven-controlled crystal oscillator, is calibrated against the NIST hydrogen maser clock. The NMR time base errors, which include jitter of transmit and receive events, give minor contributions to the overall uncertainty in relaxation time measurements.

More important factors that determine the uncertainty in relaxation time measurements are RF transmit field $B_1^+$ calibration, $B_1^+$ inhomogeneity, $B_0$ inhomogeneity, and measurement temperature. $B_1^+$ is calibrated using a nutation experiment on the sample to be calibrated, where the spins are progressively tipped, by varying the RF pulse duration, at increasing angles through at least 2 cycles (720°). Free induction decays are recorded, Fourier transformed to obtain a spectral peak, integrated and plotted against RF pulse duration. The data, which approximate a damped sinusoid, are fit to extract the pulse duration required for the desired tip angle and to extract an RF field inhomogeneity in the system. The RF field inhomogeneity is minimized during calibration by precisely centering a 10 mm long test sample within a 14 mm long RF homogeneous zone. This field inhomogeneity and uncertainty in the $B_1^+$ calibration are fed into Monte Carlo Bloch simulations, along with all other uncertainties, to calculate the uncertainties in the measured relaxation times.

$B_0$ inhomogeneity in the NMR metrology system is minimized by performing an automated shimming procedure on the sample to be calibrated. The inhomogeneous line width is required to be no more than 10 Hz larger than the homogeneous line width. Monte Carlo Bloch simulations indicate that this level of $B_0$ inhomogeneity gives a negligible contribution to the overall uncertainty in relaxation times with the pulse sequences used (inversion recovery sequence with composite 180° pulses for $T_1$ and Carr-Purcell-Meiboom-Gill (CPMG) sequence with 2.0 ms refocusing times for $T_2$).

Temperature measurement and control are a major source of uncertainty both in the primary calibrations and in phantom-based MRI scanner assessment. Phantom components can have several percent change in relaxation time per °C, Supporting Information Sec. 9. For primary calibration in the NMR metrology system, sample capillaries are immersed in a fluorinated heat transfer solution within a 5 mm diameter NMR tube. The fluorinated solution also serves as a susceptibility matching solution. A fiber optic thermometer is positioned 15 mm away from the sample and is calibrated in the same NMR tube in a water bath against two NIST-calibrated platinum resistance thermometers.

Geometric accuracy is determined at manufacture time using computer-controlled machining techniques. Post-assembly geometric accuracy can be checked with computed tomography techniques using SI traceable geometric calibrations. Here, the geometric distortion is reported relative to the 3D model values, with uncertainty given by the manufacturing tolerances.

Composition of the MR parameter arrays is verified using inductively coupled plasma- mass or optical spectroscopy techniques calibrated with traceable NIST reference standards. A reference library of all solutions, flame sealed in capillaries, is kept to establish stability over time. A test of the stability of proton relaxation times of an archived sample of Ni-12 solution at 3 T and 20.0°C, for example, gave $T_1 = 44.53 \text{ ms} ± 0.01 \text{ ms}$, $T_2 = 31.86 \text{ ms} ± 0.03 \text{ ms}$ in May 2015 and $T_1 = 44.67 \text{ ms} ± 0.003 \text{ ms}$, $T_2 = 31.97 \text{ ms} ± 0.026 \text{ ms}$ in June 2019, a difference of 0.3 % for $T_1$ and 0.3 % for $T_2$. Here, the reported errors are the SD of three consecutive measurements at each point in time and is a measure of system stability and is not the uncertainty in the values, which must incorporate all systematic errors as described in NIST SP250-97. Uncertainties will depend on the material under test, the precise pulse sequence used, the geometry of the sample, and on the values of the relaxation times. The uncertainty for proton spin relaxation time calibrations, defined as the range within which there is greater than 96 % probability of the true value, are given in the NIST calibration certificates for each material and are on the order of 1.5 %.

3 | RESULTS

3.1 | Geometric distortion, image uniformity, and B₁ homogeneity

Geometric distortion is assessed in a manner similar to that described in Ref. 4. A 3D gradient echo image, with short echo time (TE), of the whole phantom is obtained with isotropic voxels with typical voxel dimension of 1.0 mm. This sequence can be used to assess the intrinsic geometric accuracy of the scanner, whereas sequences with longer TEs will
the apparent locations of the 10.0 mm fiducial spheres are
determined by cross-correlation maximums between a fidu-
cial sphere mask and the image (discussed in the Supporting
Information). The sphere center can be determined to <0.1
mm, which is comparable to the accuracy in the construc-
tion. Determination of the center of the fiducial spheres is
sensitive to defects, such as small air bubbles in the solu-
tion. These defects can be detected manually or numerically,
and the defective sphere data discarded. After locating the
apparent sphere centers, a similarity transformation (an af-
fine transformation consisting of a translation, rotation, and
isotropic scale factor), which minimizes the distances be-
tween the prescribed and apparent locations is determined
by a generalized Procrustes algorithm. The resulting rotation
and translation define the map from phantom to scanner co-
ordinates. The scale factor provides a measure of the overall
dimensional accuracy of the scanner. We define geometric
distortion as the difference between the apparent position
$\vec{R}_a$ and the prescribed position $\vec{R}_p$ of the sphere centers after
the similarity transformation has been applied to the latter,
$\delta \vec{R} = \vec{R}_a - \vec{R}_p$. An example geometric distortion analysis for
an image without corrections for gradient non-uniformity,
is shown in Figure 2. A large (~2 mm) $x$-component of

![Figure 5]

**FIGURE 5** A, Coronal image of the NiCl$_2$ array at 3 T ~20°C using an IR protocol. B, Average signal in ROIs 1, 5, 9, and 13 in the NiCl$_2$
array, plotted as a function of TI and fits to the inversion recovery model. C, Automated analysis of $T_1$ for each of the 14 NiCl$_2$ array elements,
with the deviation from the NMR reference measurements. The analysis was done both by (1) fitting the average of the ROI signal to the model
(blue squares) and (2) fitting each voxel independently to the model, making a $T_1$ map, and then averaging (red squares). The deviations are shown
in blue and red squares, respectively. The error bars on the deviations are the standard errors in the nonlinear least squares fits and represent the
quality of the model and the fit. D, The inversion efficiency, $\delta$ determined from the fits. The line at $\delta = 1$ represents perfect inversion, and the
error bars are the standard errors in the parameter determined from the nonlinear least squares fits. Unobserved error bars are contained within the
symbol.
geometric distortion is seen on the right face, with a negative sign, in Figure 2 since the apparent position is farther along the negative x-axis than the prescribed position. Similarly, there is a large x-component of geometric distortion on the left face, with a positive sign, since the apparent position is farther along the positive x-axis. Examples of geometric distortion with non-uniform gradient corrections applied are given in the Supporting Information and show the need for the phantom geometric accuracy to be < 0.1 mm. Local volumetric distortion can be assessed by comparing cross correlation volumes.

We assess the uniformity of image intensity as part of this same analyses by calculating the integrated intensity of the 57 fiducial spheres. Figure 2 shows image uniformity obtained from a head coil, which shows signal loss in the inferior or chin region due to low receive coil sensitivity. The image uniformity analysis can also be done as part of other protocols, such as the of $T_1$, $T_2$ protocols.

Finally, $B_1^+$ mapping can be done with the system phantom fiducial array using flip angle mapping techniques.\textsuperscript{16-18} The fiducial spheres, as well as the surrounding background fill liquid (DI water), provide regions with identical properties to allow identification of RF non-uniformities. To assess the effect of fill conductivity and dielectric resonances, the fill solution can readily be swapped out with a conducting salt solution or a low dielectric constant fluorocarbon fluid. We do not perform this analysis here.

### 3.2 $T_1$ measurement

Figure 5 shows automated $T_1$ analysis of the NiCl$_2$ array using the Python-based analysis package. Figure 5A shows an image of the NiCl$_2$ array using the inversion-recovery (IR) sequence protocol described in the Table S8 with recommended 10.0 mm circular ROIs centered on the NiCl$_2$ array spheres. The suggested $T_1$-IR protocol requires 10 inversion times, TI, ranging from 50 ms to 3000 ms and TR = 4500 ms. Note, the actual inversion recovery protocol may vary slightly among the different vendors due to details of the scanner design and programmable control. Figure 5B shows a typical fit to the data using a standard IR model,\textsuperscript{19-21} appropriate when TR >> $T_1$, and the inversion and detection flip angles are precisely 180° and 90°. The average signal in each ROI is given by $S(T) = A \left\{ 1 - (1 + \delta) e^{-\frac{t}{T_1}} \right\}$, where A is the signal amplitude when the magnetization is allowed to fully relax back to its equilibrium state, and $\delta$ is the inversion efficiency.\textsuperscript{19,21,22} The signal amplitude immediately following inversion normalized to A. Here, only magnitude data are used since the phase data are often not available and can be hard to interpret. This is in contrast with the NMR IR reference protocol, which uses the real part of the complex signal for analysis. Figure 5C shows the results of the $T_1$ analysis along with the deviation relative to the NMR reference values given in Table S1. The deviation is defined as $100(T_{1\text{meas}} - T_{1\text{ref}})/T_{1\text{ref}}$, where $T_{1\text{meas}}$, $T_{1\text{ref}}$ are the measured and reference values, respectively. The large deviation for sphere Ni-14, which has the shortest $T_1$, results from having only three data points where the signal is changing before full recovery. This is reflected in the large error bar, which is the standard error in the nonlinear least squares fit. The analysis was done both by (1) fitting the average of the ROI signal to the model and (2) fitting each voxel independently to the model, making a $T_1$ map, and then averaging. Both methods agree within the standard errors. A series of $T_1$ measurements made on 3 T scanners over the course of 7 years is shown in the Figure S13 to illustrate phantom stability.

More information is obtained from these fits than simply the relaxation time. The inversion efficiency, $\delta$, is an important indicator of $B_1$ and flip angle uniformity. As seen in Figure 5D, the inversion efficiency is close to 0.85, whereas the ideal value is 1.0. Typical values obtained from NMR data on the same solutions, with a more homogenous RF field geometry and using composite inversion pulses, are between 0.97 and 1.0. A Monte Carlo study of Bloch simulations indicate that there is a correlation between the deviation of the inversion efficiency from its ideal value and the error in the $T_1$ measurements.\textsuperscript{15}

IR sequences, while fairly accurate in measuring $T_1$, require long acquisition times. Variable flip angle (VFA) sequences, with shorter TRs and smaller flip angles, are considerably faster with less RF deposition. VFA protocols typically use 3D gradient echo sequences (see Supporting Information) and vary the flip angle $\alpha$. Ideally, the signal is given by $S(\alpha) = S_{90}\sin(\alpha) \frac{1 - e^{-\frac{\alpha}{T_1}}}{1 - e^{-\frac{TR}{T_1}\cos(\alpha)}}$, where $S_{90}$ is the maximum signal given $\alpha = 90^\circ$ and long $TR$.\textsuperscript{20,21} Figure 6A shows a coronal image of the NiCl$_2$ array using the VFA protocol described in the Supporting Information, with $TR = 5.37$ ms and TE = 1.49 ms. Figure 6C shows a series of sagittal images for a series of flip angles varying from 2° to 30°. The signal in each ROI has a maximum at the Ernst angle, $\cos^{-1}(\frac{\alpha}{TR}) = e^{-\frac{\alpha}{T_1}}$, which for ROI 5 is near 10° and ROI 2 is near 5°. Figure 6B shows the mean signal for ROIs 2, 5, 8, 11 along with fits to the model listed above and the residuals. As seen by the non-Gaussian distribution of the residuals, there are systematic deviations from the simple model listed above due, in large part, to the inability to achieve the prescribed flip angle at each location in the phantom. Figure 6D shows the measured $T_1$ values and deviations from the NMR reference values. The error bars in Figure 6D are the standard error in the fits, which are large for the longer $T_1$ ROIs. The large standard errors on the fits, along with a systematic variation in the residuals, indicate
that the model, which assumes an accurately prescribed flip angle that is the same for all regions in the phantom, is limiting measurement accuracy. More advanced models and protocols exist to improve on VFA $T_1$ measurements \(^\text{23}\); however, here we are focusing on using the system phantom and simple protocols to identify and characterize scanner non-idealities.

### 3.3 $T_2$ measurement

Figure 7 shows $T_2$ measurements of the MnCl$_2$ array using the Python-based analysis package. Figure 7A shows an image of the MnCl$_2$ array using the spin echo (SE) sequence listed in the Supporting Information with TE varying from 10 ms to 320 ms by 10 ms and TR = 5000 ms. Figure 7B shows a typical fit to the data using a simple exponential model: $S(TE) = S_0 e^{-TE/T_2}$, where $S(TE)$ is average signal in each ROI and TE is the spin TE. Figure 7C shows the $T_2$ obtained from the fits along with the deviation relative to the NMR reference values given in Table S2. The error bars in Figure 7C are the standard errors derived from the non-linear least squares fitting procedure. For the ROIs with short $T_2$ times, there are not many points used in the fit, and the standard error is large.

### 3.4 Proton density and SNR measurement

A coronal spin echo image of the PD array is shown in Figure 8A with the PD ROIs shown in yellow. Substantial background nonuniformity exists; the signal changes by a factor of ~6 going from the periphery to the center along the superior/inferior direction. Figure 8B shows the average signal in each PD ROI, and the measured MR proton density plotted versus prescribed proton density. The MR proton density is calculated by normalizing the average signal to the local background, as determined by averaging the signal from four points taken 3 mm outside the spheres, and then normalizing to the 100 % water sphere. The average signal
in the PD ROIs shows a clear nonlinearity due to signal loss towards the periphery of the phantom. The normalized signal shows the expected linearity of the signal with respect to proton density. The maximum error in the measured proton density is 8.3%, and the SD of the errors is 2.8%.

In addition to measuring proton density, this array is used to measure SNR using methods similar to those described in Ref. 24. The noise is measured by taking the difference of two identical scans, \( \Delta S = S_2 - S_1 \), taken sequentially (Figure 8C), as discussed in Method 1 in Ref. 24. The SNR is calculated for each PD ROI by taking the average signal \( S_{av} = \frac{1}{2} (S_{2av} + S_{1av}) \) and dividing by the noise \( N = SD/\sqrt{2} \), where SD is the SD of the difference signal \( \Delta S \) within the ROIs. The SNR is plotted versus proton density in Figure 8D. Also shown is the average signal in each ROI normalized to the SD within each ROI. This is a measure of the ratio of the average signal to the nonuniformity within each region, only part of which comes from noise.

SNR1 ROI, shown in Figure 8 in blue, is meant to more closely mimic the geometry suggested in Ref. 24. However, the larger ROIs are more susceptible to scanner and phantom temporal drift, which gives rise to the low spatial frequency structure in Figure 8C.

3.5 | Resolution

Resolution is determined either using the manual method suggested by ACR\(^{13}\) or an automated computer algorithm that compares the MR image to synthesized images with different resolution distortions. Figure 9A shows an image of the resolution inset taken with a voxel size of 0.35 mm and a slice thickness of 4.0 mm. As per ACR guidelines, visual inspection of the image yields a resolution of 0.5 mm. Figure 9B shows a synthetic image generated by analytically computing the Fourier transform of the resolution-inset disk array, calculating a finite k-space image, and converting into a real-space image. Comparison of the synthetic image to the measured image shows additional blurring beyond that expected from the point spread function due to finite k-space sampling. Applying an additional Gaussian point spread function can improve the match between the synthetic and real image.
3.6 | Slice profile measurement

Slice profile measurements, shown in Figure 10, are done in accordance with the NEMA MS-5-2018 standard\textsuperscript{14}: the edge response function is calculated from the imaging data by placing a 4 mm x 50 mm rectangular ROI over each wedge, differentiating the data with respect to image position, and then multiplying the image position by \( \tan(\theta) \) to determine the slice profile. \( \alpha = 10^\circ + \theta \) is the corrected angle, where 
\[
\theta = \frac{1}{2} \sin^{-1} \left( \frac{(w_2 - w_1) \sin(2 \times 10^\circ)}{w_2 + w_1} \right)
\]
measures the angle of the image plane with respect to the base of the wedges (see Figure 10C inset), and \( w_1, w_2 \) are the projected thicknesses of the two wedges. The slice thickness, \( t_{sl} \), is then determined by measuring the full width at half maximum of the corrected slice profile. The suggested prescribed slice thicknesses are 3 mm and 5 mm, with the maximum being set by the wedge height of 10 mm.

The slice thickness may also be measured by fitting the edge response function with a piecewise linear function, extracting (Figure 9), with the difference between the images minimized using a point spread function with a width \( 2\sigma = 0.075 \) mm, as seen in Figure 9D. The resolution is then reported as a finite k-space sample limited resolution of 0.35 mm and a machine-specific Gaussian broadening of 0.075 mm.

(Figure 8) A. Coronal image of proton density array using a spin-echo sequence with TR = 5000 ms, TE = 10 ms. A background SNR ROI is shown in blue, in addition to the 14 yellow proton density ROIs. Line scans across the center background fill are shown in yellow and orange at the bottom to show the image inhomogeneity. B. Average intensity of PD ROIs and measured PD as a function of prescribed proton density. C. Difference of two identical images and D, plot of the SNR as a function of proton density. Also shown in D, is the ratio of the signal to the SD within each ROI.
the projected slice thicknesses, and then multiplying by the tangent of the wedge angle: 

$$t_{sl} = \frac{1}{2} (w_2 + w_1) \times \tan(10^\circ).$$

Averaging both ramp orientations corrects, in lowest order, for errors due to the misorientation of the image slice plane relative to the base of the wedges. The latter method has the advantage that it is easily automated and has a well-defined uncertainty.

4 | DISCUSSION

The system phantom described here, or modified versions of it, have been used for several studies including repeatability of MR fingerprinting, assessing changes occurring during scanner upgrades, and assessing variation in multi-site $T_1$ measurements. Many other uses are ongoing, including monitoring of scanners for quality assurance and homogenizing scan protocols across multi-vendor scanners. The phantom was designed to be used in several modes, starting from simple visual inspection for identification of geometric distortion and resolution, to semi-automated analysis to determine relaxation times and SNR, and finally for custom analysis to validate advanced quantitative protocols. While application-specific phantoms may provide better structures for particular measurements, for example, the ADNI phantom for geometric distortion or a uniform agar phantom for image homogeneity, the system phantom is meant for characterization of a large number of scanner parameters and to have sufficient accuracy, generality, and complexity to assess the accuracy of many types of advanced hardware and protocols. The standard protocols provided with the system phantom are not intended to be the best protocols for accurate measurement of geometric distortion, proton spin relaxation times, resolution, and SNR. Rather, they are meant as a common method to test the ideality of the scanner and to provide a comparison for other quantitative protocols that may be faster or more accurate.

Here we have shown how the phantom can be used for scanner diagnostics assessing geometric distortion, image homogeneity, resolution, and slice profile. The resulting data can provide both parameters to benchmark scanner performance and insight on what is determining these parameters. For example, resolution analysis can separate the pulse-sequence component from the intrinsic scanner component of the point spread function. The SNR protocol can separate temporal and spatial noise components, which contribute in different ways to SNR. Finally, the MR-parameter arrays provide a very broad range of properties to allow validation of both standard and advanced quantitative protocols.

One specific challenge encountered is that $T_2$ is not a rigorously defined quantity. $1/T_2$ is the component of the total transverse spin dephasing rate, $1/T^*_2$, that is considered to be intrinsic to the sample, with the extrinsic component, $1/T_{2e}$, being eliminated using a spin echo: $1/T_2 = 1/T_{2e} + 1/T_{2e}$. For complex materials there is ambiguity on what dephasing components are considered extrinsic and intrinsic, and the components being eliminated by the refocusing pulses will depend on the details of the refocusing pulses. $T_2$ can be a function of the refocusing time and the values measured by CPMG are not the same as measured by SE. The system phantom uses simple materials, so this is not a major issue here. However, to get traceability, a robust operational definition of $T_2$ is required. Here, we define it as the exponential signal decay measured with a CPMG sequence with a refocusing time of 2 ms. CPMG sequences also effectively null out the longitudinal magnetization, which, if present along with $B_0$...
and $B_1$ inhomogeneities, can lead to stimulated echoes. Other imperfections such as imperfect refocusing pulses, which can be present in SE sequences, can add long-time signal and can increase the measured $T_2$. The variability in $T_2$ measurements have been extensively studied in the literature.\textsuperscript{31-33} Here, we note that simple SE sequences, which may appear to be similar, can have very different spin manipulations and signal production. Using a standard phantom can identify these differences, which can be important for both conventional qualitative interpretation as well as quantitative analysis.

From the feedback of users of the prototype phantoms, several limitations and potential improvements have been identified. During the time since this phantom was conceived, head coil geometries are becoming more complex, and the size is decreasing. Newer head coils have minimum dimensions of 196 mm or less, which is smaller than the prototype system phantom design. There is a trade-off in the system phantom between sampling large volumes to assess uniformity of measurements and the ability to fit into newer, smaller coils. Smaller phantom shells have been investigated and are available in commercial versions. Another concern is the cost of the phantom. Tradeoffs can be made, such as relaxing the 0.1 mm geometric accuracy, to reduce costs to make the phantoms commercially viable, as discussed in Supporting Information Sec. 13.

The contrast array components are arranged in an orderly fashion to allow easier interpretation by users. More information may potentially be obtained by random placement of the contrast elements, given that their properties are highly determined and errors due to location may be better extracted using computer algorithms. There could be benefit as well in reducing the number of unique spheres in the contrast array to provide replicate spheres. The prototype phantoms presented here are easily reconfigurable to test these modifications.

The contrast and fiducial arrays use plastic components that have sharp signal-to-no-signal interfaces that cause Gibb’s ringing artifacts, as can be seen in Figure 6A. These artifacts can confound SNR and other measurements. While these artifacts can be accounted for in certain measurements, for instance they can be included in the kernel for the cross-correlations determining geometric distortion, it would be desirable to minimize these artifacts by making the plastic shells thinner or by allowing them to have some signal.

Dielectric resonances can give rise to image distortion, particularly for large water phantoms at high fields.\textsuperscript{34} These effects still need to be studied for the system phantom. However, the outer fill is easily changeable, without affecting the measurement protocols. Fills with different conductivities and dielectric properties can easily be explored.

Temperature control and monitoring remain a critical issue, especially when trying to verify the accuracy of relaxation time measurements to better than 2 %. While the system phantom is shipped with a thermometer to record
temperature, the bore temperatures are often different from the ambient scanner room temperature or even the control room, where the temperature is typically measured. There is no guarantee that the phantom temperature is the same as the recorded temperature. Furthermore, the temperature information is not easily encoded in the DICOM files. This establishes the need for in-situ NMR thermometers, and an MRI-readable liquid crystal thermometer has been developed for MRI phantoms.\textsuperscript{35,36} Furthermore, the system phantom would greatly benefit from the use of materials that are less sensitive to variations in temperature and magnetic field strength. Most paramagnetic salts such as MnCl\textsubscript{2} and CuSO\textsubscript{4} have considerable temperature\textsuperscript{10} and field dependence (see Supporting Information Figures S10 and S11) in their relaxivity values r\textsubscript{1}, r\textsubscript{2} (mM\textsuperscript{−1} s\textsuperscript{−1}). NiCl\textsubscript{2}, while having less temperature and field dependence, has safety/toxicity issues.

The materials used in the system phantom were chosen for their stability and simple properties, for example, exhibiting mono-exponential spin relaxation. As shown in the Supporting Information, a reference library sample, sealed in a glass capillary, has 4-year \( T_1 \) and \( T_2 \) stability better than 0.1 \%. Assessing stability of the phantom components is more challenging, since sampling the solutions for SI-traceable NMR measurements would be destructive. The plastic spheres in the phantoms have some permeability to water and may slowly absorb paramagnetic salts, leading to long-term drift in \( T_1 \) and \( T_2 \). A sampling protocol of the MR-property spheres, stored under conditions similar to those contained within the phantom, is required to assess long term stability and is in progress.

There is demand for more biomimetic materials that match multiple tissue properties, such as the apparent water diffusion coefficient as well as proton density and proton spin relaxation times. It is challenging to obtain complex biomimetic materials that are stable and also have properties that can be rigorously quantified. However, this work is under way, as illustrated by the stable fat mimic developed for the commercial version of the breast phantom developed by NIST and the University of California San Francisco.\textsuperscript{37}

While the current implementation of the MRI system phantom can be improved and will never be suitable for all calibration needs, it is a first attempt at developing and implementing a rigorous calibration structure with traceable properties, community-driven open construction details and evaluation protocols, and long-term monitoring and assessment with state-of-the-art metrology tools.

ACKNOWLEDGMENTS

The authors thank Michael Snow and William Hollander at High Precision Devices for the computer aided design and manufacturing guidance. The authors also thank Elizabeth Mirowski from Verellium and Joshua Levy from Phantom Labs for useful discussions on phantom design and implementation. The authors also thank all the ISMRM SQMR committee members, past and present, for their advice and support. This paper is dedicated to Larry Clarke, NCI Branch Chief, who helped initiate and champion this work and many other quantitative imaging programs over the past 2 decades.

CONFLICT OF INTEREST

Todor Karaulanov is an employee of CaliberMRI.

DATA AVAILABILITY STATEMENT

The analysis software can be found at https://github.com/MRISt andards/PhantomViewer. An analysis manual, 3D models, and sample datasets can be found at https://github.com/MRISt andards/SystemPhantom.

ORCID

Karl F. Stupic https://orcid.org/0000-0001-8356-1660
Michael A. Boss https://orcid.org/0000-0002-9492-767X
Kathryn E. Keenan https://orcid.org/0000-0001-9070-5255
Chun Yuan https://orcid.org/0000-0002-6154-5106
Stephen E. Russek https://orcid.org/0000-0002-8788-2442

TWITTER

Karl F. Stupic @KarlStupic

REFERENCES

1. Clarke LP, Sriram RD, Schilling LB. Imaging as a biomarker: standards for change measurements in therapy workshop summary. J Academic Radiol. 2008;15:501-530.
2. Keenan KE, Ainslie M, Barker AJ, et al. Quantitative magnetic resonance imaging phantoms: a review and the need for a system phantom. Magn Reson Med. 2018;79:48-61.
3. Price R, Allison J, Clarke G, et al. 2015 MRI Quality Control Manual. Virginia: American College of Radiology; 2015.
4. Gunter JI, Bernstein MA, Borowski BJ, et al. Measurement of MRI scanner performance with the ADNI phantom. Med Phys. 2009;36:2193-2205.
5. ADNI2. Alzheimer’s Disease Neuroimaging Initiative 3T MRI Technical Procedures Manual. California: Alzheimer’s Disease Neuroimaging Initiative; 2011.
6. Premium System Phantom, CaliberMRI. Web site. https://qmri.com/qmri-solutions/t1-t2-pd-imaging-phantom/. Accessed August 1, 2020.
7. Tofts PS. PD: proton density of tissue water. Quantitative MRI of the Brain. 2004:83-103.
8. ACS. Water, Reagent. ACS Reagent Chemicals. 2017;4A.
9. ITIS. Tissue Properties Database V4.0. Foundation for Research on Information Technologies in Society (ITIS). Zurich: ITIS Foundation.
10. Kraft KA, Fatouros PP, Clarke GD, Kishore PRS. An MRI phantom material for quantitative relaxometry. Magn Reson Med. 1987;5:555-562.
11. Kowalewski J, Maler L. Nuclear Spin Relaxation in Liquids: Theory, Experiments, and Applications. Series in chemical physics. New York: Taylor & Francis; 2006.

12. Kowalewski J, Egrov A, Kruk D, et al. Extensive NMRD studies of Ni(II) salt solutions in water and water-glycerol mixtures. J Magn Reson. 2008;195:103-111.

13. ACR. Phantom Test Guidance for Use of the Large MRI Phantom for the ACR MRI Accreditation Program. Virginia: American College of Radiology. 2018.

14. National Electrical Manufacturers Association. NEMA Standards Publication MS 5–2018 Slice Thickness. Determination of Slice Thickness in Diagnostic Magnetic Resonance Imaging. New York: The Association; 2018.

15. Boss MA, Dienstfrey AM, Gimbutas Z, et al. Magnetic resonance imaging biomarker calibration service. Proton Spin Relaxation Times. 2018;Special Publication (NIST SP) - 250-97.

16. Lutti A, Hutton C, Finsterbusch J, Helms G, Weiskopf N. Optimization and validation of methods for mapping of the radio-frequency transmit field at 3T. Magn Reson Med. 2010;64:229-238.

17. Stollberger RWP, McKinnon G, Justich E, Ebner F. RF-field mapping in vivo. In: Proceedings of the 7th Annual Meeting of ISMRM, San Francisco, CA, USA. 1998. p. 106.

18. van Schie JJ, Lavini C, van Vliet LJ, Vos FM. Feasibility of a fast method for B1-inhomogeneity correction for FSPGR sequences. Magn Reson Imaging. 2015;33:312-318.

19. Kingsley PB. Methods of measuring spin-lattice (T1) relaxation times: an annotated bibliography. Concepts Magn Reson. 1999;11:243-276.

20. Bojorquez JZ, Bricq S, Acquitier C, Brunotte F, Walker PM, Lalande A. What are normal relaxation times of tissues at 3 T? Magn Reson Imaging. 2017;35:69-80.

21. Cercignani M, Dowell NG, Tofts P. Quantitative MRI of the Brain: Principles of Physical Measurement. Boca Raton: CRC Press, Taylor & Francis Group; 2018. xvii, 342 pages p.

22. Rakow-Penner R, Daniel B, Yu H, Sawyer-Glover A, Glover GH. Relaxation times of breast tissue at 1.5T and 3T measured using IDEAL. Magn Reson Imaging. 2006;23:87-91.

23. Taylor AJ, Salerno M, Dharmakumar R, Jerosch-Herold M. T1 mapping: basic techniques and clinical applications. JACC Cardiovasc Imaging. 2016;9:67-81.

24. National Electrical Manufacturers Association. NEMA Standards Publication MS 1–2008 Determination of Signal-to-Noise Ratio SNR in Diagnostic MRI. New York: The Association; 2014.

25. Jiang Y, Ma D, Keenan KE, Stupic KF, Gulani V, Griswold MA. Repeatability of magnetic resonance fingerprinting T1 and T2 estimates assessed using the ISMRM/NIST MRI system phantom. Magn Reson Med. 2017;78:1452-1457.

26. Keenan KE, Gimbutas Z, Dienstfrey A, Stupic KF. Assessing effects of scanner upgrades for clinical studies. J Magn Reson Imaging. 2019;50:1948-1954.

27. Bane O, Hectors SJ, Wagner M, et al. Accuracy, repeatability, and interplatform reproducibility of T1 quantification methods used for DCE-MRI: results from a multicenter phantom study. Magn Reson Med. 2018;79:2564-2575.

28. Abramag A. The Principles of Nuclear Magnetism. Oxford: Oxford Science Publications; 1983:57.

29. Bartha R, Michaeli S, Merkle H, et al. In vivo 1H2O T2+ measurement in the human occipital lobe at 4T and 7T by Carr-Purcell MRI; detection of microscopic susceptibility contrast. Magn Reson Med. 2002;47:742-750.

30. Ye FQ, Martin WRW, Allen PS. Estimation of the iron concentration in excised gray matter by means of proton relaxation measurements. Magn Reson Med. 1996;35:285-289.

31. Vold RL, Vold RR, Simon HE. Errors in measurements of transverse relaxation rates. J Magn Reson (1969). 1973;11:283-298.

32. Majumdar S, Orphanoudakis SC, Gmitro A, O'Donnell M, Gore JC. Errors in the measurements of T2 using multiple-echo MRI techniques. II. Effects of static field inhomogeneity. Magn Reson Med. 1986;3:562-574.

33. Majumdar S, Orphanoudakis SC, Gmitro A, O'Donnell M, Gore JC. Error in the measurements of T2 using multiple-echo MRI techniques. I. Effects of radiofrequency pulse imperfections. Magn Reson Med. 1986;3:397-417.

34. Collins CM, Liu W, Schreiber W, Yang QX, Smith MB. Central brightening due to constructive interference with, without, and despite dielectric resonance. J Magn Reson Imaging. 2005;21:192-196.

35. Mirowski E, Snow M, Keenan K. inventors; Qalibre MD, Inc. (Boulder, CO, US), National Institute of Standards and Technology. (Gaithersburg, MD, US), assignee. Liquid Crystal Thermometer for MRI. US patent 10,809,331. October 20, 2020.

36. Keenan KE, Stupic KF, Russek SE, Mirowski E. MRI-visible liquid crystal thermometer. Magn Reson Med. 2020;84:1552-1563.

37. Keenan KE, Wilmes LJ, Aliu SO, et al. Design of a breast phantom for quantitative MRI. J Magn Reson Imaging. 2016;44:610-619.

SUPPORTING INFORMATION
Additional Supporting Information may be found online in the Supporting Information section.

FIGURE S1 A, The hemispherically machined surface on the post matches with the inclined notches, B, to precisely locate the plates. Photos of blue fiducial, C, and red MnCl2 array, D, spheres showing the sealing and mounting methods. D, inset shows micro-computed tomography image of contrast sphere

FIGURE S2 Relaxation rates as a function of paramagnetic salt concentration measured by inductively coupled Plasma (ICP) mass spectrometry at 1.5 T and 3.0 T. The solid lines are fits assuming a linear increase in relaxation rate with concentration and a zero-concentration intercept given by the measured values for high purity water

FIGURE S3 Schematic of the fiducial array analysis showing input 3D image of the system phantom, cropped image with just the fiducial spheres, image of fiducial sphere, synthetic k-space image and real-space image used as a convolution mask, slice of 3D convolution image, and convolution profiles with fits used to obtain sphere center

FIGURE S4 Fiducial analysis on a 1.5 T scanner with a gradient echo sequence. A, Coronal slice, B, axial slice, C, sagittal slice. D, magnified image of fiducial sphere image with ROI location after automated location. E, Normalized integrated intensity for all 57 fiducial spheres. F, Difference between center of mass and convolutional sphere centers
FIGURE S5 A, B, C, Geometric distortion of the 57 fiducial sphere centers in x, y and z directions, respectively

FIGURE S6 NMR signal for Ni-12 versus inversion time for the T1-IR protocol along with fits to the model described in the text. Data for three consecutive measurements are shown along with the residuals for each measurement (top plot). The errors listed for $T_1$ and the inversion efficiency are the standard deviation of the 3 values obtained for each of the measurements.

FIGURE S7 NMR signal from Ni-12 using a CPMG sequence as a function of acquisition time along with exponential fits. Data for three consecutive measurements are shown along with the residuals for each measurement (top plot). The errors listed for $T_2$ are the standard deviation of the 3 values obtained for each of the measurements.

FIGURE S8 Magnetic field dependence of $T_1$ and $T_2$ for the MnCl$_2$ and NiCl$_2$ arrays.

FIGURE S9 Temperature dependence of $T_1$, $T_2$ for the Ni-12 solution measured in a metrology NMR at 3.0 T. The plot shows data from a flame-sealed borosilicate-capillary library sample over the course of 4 years.

FIGURE S10 Variation of normalized relaxation times with temperature for the NiCl$_2$ array at 3.0 T.

FIGURE S11 Temperature coefficient of spin relaxation times for the MnCl$_2$ array at 3.0 T.

FIGURE S12 Variation of relaxation times with temperature for the CuSO$_4$ fiducial solution at 3.0 T.

FIGURE S13 Deviation of T1-IR values from NMR reference values at 3 T over the course of 7 years. The gray bar indicated the range of values given a phantom temperature that can vary between 18°C and 22°C.

FIGURE S14 Water mass uptake for various plastics: nylon/polyamide (PA), poly(methyl methacrylate) (PMMA), polycarbonate (PC), polyvinyl chloride (PVC), polyvinylidene fluoride (PVDF), polyethylene sulfide (PPS), polypropylene (PP). The samples were 25 mm diameter, 6 mm thick disks.

FIGURE S15 Geometric distortion during water soaking of the same samples used in Figure S13. The horizontal line indicates the threshold for maintaining the specified geometric distortion of the phantom plates.

FIGURE S16 Spin relaxation times for NiCl$_2$-3, NiCl$_2$-5, NiCl$_2$-10 at 3 T, 20°C as a function of pH.

FIGURE S17 Relaxation times for ACS-grade and deionized water as a function of temperature.

TABLE S1 NiCl$_2$ Array

TABLE S2 MnCl$_2$ Array

TABLE S3 Proton Density Array

TABLE S4 Water proton spin relaxivities determined from the slope of the data in Figure S2.

TABLE S5 Isotropic Volume Series

TABLE S6 Section Thickness Series

TABLE S7 Resolution Inset Series

TABLE S8 Proton Density and Signal to Noise Series

TABLE S9 $T_1$ Inversion Recovery Series

TABLE S10 $T_1$ Variable Flip Angle Series

TABLE S11 $T_2$ Series

TABLE S12 Starting Chemicals for Contrast Fluids

TABLE S13 NiCl$_2$ Solutions

TABLE S14 MnCl$_2$ Solutions

TABLE S15 Proton Density Solutions

How to cite this article: Stupic KF, Ainslie M, Boss MA, et al. A standard system phantom for magnetic resonance imaging. Magn Reson Med. 2021;86:1194–1211. https://doi.org/10.1002/mrm.28779