B₁ inhomogeneity correction of RARE MRI with transceive surface radiofrequency probes

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Purpose: The use of surface radiofrequency (RF) coils is common practice to boost sensitivity in (pre)clinical MRI. The number of transceive surface RF coils is rapidly growing due to the surge in cryogenically cooled RF technology and ultrahigh-field MRI. Consequently, there is an increasing need for effective correction of the excitation field (B₁⁺) inhomogeneity inherent in these coils. Retrospective B₁ correction permits quantitative MRI, but this usually requires a pulse sequence-specific analytical signal intensity (SI) equation. Such an equation is not available for fast spin-echo (Rapid Acquisition with Relaxation Enhancement, RARE) MRI. Here we present, test, and validate retrospective B₁ correction methods for RARE.

Methods: We implemented the commonly used sensitivity correction and developed an empirical model-based method and a hybrid combination of both. Tests and validations were performed with a cryogenically cooled RF probe and a single-loop RF coil. Accuracy of SI quantification and T₁ contrast were evaluated after correction.

Results: The three described correction methods achieved dramatic improvements in B₁ homogeneity and significantly improved SI quantification and T₁ contrast, with mean SI errors reduced from >40% to >10% following correction in all cases. Upon correction, images of phantoms and mouse heads demonstrated homogeneity comparable to that of images acquired with a volume resonator. This was quantified by SI profile, SI ratio (error < 10%), and percentage of integral uniformity (PIU > 80% in vivo and ex vivo compared to PIU > 87% with the reference RF coil).

Conclusion: This work demonstrates the efficacy of three B₁ correction methods tailored for transceive surface RF probes and RARE MRI. The corrected images are suitable for quantification and show comparable results between the three methods, opening the way for T₁ measurements and X-nuclei quantification using surface
transceiver RF coils. This approach is applicable to other MR techniques for which no analytical SI exists.

**KEYWORDS**

B₁ correction, B₁ inhomogeneity, MRI, RARE, signal intensity equation, transceive surface RF coils

1 | INTRODUCTION

The ability of MRI to provide high spatial resolution images within short acquisition times is governed by the sensitivity conundrum, which balances the constraints of signal-to-noise (SNR), image contrast, spatial resolution, and temporal resolution. Numerous approaches have been developed to improve SNR per scan time from the development of novel software-driven approaches (e.g., parallel imaging, compressed sensing), to hardware improvements, including higher magnetic field strengths (B₀) and the optimization of radiofrequency (RF) technology. The use of surface RF coils is common practice to boost sensitivity in (pre)clinical MRI, predominantly with a receive-only RF coil design in combination with a volume RF coil used for excitation. The use of transceiver (transmit-receive, TxRx) surface RF coils is increasing, in particular in human MRI at ultrahigh fields where large volume body RF coils are not used for signal excitation and are not provided with ultrahigh field-MR scanners.

In preclinical research, the use of transceiver RF configurations has been dominated by cryogenically cooled RF probes (CRP) that provide significant SNR gains. CRPs are sometimes also available as decoupled Rx-only configurations in combination with a room-temperature (RT) volume resonator for RF excitation, but are not as common as the Tx/Rx configuration. By reducing thermal noise in the receiver circuitry (RF probe and preamplifier), SNR can be enhanced by a factor of up to 3-4 compared to conventional RT RF coils. The SNR gain of a CRP can be exploited to increase spatial resolution, to reduce scan time, or to lower detection limits, especially for X-nuclei MRI.

A constraint of TxRx surface RF coil technology is the strong intrinsic spatial gradient (inhomogeneity) in both excitation (B¹⁺) and field sensitivity (B⁻¹). Although the latter can be easily corrected, non-uniform B¹⁺ fields induce significant spatial variations in the excitation flip angle (FA), with the effective FA decreasing with increasing distance from the RF coil surface. The resulting B¹⁺ inhomogeneities are more pronounced at higher field strengths. This adverse effect reduces image homogeneity and affects the T₁ image contrast, representing a major challenge for applications for which absolute signal intensities are needed, such as T₁ mapping and quantification techniques in X-nuclei MRI.

Although partial mitigation of B¹⁺ inhomogeneity can be achieved with adiabatic pulses, dielectric materials, or B¹⁺ shimming, retrospective B¹⁺ correction approaches are most commonly used to achieve signal uniformity. First, the actual FA is measured using magnitude- or phase-based B¹⁺ mapping techniques, such as the double angle method, the phase sensitive technique, the actual FA method, or any of their improvements. Then, an analytical description of the signal intensity (SI) dependency on the FA for the RF pulse sequence (SI equation) or numerical simulations are used to perform the SI correction.

Retrospective B¹⁺ correction has been successfully applied to gradient echo imaging techniques such as fast low angle shot or steady-state free precession, which are inherently less sensitive to RF inhomogeneity and for which SI equations are given. Retrospective B¹⁺ correction was also reported for spin-echo imaging methods. For fast spin-echo techniques such as Rapid Acquisition with Relaxation Enhancement (RARE), there is no exact analytical SI equation. This extends to pulse sequences employing simultaneous multislice parallel imaging, non-Cartesian trajectories, variable FA 3D turbo spin-echo, water-fat separation using Dixon approaches, and hybrid imaging techniques like half-Fourier single shot turbo spin-echo/gradient and spin-echo/turboGRASE (HASTE/GRASE/TGSE). Other complex techniques with no SI equation include those derived from ultrashort echo time or echo-planar imaging (EPI). As a consequence, retrospective B¹⁺ correction for these MRI techniques demands novel solutions.

To address this need, we developed, implemented, and applied three B₁ correction approaches for RARE MRI with transceive surface RF probes with the goal to reduce errors to less than 10% for SI quantification and for T₁ contrast. All three methods were applied and validated in test phantoms and mouse brains, in vivo and ex vivo. For performance evaluation, the corrected images were benchmarked against reference images obtained with a uniform TxRx volume resonator. The starting point was the commonly used sensitivity correction that uses a uniform phantom image to correct for B¹⁻ inhomogeneities. This method does not take spatial FA variations and T₁ relaxation times into account. Given this limitation and the unavailability of an analytical SI equation for fast spin-echo imaging, we modelled the SI of RARE as a function of FA and T₁ based on empirical measurements obtained through MR experiments. This model-based correction uses the SI model to correct B¹⁺, followed by a B¹⁻ correction. We also implemented a hybrid correction using a combination of the model-based and sensitivity correction approaches. These methods are valuable not only for conventional °H-MRI when
accurate FAs are needed (e.g., for well-defined T₁ contrasts), but also in X-nuclei MRI, for which absolute SI is essential for signal quantification.

2 | METHODS

The MR hardware, sample preparation, and measurements are summarized in Table 1.

2.1 | MR hardware

All experiments were performed on a 9.4 T small animal MR scanner (BioSpec 94/20, Bruker BioSpin, Ettlingen, Germany) operating at 400 MHz (¹H).

For creating reference images, we chose RF coils with approximately uniform excitation and reception fields:

- Small reference RF coil: in-house built volume resonator tailored for mouse head imaging (inner diameter [ID] = 18.4 mm).
- Large reference RF coil: rat body linear volume resonator (Bruker BioSpin) with an ID = 72 mm.

The correction methods were applied, evaluated, and validated for 2 transceive surface RF coils:

- Cryocooled surface RF coil (CRP): 2-element transceive RF probe (CryoProbe®, Bruker BioSpin) operating in quadrature mode with an ID = 20 mm and a saddle-shaped ceramic surface.
- RT surface RF coil: planar transceive single loop (ID = 20 mm) surface RF coil (Bruker BioSpin).

2.2 | Sample and animal preparation

To characterize the excitation and receive fields of the transceive surface RF coils (B₁ mapping), we used samples that ensured full field of view coverage and had low T₁ values (T₁ ≈ 300 ms) to reduce the needed TR (TR > 5·T₁) in our measurements:

- Cylindrical uniform phantom with low T₁: 15-mL tube (ID = 14.6 mm, length = 120 mm; Thermo Fisher Scientific, Waltham, Massachusetts) filled with a mixture of water and copper sulfate (Carl Roth GmbH & Co. KG, Karlsruhe, Germany).
- Rectangular uniform phantom with low T₁: 50-mL cell culture flask ((79.7 x 42.6 x 25) mm³; Fisher Scientific) filled with a doped solution of water.

Samples with different T₁ (NMR tubes, Thermo Fisher Scientific) filled with aqueous solutions of gadolinium (Magnevist® 0.5 mmol/ml; Bayer Vital, Leverkusen, Germany) at different concentrations (0.0-0.5 mM) yielding T₁ between 190 and 2871 ms were used to produce the RARE SI models.

The test phantoms used for correction and evaluation of the B₁ correction methods were:

- Cylindrical uniform phantom: containing water doped with gadolinium embedded in a 15-mL tube (T₁ ≈ 800 ms).
- Ex vivo mouse: the central nervous system of a SJL/J female mouse, perfused with a phosphate-buffered saline (Biochrom GmbH, Berlin, Germany), fixed in paraformaldehyde (PFA; Santa Cruz Biotechnology, Inc., Dallas, Texas), and placed into a 15-mL tube filled with 4% PFA.
- In vivo mouse: a healthy SJL/J mouse anesthetized with 2.7% isoflurane and stabilized with 1.6% during scanning. Heart rate, respiration rate, and temperature were monitored and kept constant during the examinations.

- Rectangular uniform phantoms: four 50-mL cell culture flasks filled with solutions of two different ¹H-atom concentrations: 100% water, 50% water and 50% deuterium oxide (Sigma-Aldrich, Saint Louis, Missouri). Gadolinium was added to the mixtures to achieve two different T₁ values (490 and 1525 ms).

All animal experiments were approved by the Animal Welfare Department of the LAGeSo State Office of Health and Social Affairs in Berlin and in accordance with international guidelines on reduction of discomfort (86/609/EEC).

2.3 | MR measurements

To characterize the B₁ fields of both transceive surface RF coils we used:

- FA mapping: fast low angle shot (FLASH) measurements with nominal excitation FAs of 60°/120° (RT) and 60°/120°/240° (CRP) with echo time/pulse repetition time (TE/TR) = 2.49/2000 ms, matrix = 128 x 128, 3 slices with a gap of 0.5 mm and a thickness of 2 mm each, TA = 1h30. We used a field of view = (25 x 25) mm² for the CRP and field of view = (35 x 35) mm² for RT.
- B₁ mapping: FLASH measurement with a nominal FA of 5° (same parameters as above).

B₁ field characterization can be performed prior to or after the image acquisition and does not entail extra acquisition time on the day of image acquisition, for example, in vivo scans.
To compute the models, we performed RARE measurements both with and without flipback and studied the effect of the extra pulse (which restores longitudinal magnetization, improving SNR) on the SI:

- **T$_1$-weighted (T$_1$w-) RARE scans (TE/TR = 2.49/1000 ms, echo train length (ETL) = 8, receiver bandwidth = 50 kHz, centric phase encoding, field of view = (25 × 25) mm$^2$, matrix = 128 × 128, 3 slices of 2 mm thickness, 

| Purpose | MR protocol | RF coil type | RF coil | Sample(s) | Acq. time |
|---------|-------------|--------------|---------|-----------|-----------|
| Test images | T$_1$w-RARE (with flipback) | Surface TxRx | CRP | Cylindrical uniform phantom | 60 min |
| | | | | Ex vivo mouse | 60 min |
| | | | | In vivo mouse | 30 min |
| Validation images | T$_1$w-RARE (with and w/o flipback) | Surface TxRx | RT | Rectangular uniform phantoms | 30 min |
| Sensitivity correction | T$_1$w-RARE (with and w/o flipback) | Surface TxRx | CRP | Cylindrical uniform phantom with low T$_1$ | 60 min |
| | | | | Rectangular uniform phantom with low T$_1$ | 30 min |
| Model-based correction | T$_1$w-RARE (with and w/o flipback) | Surface TxRx | CRP | Cylindrical uniform phantom with low T$_1$ | 90 min per FA |
| | | | | Rectangular uniform phantom with low T$_1$ | 30 min per FA |
| RARE SI modelling | T$_1$w-RARE (with and w/o flipback) | Volume TxRx | Small reference RF coil | Samples with different T$_1$ | 5 min 40 s per FA |
| T$_1$ mapping for modelling | RARE with variable TR | Volume TxRx | Small reference RF coil | Samples with different T$_1$ | 90 min |
| T$_1$ mapping for test images | RARE with variable TR | Volume TxRx | Small reference RF coil | Cylindrical uniform phantom | 100 min |
| | | | | Ex vivo mouse | 30 min |
| | | | | In vivo mouse | 55 min |
| T$_1$ mapping for validation images | RARE with variable TR | Volume TxRx | Large reference RF coil | Rectangular uniform phantoms | 30 min |
| Hybrid correction | Mapping of FA and B$_1^+$ FLASH | Surface TxRx | CRP | Cylindrical uniform phantom with low T$_1$ | 90 min per FA |
| | FLASH | Surface TxRx | RT | Rectangular uniform phantom with low T$_1$ | 30 min per FA |
| RARE SI modelling | T$_1$w-RARE (with and w/o flipback) | Volume TxRx | Small reference RF coil | Samples with different T$_1$ | 5 min 40 s per FA |
| T$_1$ mapping for modeling | RARE with variable TR | Volume TxRx | Small reference RF coil | Samples with different T$_1$ | 90 min |
| T$_1$ mapping for test images | RARE with variable TR | Volume TxRx | Small reference RF coil | Cylindrical uniform phantom | 100 min |
| | | | | Ex vivo mouse | 30 min |
| | | | | In vivo mouse | 55 min |
| T$_1$ mapping for validation images | RARE with variable TR | Volume TxRx | Large reference RF coil | Rectangular uniform phantoms | 30 min |
| T$_1$ mapping uniform phantom | RARE with variable TR | Volume TxRx | Large reference RF coil | Rectangular uniform phantom with low T$_1$ | 100 min |

CRP, cryogenically cooled radiofrequency probes; FA, flip angle; FLASH, fast low angle shot; RARE, Rapid Acquisition with Relaxation Enhancement; RT, room temperature; TxRx, transmit-receive.
TA = 5m40s). Thirty-five reference RF powers were used to vary the excitation FA in 5° increments, between 5° and 160° (flipback) and between 5° and 110° (without flipback).

- T1 maps of all phantoms (RARE with variable TR (120-15 000 ms); ETL = 2, linear phase encoding, other parameters same as RARE scan).

T1w-RARE images were acquired using the same parameters as above with flipback (CRP) and with/without flipback (RT) for validation purposes. Corresponding T1 maps for all samples were measured using RARE with TR ranging from 150 to 14500 ms.

All reference RF power adjustments were performed on a 2-mm slice located parallel and close to the RF coil surface.

RARE modeling can be equally performed prior to or after the image acquisition and does not entail extra acquisition time on the day of image acquisition, for example, in vivo scans.

## 2.4 | Approach 1: Sensitivity correction

All post-processing was performed using customized software developed in MATLAB (MathWorks Inc., Natick, Massachusetts).

This straight-forward method only requires a uniform phantom image to correct for the B1- inhomogeneities. The following steps were performed (Figure 1A):

- MRI study. Images (sample and uniform phantom) were acquired.
- Correction factor computation. We normalized and calculated the inverse of the uniform phantom image.
- Sensitivity correction. We multiplied the uncorrected image by the estimated correction factor to correct for the B1- inhomogeneities.

This method requires neither the characterization of the transceive RF coil used, nor the calculation of a RARE SI model and it is, therefore, directly applicable after image acquisition with little post-processing.

## 2.5 | Approach 2: Model-based correction

Figure 2 shows the workflow of the model-based correction, consisting of the following steps, starting with the quantification of the B1 inhomogeneities:

![Figure 1](image-url)  
**Figure 1** Workflows of (A) sensitivity correction and (B) hybrid B1 correction. The sensitivity correction merely requires dividing the sample image by that of a normalized uniform phantom. The hybrid method combined the model-based approach to perform a B1+ correction on the sample image and a uniform phantom image. The latter is then used to perform a B1- correction using the sensitivity correction method.
RF coil characterization. FA maps were calculated using the double angle method. To increase the SNR distal to the CRP we added a measurement at a higher FA and merged the 60°/120° and 120°/240° maps using an SNR cutoff. All maps were denoised using a polynomial fitting tool (polyfitn, 10th-order polynomials).

The transmit field (B⁺) maps were computed using:

\[
    \text{FA} = \gamma \cdot B^+ \cdot \tau
\]

with \( \gamma \) being the gyromagnetic ratio (\( \gamma = 267.522 \cdot 10^6 \text{ rad s}^{-1} \text{ T}^{-1} \) for \( ^1\text{H} \)) and \( \tau \) the pulse length of a rectangular RF pulse. Because calculated RF pulses were used, each one has a complex shape tailored to the sequence parameters. We therefore approximated the RF pulse length \( \tau \) using the product of the RF pulse duration, the area under the RF pulse \( (S_{\text{int}})^b \) and the related voltage \( (V) \):

\[
    B^+ = \frac{\text{FA} \cdot \pi / 180}{\gamma \cdot \tau \cdot S_{\text{int}} \cdot V}
\]

The RF coil sensitivity maps (B⁻) were calculated using the low FA approximation:

\[
    S_{\text{low FA}} \propto \left| B^+ \right| \cdot \left| B^- \right|
\]

where \( S_{\text{low FA}} \) was the 5° fast low angle shot measurement. The low FA image and \( B^+ \) map were normalized by their respective maximum values and \( B^- \) calculated as:

\[
    B^- / \max (B^-) \propto \frac{B^+ / \max (B^+)}{S_{\text{low FA}} / \max (S_{\text{low FA}})}
\]

Ultimately, the \( B^- \) map was denoised using a 10th-order polynomial fit.

- Modeling of the RARE SI equation. The relationship between SI, FA, and \( T_1 \) was estimated using experimental data and a fitting tool:

\[
    \text{FIGURE 2} \quad \text{Workflow of model-based B}_1 \text{ correction. The necessary images and maps to be acquired are described in MR Measurements & Post-processing column. Then the flip angle (FA) and sensitivity (B⁻) maps were calculated using the double angle method and the low FA approximation, respectively. The Rapid Acquisition with Relaxation Enhancement (RARE) signal intensity model was derived from a 2D fit of the signal intensities measured for different FAs and \( T_1 \) relaxation times using a volume resonator. The B⁺ correction factor was computed pixel-wise for the actual FA and \( T_1 \) using the RARE signal intensity model. Applying this correction factor and the B⁻ map derived correction factor yielded the final B₁ corrected image.}
\]
by fitting \( S = S_0(1 - \exp(-TR/T_1)) \) to the SIs using in-house developed software in MATLAB. We assumed \( SI(FA = 0^\circ) = 0 \) for all \( T_1 \). For each \( T_1 \) sample, a circular region of interest (ROI) was drawn to extract average SI and \( T_1 \) values from the images and maps respectively.

RARE modeling: To model the SI = f(FA, \( T_1 \)) relationship a 7th-order 2D polynomial was fitted to the experimental data using MATLAB’s `polyfitn` function. This was the lowest polynomial order that gave an \( R^2 > 0.99 \) and a faithful representation of the measured data.

- MRI study. Images and corresponding \( T_1 \) maps of the test samples were acquired.
- Retrospective correction. All images and maps \((B_{1+}/B_{1-}/T_1)\) were spatially aligned, either by careful slice planning or by image registration.

The \( B_{1+} \)-correction factor \((f_{corr})\) was calculated as the modeled RARE SI for a perfect 90° excitation \((SI_{nominal})\) divided by the modeled RARE SI for the actual excitation FA \((SI_{actual})\) obtained from the FA map:

\[
f_{corr} = \frac{SI_{nominal}}{SI_{actual}}
\]

Applying this correction factor yielded a \( B_{1+} \)-corrected image:

\[
image_{B1+corr} = image \cdot f_{corr}
\]

In the few cases where the algorithm produced negative values (low-SNR regions), the correction factor was set to zero.

Dividing this \( B_{1+} \)-corrected image by the \( B_{1-} \) map produced the final \( B_{1-} \)-corrected image:

\[
image_{corr} = image_{B1+corr}/B_{1-}
\]

2.6 | Approach 3: Hybrid correction

This method combines the sensitivity and model-based correction (workflow in Figure 1B), and involves:

- RF coil characterization (as in Model-based Correction).
- Modeling of the RARE SI equation (as in Model-based Correction).
- MRI study. Images and \( T_1 \) maps of the samples and of a uniform phantom were measured.
- Model-based \( B_{1+}^c \) correction (as in Model-based Correction) was performed on the sample and uniform phantom image.

- \( B_{1-} \) correction (as in Sensitivity Correction). The inverse of the \( B_{1+}^c \)-corrected uniform phantom image was applied as the \( B_{1-}^c \) correction factor to the \( B_{1+}^c \)-corrected sample image.

Both the model-based and the hybrid correction methods need a prior/posterior characterization of the transceive RF coil used and the calculation of a RARE SI model. The post-processing needed is rather simple in both cases.

2.7 | Correction method evaluation and validation

The presented \( B_1 \) correction techniques were validated using the following methods:

2.7.1 | Central profile plots

The SI profile along a central line perpendicular to the RF coil surface was plotted against distance to the RF coil surface. Seven pixels across the width of the line were averaged, and the SIs were normalized to \([0,1]\) to allow a better comparison. A quantitative comparison was performed by calculating the root-mean-square-error (RMSE) between each profile and the reference. Each profile was scaled to minimize the RMSE against the reference, in order to compensate for the arbitrary scaling and to provide a fair comparison.

2.7.2 | Image homogeneity assessment

To quantitatively assess the uniformity of the corrected images, the percentage of integral uniformity (PIU)\(^8\) was computed for several ROIs of different sizes. A PIU of 100% represents perfect image homogeneity. In the uniform phantom, we defined 5 internally tangential circular regions of interest (ROIs) with increasing diameter on the central vertical line. For the brain images (ex vivo, in vivo), we manually outlined the cortex and basal ganglia/thalamus (left and right), achieving 3 ROIs.

2.7.3 | \( T_1 \)-contrast and quantification performance

We used the experimental setup (Figure 3A) to compare substances with different water content (100% or 50%, respectively) and different \( T_1 \) relaxation times (490 or 1525 ms, respectively). All acquired images with and without flipback were corrected using the three \( B_1 \) correction methods. Five ROIs were drawn at
pseudo-randomized positions (Figure 3B) on all sets of images (three corrections, original and reference) for all flasks. For each of the flask image pairs described in Figure 3A, mean SI ratios were calculated using all possible ROI combinations for all sets. Relative ratio errors were computed:

\[
\text{Ratio error} = \frac{\text{abs}(\text{SI}_{\text{reference}} - \text{SI}_{\text{corrected}})}{\text{SI}_{\text{reference}}} \times 100(\%)
\]

With \(\text{SI}_{\text{reference}}\) being the mean SI ratio computed using all ROI combinations on the reference image pairs, and \(\text{SI}_{\text{corrected}}\) being that achieved using the corrected image pairs. Finally, the mean error and mean SD were calculated. An example of the workflow is shown in Figure 3B.

Statistical analysis. A nonparametric 1-way analysis of variance Friedman repeated measures test was performed (mean errors on the original data did not have a Gaussian distribution) followed by Dunn’s test where all corrections were compared to the original data (\(P\) values < .05 were considered significant). All statistical assessments were performed using GraphPad Prism 5 (GraphPad Software, La Jolla, California).

3 | RESULTS

3.1 | RF coil characterization

The maps of the receive field (\(B_{r}^-\)) (Figure 4A) and transmit field (\(B_{t}^+\), here as FA) relative to a 90° excitation FA (Figure 4B) demonstrate the spatially varying sensitivity and FA for the CRP. A closer look at the vertical midline profile reveals a strong deviation from the target of FA = 90° (nominal FA) with increasing distance from the surface of the CRP (Figure 4C). These field maps show the typical inhomogeneity inherent to transceive surface RF coils, which was very similar in the \(B_{r}^-\) and \(B_{t}^+\) maps and FA profiles for the single loop RF coil (Figure 4D-F). The minor deviation of the FA profiles at 20-30 mm from the coil surface (in gray) reflects a mathematical artifact of the polynomial fit at low-SNR regions.
3.2 | Modeling of the RARE SI equation

The RARE SI dependency on FA and $T_1$ ($SI = f(FA, T_1)$) was modeled by fitting a polynomial to the experimental data acquired with these parameters, either incorporating a flipback pulse to restore longitudinal magnetization and hence improve SNR (Figure 5A-C), or excluding flipback to allow natural relaxation (Figure 5D-F). The fitted 3D-surfaces are shown in Figure 5A,D. Two-dimensional projections of the RARE models show the relationships between SI and $T_1$ for several FA values (Figure 5B,E) and between SI and FA for several $T_1$ values (Figure 5C,F). As expected, the fitted SI data predicts lower SI with increasing $T_1$ and maximal SI for FAs around 90°. The surface fits modeled the experimental data well ($R^2 = 0.997$ in both cases).

3.3 | Correction method evaluation and validation

We acquired $T_1$ maps (needed for $B_{1}^\dagger$ correction) and reference images of a uniform phantom, an ex vivo mouse phantom, and an in vivo mouse brain using a volume resonator (Figure 6A-B). The original uncorrected CRP images show the strong spatial SI gradient typical of transceive surface RF coils (Figure 6C). The results obtained with the three $B_1^\dagger$ correction methods are shown in Figure 6D-F. The strong spatial SI gradient present in the CRP images was removed by all $B_1^\dagger$ correction methods, yielding a uniform SI throughout the entire field of view for all investigated samples, including the in vivo mouse head. With the sensitivity and model-based corrections we observed an overshoot in SI in some regions, particularly distal to the CRP. This was due to a combination of increasing inaccuracies in the FA and SI data at low SNR. This overshoot in SI was resolved by combining both methods in the hybrid correction approach.

3.3.1 | Central profile plots (CRP)

To quantitatively assess the correction of the image inhomogeneity, we plotted normalized vertical SI profiles (Figure 7A-C). For all three approaches, the corrected SI profiles showed close correspondence with the reference RF coil (plotted as a surface in green). From these profiles one can determine how far away from the RF coil it is still viable to perform $B_1^\dagger$ correction. This depends on the specific scanning parameters and the dimensions of the RF coil; here this distance was approximately 17 mm (for a nominal FA of 90°, an actual FA of up to 8° could be
FIGURE 5. Signal intensity (SI) models for Rapid Acquisition with Relaxation Enhancement (RARE) with and without flipback. (A,D) 3D-plots of the modelled RARE signal intensity (SI) as a function of the T1 relaxation time and flip angle (FA) with and without flipback, respectively (R^2 = 0.997 for both). (B,E) show the SI vs FA projection in both models, whereas (C,F) depict the SI vs T1 projection. Selected FA and T1 values are plotted to demonstrate the fidelity of the experimental data and the model. Each colored line depicts a different T1 and FA, respectively. The dots represent the measured data points.
corrected). For our experimental setup, the region beyond 17 mm showed increasing inaccuracies in the field maps and SI measurements, leading to unacceptable errors in all corrected images.

Quantitative examination revealed that all correction methods considerably reduced the RMSE computed on the profiles to a maximum of 0.18 (uniform), 0.12 (ex vivo) and 0.26 (in vivo), with respect to the reference. For the uniform phantom, the sensitivity and hybrid approaches performed equally well (0.11). For the ex vivo phantom the sensitivity and model-based correction performed similarly (0.11). In vivo, the sensitivity correction achieved the best result (0.21). In comparison, the uncorrected profiles revealed an average RMSE of 0.53 ± 0.07 for all test phantoms.

3.3.2 | Image homogeneity assessment (CRP)

For the uniform phantom, we found the calculated PIU (Figure 7D-F) to be 95.7% within the largest ROI using the volume resonator, indicating no significant inhomogeneities across the image, as expected. Conversely, a PIU of 0.9% was obtained within the same ROI on the uncorrected image. The PIU degradation scaled with increasing ROI diameter. After correction, the model-based approach showed a PIU of 65% on the fourth ROI (up to a distance of 16.2 mm from the RF coil surface). Beyond that distance, the observed overshoots confounded the PIU, which decreased to 0% in the largest ROI.

For the mouse brain images the PIUs showed the expected high homogeneity for the reference RF coil: ex vivo 87.0 ± 4.4% and in vivo 87.7 ± 9.1%. The original surface RF coil images displayed substantial inhomogeneities: averages of 35.4 ± 9.2% ex vivo and 33.2 ± 11.8% in vivo. A significant improvement in image homogeneity was achieved with all three correction methods, both in vivo and ex vivo. The model-based method performed best on average (85.0 ± 3.8% ex vivo and 80.5 ± 11.3% in vivo), closely followed by the hybrid (81.6 ± 6.9% ex vivo and 79.7 ± 11.2% in vivo) and sensitivity (80.8 ± 5.7% ex vivo and 76.5 ± 10.3% in vivo) corrections.

3.3.3 | T1-contrast and quantification performance (RT)

We studied the errors in SI ratios between several fixed locations for all four phantoms, comparing original (uncorrected) RARE images and their three corrections, relative to the ground truth (reference images). These validation assessments were performed for RARE without flipback (Figure 8) and with flipback (Figure 9). The box plots (whiskers at 5-95 percentile) depict the mean errors for quantification at low and high T1 relaxation times, and for T1 contrast measurements with low and high proton
density. Errors below 10% (dashed line) were considered acceptable.

Correction of RARE MR images without flipback (Figure 8): All correction methods reduced the errors to less than 10% for both quantification and contrast, contrary to uncorrected images that showed substantial errors (41-45%) and variabilities (37-42%). None of the calculated mean errors reached a value >8.3% after correction.

The sensitivity correction performed best when calculating water content proportions at low T1 values (5.0 ± 2.9%), followed closely by the hybrid (6.0 ± 2.7%) and model-based (6.6 ± 4.5%) methods. All three methods behaved similarly for higher T1 values, with mean errors of approximately 8% (sensitivity 8.1 ± 2.9%, model-based 8.3 ± 5.9%, hybrid 8.1 ± 3.3%). All correction methods improved quantification significantly (P value < .0001) when compared to the original data.

When measuring T1 contrast, the hybrid method performed best for both water content phantoms (2.4 ± 1.7% high, 4.7 ± 3.8% low). The sensitivity correction method performed better than the model-based method for the high water content phantom (3.5 ± 2.5% vs. 6.2 ± 5.5%). However, for the low water content comparison, the model-based correction method performed better than the sensitivity correction (5.2 ± 3.9% vs. 6.1 ± 3.1%). Similarly, the three described correction methods significantly improved T1 contrast, when compared to the original data (P value < .0001).

Correction of RARE MR images with flipback (Figure 9): In general, all correction methods performed worse when flipback was enabled in RARE measurements, compared to RARE without flipback. The errors without correction were comparable to the case without the flipback option (40-58%). Their variabilities, however, were spread along a wider range (40-62%).

For quantification, the correction methods performed worse at low T1 relaxation times (overall about 10%: sensitivity 11.0 ± 7.6%, model-based 10.7 ± 7.9%, hybrid 12.2 ± 8.6%) than at higher ones (sensitivity 4.8 ± 4.0%, model-based 11.4 ± 10.1%, hybrid 7.2 ± 6.0%). All correction methods significantly improved quantification when compared to the original data (P < .0001).

T1 contrast accuracy was considerably reduced when using flipback during the measurements, with errors approaching 20-30% for high water content. The sensitivity correction
method (19.5 ± 9.7%) performed marginally better than the model-based (28.9 ± 19.4%) and hybrid (28.4 ± 14.5%) methods. For higher water content the errors were smaller (8-15%). Similarly, the sensitivity correction method (8.3 ± 5.0%) performed slightly better than the other two (model-based 15.2 ± 13.2%, hybrid 15.2 ± 8.7%). Only the sensitivity method significantly improved T1 contrast (P value = .0002 and .0003 for high and low proton density, respectively).

FIGURE 8 Assessment of quantification and contrast accuracy for Rapid Acquisition with Relaxation Enhancement (RARE) without flipback. Box plot of relative quantification and contrast errors for the original uncorrected images and those corrected with each of the three B1 correction methods. All B1 correction methods reduced the median error from well above 25% to below 10% (dashed line). Whiskers represent the 5 and 95 percentiles. Asterisks indicate statistically significant differences compared to the uncorrected images.

4 | DISCUSSION

Several methods have been described in the literature to correct B1 inhomogeneities. These methods are especially crucial for images acquired with transceive surface RF coils. The current study extends this work by demonstrating the feasibility and efficacy of B1 field inhomogeneity correction methods for RARE MRI, for which an analytical SI equation is not available. Our phantom results showed a substantial improvement in image homogeneity after B1 correction using the methods we investigated. We also establish the feasibility of these approaches for samples with more complex structures (ex vivo and in vivo mouse) and in time-constrained scenarios (in vivo). These results demonstrate that images derived from the correction procedures are suitable for accurate T1 contrast and SI quantification purposes, thus opening the way for parametric T1 mapping and X-nuclei quantification using surface transceiver RF coils/probes. Compared to previously developed correction methods, the approaches presented and evaluated here are applicable to MR imaging techniques for which no analytical SI equation exists, including but not limited to echo-planar imaging and ultrashort echo time imaging techniques.
The sensitivity correction method is well established in the literature for correction of sensitivity-related inhomogeneities. A typical application is the correction of $B_1^-$ inhomogeneities in a RF coil setup where a volume resonator is used for transmission and a surface RF coil (with or without cryocooled technology) for MR signal detection. We demonstrate here that this method is also effective for correction of $B_1^+$ inhomogeneities. The sensitivity correction method includes an inherently linear $B_1^+$ correction, because all images are the product of the transmission and reception capabilities of an RF coil. This concept is supported by the quasi-linear trends shown in our SI model for SI vs. T1 relaxation time, and the linear trends present for the majority of the SI vs. FA range (e.g. between 30°-70° and between 90°-140°).

The two novel $B_1$ correction methods (model-based, hybrid) we propose use an empirical SI model of the RF pulse sequence. The correction workflow involves using the calculated SI model to adjust the SI to that of the nominal FA, based on the actual FA and $T_1$. This rectifies the inhomogeneities related to RF transmission ($B_1^+$), whereas those related to the RF coil sensitivity ($B_1^-$) are addressed in a separate step using a previously calculated $B_1^-$ map (model-based) or using a $B_1^+$-corrected uniform phantom (hybrid).

Homogeneity was first assessed calculating the PIU and central SI profiles in the corrected and reference images. These tests revealed a high homogeneity, maintained when comparing the ex vivo phantom to the in vivo situation (difference in mean below 5%). A clear difference was found in the profile comparison (RMSE = 0.12 vs 0.26, ex vivo and in vivo respectively), which might be related to a change in animal position when transferring the animal from $^1$H-CRP to reference RF coil. These differences might be also caused...
by motion (eg, due to misalignment of the FA map, worse \(B_0\) shimming, etc.). Although the option motion averaging was used, it might not have been enough to compensate for bulk motion. Because we were using a RARE-based imaging sequence where the blood signal in large vessels is inherently suppressed due to the use of a spin-echo train, we do not believe the changes in performance to be related to blood flow.

Assessing the accuracy of SI quantification and \(T_1\) contrast measurements yielded different results for RARE with and without flipback that drives the equilibrium regimen. Without the driven equilibrium regimen all correction methods reduced the errors to less than 10\% for both quantification and \(T_1\) contrast, and produced statistically significant improvements compared to the original data. For the driven equilibrium regimen, the errors in the original data were more pronounced, which translated into higher SI quantification and \(T_1\) contrast errors after correction. For all three \(B_1\) correction methods, errors were around 10\% for quantification, but the accuracy of \(T_1\) contrast was considerably reduced, with errors up to 20-30\% (for high water content). Only the sensitivity method improved \(T_1\) contrast significantly.

When flipback was inactive, all three methods performed similarly for SI quantification purposes, and yielded improved performance for the low \(T_1\) mode. This can be attributed to the reduced \(T_1\)-weighting at the repetition times used, so that less correction was needed. The sensitivity correction method performed slightly better than the other two for SI quantification purposes. The simplicity of this approach makes it preferable for absolute SI quantification. Conversely, our results showed that the hybrid correction provides more accuracy when \(T_1\) contrast is essential (eg, for contrast-enhanced imaging in inflammatory disease).

Overall, the hybrid method performed better than the model-based one. Because the only difference between them is the sensitivity profile calculation, we conclude that the simple sensitivity correction performs better than the low FA approximation when computing a \(B_1^+\) map from measurements. The minor artifacts produced at regions distal to the coil are caused by inaccuracies in the FA information associated with low SNR.

The described model-based approach is fundamentally limited by SNR constraints at larger distances from the RF coil, and by the accuracy of the \(B_1\) and \(T_1\) maps and the polynomial fit. Determining the distance until which a meaningful correction can be achieved is challenging, since it depends on the conditions and scanning parameters used (eg, coil dimensions, SNR, acquisition time). Hence, this distance should be determined by each user, for each specific setup: (1) calculate the central profile plots for each correction and (2) determine at what distance from the coil the corrected profile still follows that of the reference volume RF coil. This will not require extra time, because \(T_1\) mapping with the reference RF coil is anyway required for the \(B_1\) correction.

Accurate knowledge of \(T_1\) and FA is crucial for the precise correction of the \(B_1\) inhomogeneities using the model-based and hybrid methods. For our workflow, we selected readily-available MR imaging protocols (eg, double angle mapping, RARE with variable TR). Limitations are related to the inherent instability associated with the FA and \(T_1\) mapping techniques, the lack of an established gold standard, and substantial variability among the different methods. FA mapping depends on the slice excitation profile, \(B_0\) homogeneity and other factors, which produce additional uncertainties. Moreover, FA mapping techniques are usually imprecise for low FAs, increasing the FA error at large distances from the RF coil.

\(T_1\) mapping is equally challenging and subject to many sources of error. Fundamentally dependent on the FA, it is usually performed using volume resonators or a combination of RF coils for transmit-receive (volume for transmit, surface for receive) to attenuate the effects of \(B_1^+\) inhomogeneity. A caveat of these methods (model-based and hybrid) is the need to acquire a \(T_1\) map with each image (in order to consider the \(T_1\) contrast of tissues) when removing the field inhomogeneities in \(^1\)H images. Although \(T_1\) mapping is feasible using a cryocooled RF probe, we invested the extra time and used a volume resonator to reduce \(T_1\) map errors.

An alternative to calculate the signal evolution (SI model) would be to use extended phase graph\(^87\) or Bloch\(^88\) simulations. Equally, magnetic resonance fingerprinting\(^89\) could be used not only to create the model but also to acquire a \(T_1\) and \(B_1\) map altogether by changing FA and TR, reducing the amount of scan time needed and producing a tailored correction (“real” \(B_1\) map of the phantom/mouse). To our knowledge there are no magnetic resonance fingerprinting-RARE techniques available to date and the development of such MR sequences was out of the scope of this study.

When considering SNR, it is important to bear in mind that these correction methods entail multiplication by a position-dependent matrix of correction factors. Thus, both the signal and the noise will be increased; furthermore, this effect will be different for each image pixel. Therefore, SNR calculations must be performed on the original uncorrected images.

The \(B_1\) correction methods presented here have widespread implications. We demonstrated that these methods are not only useful for the specific case of cryogenically cooled RF probes, frequently used to boost SNR in preclinical MRI,\(^24,40,90\) but are also generally applicable for transceive surface RF coils like single-loop RF coils. We demonstrated the applicability of the correction methods in conventional \(^1\)H brain imaging; however, these methods can also be applied to moving organs, for example, cardiovascular research, as long as the calculated reference power is correct and the maps and images are acquired using a trigger and spatially aligned. These approaches are also highly relevant for quantitative MR of X-nuclei, where absolute SI is important.\_ENREF_39,40 For the low-SNR scenarios that are
prevalent in X-nuclei imaging, the procedures that are used to validate the correction methods described in this manuscript (e.g., PIU, central profile plots) might not be entirely valid. In these cases, we suggest performing error propagation simulations associated with inaccuracies due to the low SNR or to simply use the sensitivity correction method.

Interestingly, all correction methods we studied here greatly improved SI quantification and image contrast, with only minor differences in performance of the three approaches. The best results were obtained with the hybrid correction, but contrary to expectations, even the straightforward sensitivity correction performed well. Therefore, one could recommend this last method due to its simplicity. These $B_1$ correction methods permit quantitative SI and $T_1$ contrast measurements with transceiver surface RF coils, using MRI techniques for which analytical SI equations do not exist. This circumvents a key limitation and offers a new approach for correcting $B_1$ inhomogeneity that may be applied for a broad range of biomedical research applications.

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CONFLICT OF INTEREST

Andre Kuehne is an employee of MRI.TOOLS GmbH, Berlin, Germany. Thoralf Niendorf is founder and chief executive officer of MRI.TOOLS GmbH, Berlin, Germany. All other authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT

The code and data that support the findings of this study are openly available in GitHub at https://github.com/pramosdelgado/B1correction-toolkit.

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