**MP2RAGEME: T₁, T₂*, and QSM mapping in one sequence at 7 tesla**

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**Abstract**
Quantitative magnetic resonance imaging generates images of meaningful physical or chemical variables measured in physical units that allow quantitative comparisons between tissue regions and among subjects scanned at the same or different sites. Here, we show that we can acquire quantitative T₁, T₂*, and quantitative susceptibility mapping (QSM) information in a single acquisition, using a multi-echo (ME) extension of the second gradient-echo image of the MP2RAGE sequence. This combination is called MP2RAGE ME, or MP2RAGEME. The simultaneous acquisition results in large time savings, perfectly coregistered data, and minimal image quality differences compared to separately acquired data. Following a correction for residual transmit B₁+-sensitivity, quantitative T₁, T₂*, and QSM values were in excellent agreement with those obtained from separately acquired, also B₁+-corrected, MP2RAGE data and ME gradient echo data. The quantitative values from reference regions of interests were also in very good correspondence with literature values. From the MP2RAGEME data, we further derived a multiparametric cortical parcellation, as well as a combined arterial and venous map. In sum, our MP2RAGEME sequence has the benefit in large time savings, perfectly coregistered data and minor image quality differences.

**KEYWORDS**
QSM, quantitative imaging, T₁ mapping, T₂* mapping

## 1 | INTRODUCTION

Quantitative magnetic resonance imaging (MRI) is becoming a popular tool in neuroimaging, in a large part due to the increasing availability of 7 T MR scanners, where the increased signal-to-noise ratio (SNR) accommodates more complex and generally longer acquisitions (van der Zwaag, Schäfer, Marques, Turner, & Trampel, 2016). With quantitative MRI, we depart from volumetric representations of the underlying tissues, and obtain a directly comparable measure of more tissue-specific MRI quantities that may not be captured by studying volume alone (Draganski et al., 2011). With such measures, we can study changes due to aging, disease, or learning-induced plasticity with increased specificity (Keuken et al., 2017; Tardif et al., 2016; Vargas et al., 2018; Yeatman, Wandell, & Mezer, 2014). Moreover, the measures are more directly reproducible across scanners and sites, even for different sequences (Stikov et al., 2015; Weiskopf et al., 2013).

The main compounds that can be quantitatively measured in the brain are myelin and iron. In white matter (WM), the longitudinal relaxation rate R₁ = 1/T₁ has been shown to be linearly related to myelin concentration (Stüber et al., 2014) and axon diameter (Harkins et al., 2016). In gray matter, both R₂* and quantitative susceptibility mapping (QSM) are linearly related to iron concentration (Deistung et al., 2013a). Cortical gray matter shows a spatial distribution of both myelin and iron, apparent in T₁- and T₂*-maps (Waehnert et al., 2016).

While T₁ contrast is used widely to define the gray-WM border along the cortical surface (Bazin et al., 2014; Fischl et al., 2002), T₂* and derived contrasts are widely used to segment deep gray matter structures (Keuken, Isaacs, Trampel, van der Zwaag, & Forstmann, 2018).

T₁ relaxometry can be done with multiple 3D FLASH (fast low angle shot) acquisitions (Frahm, Haase, & Matthaei, 1986) with variable excitation flip angles (Helms, Dathe, & Dechent, 2008). An
additional B1-map may be obtained to correct for transmit field (B1\(+\)) inhomoogeneities, such as by means of a dual refocusing echo acquisition mode in the DREAM-sequence (Nehrke & Börnert, 2012). Other work performed a segmentation-based correction, alleviating the need of a separate B1-map (Weiskopf et al., 2011). Acquiring multiple multi-echo gradient echo (ME-GRE) readouts enabled quantitative multi-parameter mapping of R2 and R2\(+\) (Weiskopf et al., 2013). These parameters could be robustly estimated over multiple sites. In order to reduce the sensitivity to B1\(+\)-inhomoogeneities and alleviate the need to coregister separately acquired volumes, the MP2RAGE-sequence was proposed. In this inversion-recovery sequence, two GRE-readouts follow after optimized inversion times. This allows for high-resolution imaging at high field (Marques et al., 2010; Marques & Gruetter, 2013). The resulting image is free of T2\*, M0 (net magnetization or proton density) and B1\(+\) effects, but a small and protocol-dependent amount of B1\(+\) contrast remains. This can be removed by acquiring a B1\(+\)-map in addition to T1-weighted data (Marques & Gruetter, 2013) and use it in the parameter estimation step. Although the MP2RAGE sequence is robust and widely used at 7 T, the acquisition is relatively inefficient because of the long TR required for the magnetization to return to equilibrium. Longer readouts, containing more k-space lines, would lead to shorter scan times, but also result in more T1\(-\)relaxation during the readout and, hence, incur more T1\(-\)-induced blurring in the images, leading to a poorer PSF. Therefore, most acquisitions opt for relatively short readouts of typically one k-space plane, limiting blurring and accepting the acquisition dead time. Optimally exploiting the dead time, as we will propose in this article, will result in a time-efficient sequence.

Faster imaging at high field was achieved by using a multislice echo planar imaging (EPI) readout (Polders, Leemans, Luijten, & Hoogduin, 2012; van der Zwaag et al., 2018; Wright et al., 2008), but EPI imaging comes at the cost of spatial distortions due to the lower readout bandwidth. Where the MP2RAGE sequence is limited to two readouts, the MPnRAGE performs many more radial readouts with view sharing, sampling the relaxation curve over a wider range (Kecskemeti et al., 2015).

\(T_2\) relaxometry is commonly performed using a FLASH sequence with a readout comprising of ME, QSM is performed by a dipole deconvolution of the magnetic field, obtained from the phase data of this sequence (Wang & Liu, 2015).

Recent work showed that combined T1, T2\*, and QSM mapping is feasible by extending the MP2RAGE sequence to have ME at both readouts (Metere, Kober, Möller, & Schäfer, 2017). This makes efficient use of the dead time in the MP2RAGE sequence, although the ME on the first image following the inversion necessarily leads to a longer readout block and, subsequently, to a nonoptimal inversion time for T1\(-\)-sensitivity as well as a too-short TE for optimal T2\* contrast (TEs should at least match the expected T2\* values of tissues).

T1, T2\*, and QSM are thus widely used contrasts at 7 T and mapping these properties quantitatively is more and more sought after. While indeed many approaches can be used to measure these in reasonable times, challenges arise in terms of inhomogeneities or bias in the estimated quantities and precision of their alignment. Simultaneous acquisition avoids the need for coregistration and subsequent resampling of separately acquired scans and allows time savings while compromises to both maps are minimal.

Here, we present an extension of the MP2RAGE sequence in which ME are acquired on the second inversion. This allows measurement of T1, T2\*, and QSM simultaneously and efficiently, with optimized inversion and longer echo times, at high resolution in maximally 17 min, reducing the sequence dead time to 6%, and show that the measured quantities agree well with expected measures after correction of B1\(+\)-inhomoogeneities.

## 2 METHODS

The MP2RAGE sequence (Marques et al., 2010) was modified so as to acquire ME in the second inversion while maintaining the single-echo acquisition in the first inversion (Figure 1). Specifically, the longest TE of the second inversion will be designed to be in the range of reported T2\* values of GM and WM in the brain.

The signal model for the MP2RAGEME sequence for given inversion, echo and repetition times can be analytically written out, as given in the Appendix. T1 values were computed from this model through a lookup table with sequence parameter values.

### 2.1 Subjects

Eight healthy volunteers were scanned at 7 T (Philips, Best, NL). The Multiple Interleaved Scanning Sequences environment provided by Philips was used to execute the alternation between the two gradient echo blocks. All subjects provided written informed consent and the study was approved by the local Medical Ethical Committee. From two subjects, the data acquisition was not completed due to hardware failure. For the other six subjects (age range 18–38, four males, and two females), the data were further analyzed.

### 2.2 Sequence

The MP2RAGEME data were compared with separately acquired MP2RAGE and ME-GRE data in the same session. Parameters in common between all three sequences were: field of view: \(205 \times 205 \times 164\) mm, matrix size: \(320 \times 320 \times 256\) and hence resolution: \(0.64 \times 0.64 \times 0.64\) mm. All imaging slabs were oriented sagittally and rotated 10° around the left–right axis to avoid overlap between the wrapped around signal from the nose and the cerebel- lum. A SENSE acceleration of \(1.7(LR) \times 1.7(AP)\) was used in combination with a circular k-space window to reduce scan time. The readout bandwidth was 405 Hz for all scans.

We required the longest TE of the second inversion block of the MP2RAGEME to be at least 26 ms, to match the T2\* of gray matter (Marques & Norris, 2017; van der Zwaag et al., 2016). TE\(_{2\alpha}\), the first echo of the second inversion, has to match TE\(_{1}\) to be able to cancel out the T2\* weighting in the T1\(-\)-weighted MP2RAGE data. A shorter TE results in higher SNR in the T1\(-\)-weighted image. Hence, TE\(_{2\alpha}\) was chosen as short as possible. This led to the range of TE\(_{2\alpha}\) of 3/11.5/20/28.5 and a minimal TR\(_{2}\) of 31.4 ms. The resulting longer readout of the second GRE block of the MP2RAGEME necessitated

| Sequence | Parameters |
|-----------|------------|
| MP2RAGEME | \(205 \times 205 \times 164\) mm, \(320 \times 320 \times 256\) |
that TRMP2RAGE was extended from 6 to 6.7 s. In summary, the following sequences were acquired:

- **MP2RAGEME**: TRMP2RAGE = 6.72 s, TR1 (the repetition time of the first GRE block) = 6.2 ms, TR2 = 31.4 ms, TE1 = 3 ms, TE2A-D = 3/11.5/20/28.5 ms, TI1/TI2 = 670/3855 ms, $\alpha_1/\alpha_2 = 7/6/C_{14}$. Number of lines/shots per readout: 150, number of readouts: 147. Scan duration: 16:30. Dead time: 8%.
- **MP2RAGE**: TRMP2RAGE = 6 s, TR = 6.2 ms, TE = 3 ms, TI1/TI2 = 1,000 ms/3200 ms, $\alpha_1/\alpha_2 = 7/6/C_{14}$. Number of lines/shots per readout: 150, number of readouts: 147. Scan duration: 14:45. Dead time: 52%.
- **ME-GRE**: TR = 31.4 ms, $\alpha = 12^\circ/C_{14}$, TEA,D = 3/11.5/20/28.5 ms. Scan duration: 11:35. Dead time: 0%; and
- **$B_1^+$-map**: DREAM sequence (Nehrke & Börnert, 2012), voxel size = 8.5 mm$^3$, flip angle 60$^\circ$.

### 2.3 Image processing and quantification

$T_2^*$ maps were obtained from the ME-GRE and MP2RAGEME data using a single-exponential fit. QSMs were obtained and averaged from TE2-4 of the ME-GRE and MP2RAGEME data with respect to cerebrospinal fluid (CSF) using STI Suite (Liu, Li, Tong, Yeom, & Kuzminski, 2015). Here, Laplacian-based phase unwrapping was performed (Li, Avram, Wu, Xiao, & Liu, 2014). A brain mask was obtained from the first echo TE1 magnitude image using SPM8 (Ashburner & Friston, 2005) and eroded by five voxels to remove veins, CSF and regions with low SNR due to susceptibility-induced intravoxel dephasing. The resulting susceptibility maps were normalized to zero with respect to the whole brain average.

$T_1$-weighted images were corrected for residual transmit $B_1^+$ inhomogeneities using the separately acquired low-resolution $B_1^+$-map following (Marques & Gruetter, 2013). Corrected MP2RAGE and MP2RAGEME data were subsequently processed using a dedicated MP2RAGE segmentation pipeline, including skull stripping (Bazin et al., 2014), whole brain segmentation with the multigeometric deformable model (MGDM) algorithm (Bazin et al., 2014; Bogovic, Prince, & Bazin, 2013), cortical reconstruction with the CRUISE algorithm (Han et al., 2004), volume-preserving cortical depth estimation (Waehnert et al., 2014), and vascular segmentation (Bazin, Plessis, Fan, Villringer, & Gauthier, 2016) to obtain cortical surface reconstructions at three different cortical depths, subcortical and cerebellar regions, and the arterial and venous vasculature around the brain. To explore the cortical parcellations that can be derived from MP2RAGEME-data, for a single subject, midcortical maps of $T_1$, $T_2^*$, and QSM were z-scored parallel to the cortex, allowing for visual inspection of the relative contrast. For quantitative comparison, average distances between the MP2RAGE and MP2RAGEME-based cortical surfaces over all cortical regions (left and right cerebrum, cerebellum) were computed per subject.

From the MGDM segmentation, median $T_1$, $T_2^*$, and QSM values were obtained from the following regions of interest (ROIs): WM (all), nucleus caudate, putamen, thalamus, cortical gray matter (all). For the
red nucleus, substantia nigra and subthalamic nucleus (STN), ROIs were defined by coregistration of an atlas, using the maximum probability labels (Keuken et al., 2017), that were thresholded at 10%. The image intensity distributions were subsequently visualized using histograms averaged over all subjects in MATLAB (The MathWorks, Inc., Natick, MA). To assess reproducibility, correlation plots were made and the Pearson correlation coefficient $r^2$ was calculated. Bland–Altman plots were generated (Bland & Altman, 1999), and reproducibility coefficients and coefficients of variation were computed, with a Kolmogorov–Smirnov (KS) test on non-Gaussianity of the difference data.

2.4 | Simulations

The sensitivity to $B_1^+$ inhomogeneity was simulated by numerically calculating signal intensities using the Bloch equations for both MP2RAGE and MP2RAGEME protocols (Marques et al., 2010). Contrast curves were simulated for $B_1$ values of 0.8, 1.0, and 1.2 times the nominal $B_1$ value. Also, the amount of $T_1$-induced blurring across the slice direction was simulated for both acquisition protocols. An artificial image was made containing a square of $21 \times 21$ gray matter voxels ($T_1 = 1.85$ s) in a WM ($T_1 = 1.15$ s) or CSF ($T_1 = 4$ s) background of $150 \times 150$ voxels, in order to be able to simulate the encoding used in Experiment 1. The signal in each compartment at each excitation during the acquisition of the first and second inversion readouts was computed using Bloch equations. All these images (two inversions times number of phase encoding steps per readout block) were Fourier transformed. Synthetic k-space data were created where each k-space encoding line was obtained from the k-space associated with its actual inversion time. Finally, these were inverse Fourier transformed and combined using the MP2RAGE image combination. The amount of blurring was visualized by comparing the readout direction profile with the slice-direction profile of the center one-dimensional image profile.

3 | RESULTS

For this 0.64 mm resolution protocol, MP2RAGEME offers a 40% time saving over separate scans (16 instead of 26 min) as the ME-GRE is acquired in the empty time of the TRMP2RAGE. The dead time of the MP2RAGE protocol was approximately half the scan time, while that of the MP2RAGEME protocol was only 6%.

The simulations showed that the $B_1^+$-sensitivity of the MP2RAGEME- and MP2RAGE-derived $T_1$-weighted images and $T_1$-maps were similar (Figure 2), though there are some differences for the different tissue types. For WM, a wider spread of image intensity values is seen in the MP2RAGEME protocol compared to the MP2RAGE protocol. Comparing 0.8 to 1.2 times the nominal $B_1^+$, the range of intensity values was 0.1 compared to 0.05 AU, and $T_1$ ranged from 0.9 to 1.4 versus 1.0 to 1.3, indicating more $B_1^+$-sensitivity. For GM, the image intensity ranges are comparable and for the CSF the range of intensities is narrower for the MP2RAGEME protocol, indicating equal and less $B_1^+$-sensitivity, respectively.

The simulations of the amount of $T_1$-induced blurring for both protocols are shown in Figure 3. As the $T_1$ of the MP2RAGEME is shorter than the $T_1$ of the MP2RAGE sequence, here is more $T_1$-evolution expected during the (equally long) readout, and, hence, increased blurring in the slice direction. The simulated slice profiles of the synthetic images show that there is indeed some blurring at the hard boundaries. The blurring is not visible in the complete profiles (Figure 3a,b), but in the zoomed panels in Figure 3b some effects are seen. The MP2RAGE protocol has minimal blurring at the WM/gray matter boundary, and shows moderate effects at the larger $T_1$-difference boundary between gray matter and CSF. In the MP2RAGEME differences in signal intensity of the voxels immediately neighboring the boundary can be observed at both frontiers. For both boundaries and acquisition protocols, the contrast-to-noise ratio (CNR) necessary to observe such blurring is higher than what is typically achieved in 0.6 mm acquisitions. The CNR for GM/WM was measured to be 10 in one data set, while the blurring induced was less than 10% of the difference between these two tissues.

Regarding the contrast between tissue types, for MP2RAGE-ME, the GM/WM contrast is maintained. The GM/CSF contrast, that was relatively high in the MP2RAGE sequence, is reduced by 40% to approximately the level of the GM/WM contrast.

The image quality overall was excellent for the derived $T_1$-weighted images, as well as for the quantitative $T_1$, $T_2^*$, and QSM maps. Figure 4 shows example slices of an MP2RAGEME data set at each orientation for all four derived images, including enlarged sections of specific ROIs in the axial plane. Because of the differences in the acquisition parameters, the $T_1$-weighted image intensity distributions differ significantly between the MP2RAGE and MP2RAGEME. Despite the different signal intensity distributions, the $T_1$-maps showed good agreement (see also Figure 5).

Table 1 summarizes the obtained quantitative $T_1$, $T_2^*$ and QSM values over all volunteers, in comparison with values obtained from the recent literature and from the separately acquired scans (see Figure S1 for an example of the studied ROIs). Most MP2RAGEME-obtained values were within one SD difference to the values obtained from the MP2RAGE and ME-GRE from the same subjects. For $T_2^*$, the substantia nigra and STN on average differed 1.5 and 1.6 ms, accounting to 10% of the mean reported values. For QSM, the most noticeable effect is the higher variability of 20 ppb in the red nucleus, substantia nigra, and STN. Here, literature values also vary more strongly. Other reported values reside within the range of values reported in literature.

Figure 5 shows histograms of the image intensities in the MP2RAGEME and separately acquired quantitative maps. Because of the differences in the acquisition parameters, the $T_1$-weighted image intensity distributions differ significantly between the MP2RAGE and MP2RAGEME (Figure 5a,d). In contrast, the $T_1$-maps generated with the appropriate lookup table and following $B_1^+$-correction are highly similar (Figure 5b,e). Both gray matter and WM peaks are slightly sharper for the MP2RAGE than for the MP2RAGEME. There are negligible differences in the $T_2^*$ values seen (Figure 5c,f).

Bland–Altman plots are displayed in Figure 6, with mean ROI values of all subjects, showing high $r^2$ values of 0.97, 0.94, and 0.90 for $T_1$, $T_2^*$, and QSM, respectively. Mean differences between sequences were nonsignificant for $T_2^*$ and $T_2^*$ ($p = .77; p = .94$) while a small significant difference for QSM was observed ($p = .01, \Delta T = -4.4$ ppb).
No non-Gaussian data distributions were observed, since KS p-values were .149, .261, and .090 for T1, T2*, and QSM, respectively. Repeatability coefficients reflecting the absolute difference’s 95% confidence interval were 99 ms, 3.0 ms, and 23 ppb for T1, T2*, and QSM, the latter being heightened by variable findings in deep brain nuclei, possibly caused by partial voluming or coregistration errors. Also, susceptibility maps are less robust and sensitive to streaking artifacts that can be due to the ill-posed nature of the problem or originating from phase errors in vessels. The CV was low in T1 (3.3%) and T2* (6.6%), and higher in QSM (33%), again because of variable results in the deep nuclei. For T1,

![Image 1](https://example.com/image1.png)

**FIGURE 2** Sensitivity to B1*-inhomogeneity. The signal intensity curves of MP2RAGE depend on the local B1*. The simulated image intensities for the MP2RAGEME (left) and MP2RAGE (right) protocols used are compared here for different B1* values. The spread of image intensity values (x axis) for the white matter (shaded area, continuous line) is larger in the MP2RAGEME protocol for a B1-range of 0.8–1.2 times nominal B1. For GM (broken line) the image intensity ranges are comparable for the two protocols. For the cerebrospinal fluid (CSF, dotted line), the simulated range of intensities is wider for the MP2RAGE protocol. Note the slightly different ranges of image intensities along the x axis.

![Image 2](https://example.com/image2.png)

**FIGURE 3** T1-induced blurring. The signal development during the acquisition leads to a blurring in the final image. As most of the T1-evolution happens during the first inversion, this is the main culprit. And as the TI1 of the MP2RAGEME is shorter, there is more T1-evolution during the (equally long) readout. This leads to slightly increased blurring in the slice direction (SL, dashed lines), which is compared to the unaffected readout (RO) direction. Note that signal levels differ between the MP2RAGE and MP2RAGEME acquisitions and hence their signal profiles do not overlap. (a) Simulated image containing a square of 30 x 30 GM voxels in a WM background. (b) Simulated image containing a square of 30 x 30 GM voxels in a cerebrospinal fluid (CSF) background. The larger T1-difference between gray matter and CSF means this border is more affected than the WM/GM one [Color figure can be viewed at wileyonlinelibrary.com]
a small positive bias of 50 ms is visible but otherwise similar values were observed, as was the case for $T_2^*$ values throughout the brain. QSM values for caudate, putamen, and thalamus differed by 4 ppb.

Figure 7 shows for a single subject a mapping of $T_1$, $T_2^*$, and susceptibility values onto the cortical surfaces. The observed spatial patterns are similar for both sequences, showing, for example, lower $T_1$ values in the primary, auditory, and visual motor cortices for both sequences. Furthermore, distinct spatial patterns for the different parameters can be observed, for example, in transversal projections, showing lower $T_2^*$ values in posterior and lower susceptibility values in anterior parts of the cortex, respectively.

The average cortical boundary distances generated from the MP2RAGE and MP2RAGEME data differed over all subjects on average by $0.1 \pm 1.1$ mm (WM/GM) and $0.1 \pm 1.2$ mm (GM/CSF). There is thus negligible bias, that is, much smaller than the voxel size and intersubject variation.

Figure 8 shows a combined arterial and venous map, by taking the maximum intensity projections of the CBS tools (Cognitive and Brain Sciences tools) software generated vasculature maps. The proximity of the veins and arteries clearly shows the importance of perfect coregistration between the data sets these vessels are derived from. Differences in the vascular trees that are visible between both views.
are attributed to small motion leading to displacement between subsequently scanned series. Due to this, certain arteries or veins may or may not be included in the ROI used for intensity projection. Also, locally varying shading patterns may be caused by subject motion.

4 | DISCUSSION

We presented the MP2RAGEME sequence, which allows simultaneous measurement of T₁, T₂*, and QSM at high resolution. The sequence’s two inversion readouts were flexibly designed to have a single echo and ME, respectively, leading to a time-efficient sequence of 16 min, maintaining a relatively long TE for good T₂* and QSM contrasts. The reconstructed parameter maps are naturally aligned, allowing for cortical reconstructions and providing parameter values that are in range with reported literature values.

While the optimal parameters for maximum GM/WM and CSF/GM contrast in MP2RAGE can be simulated using the Bloch equations (Marques et al., 2010), the number of parameters to be set is higher in the MP2RAGEME sequence, complicating these simulations. Moreover, as there are multiple contrasts-of-interest, the target of such simulations is also not clear (Metere et al., 2017). Hence, we chose sequence parameters for the MP2RAGEME to deliver: (1) optimal contrast in the T₂* map and QSM data, (2) optimal GM/WM contrast in the first inversion image, and (3) a maximal scan time of 17 min. Further optimization could be achieved by employing a discrete-time dynamic system to model spin dynamics (Zhao et al., 2018).

TI₁ was chosen to match the TI of a brain-stem specific protocol (670 ms, (Tourdias, Saranathan, Levesque, Su, & Rutt, 2014)), to facilitate midbrain segmentation using multiple-contrast data (Bazin et al., 2014). Because the second inversion image cannot overlap in time with the first, the minimum possible value for TI₂ became nearly 4 ss. The longer TI₂ reduces somewhat the T₁-sensitivity of the T₁-weighted images.

The flip angles in the MP2RAGEME of 7 and 6° were set for optimal T₁ contrast and minimal B₁+-sensitivity of the T₁-weighted images but are lower than the Ernst angle of 12° that was used in the ME-GRE. This implies that the MP2RAGEME was acquired with a lower SNR than the ME-GRE, calculated to be 25% lower in WM and 12% lower in GM, respectively. The impact on, for example, manual delineation of deep brain nuclei is thus small.

The shorter TI₁ resulted in minimally higher B₁+-sensitivity in the MP2RAGEME T₁-map (Figure 2), which was negligible in both...
| T1 (s) | (1) | (2) | (3) | (4) | (5) | (6) | T1 (s) | MP2RAGE | MP2RAGEME |
|-------|-----|-----|-----|-----|-----|-----|--------|---------|-----------|
| WM    | 1.15 ± 0.06 | 1.22 ± 0.03 | 1.13 ± 0.10 | 1.22 ± 0.03 | 1.1–14 | WM | 1.22 ± 0.01 | 1.19 ± 0.02 |
| CN    | 1.63 ± 0.09 | 1.75 ± 0.06 | 1.68 ± 0.07 | 1.44 ± 0.03 | 1.68 ± 0.06 | 1.6–17 | CN | 1.77 ± 0.05 | 1.80 ± 0.08 |
| Putamen | 1.52 ± 0.09 | 1.70 ± 0.07 | 1.64 ± 0.16 | 1.66 ± 0.04 | 1.5–17 | Putamen | 1.63 ± 0.04 | 1.63 ± 0.05 |
| Thalamus | 1.43 ± 0.10 | 1.70 ± 0.04 | 1.85 ± 0.04 | 1.9–21 | Thalamus | 1.59 ± 0.04 | 1.58 ± 0.04 |
| GM    | 1.97 ± 0.15 | 1.87 ± 0.17 | 2.13 ± 0.10 | 1.93 ± 0.15 | 1.80 ± 0.05 | 1.9–21 | GM | 1.95 ± 0.04 | 2.00 ± 0.08 |
| RN    | 1.23 ± 0.04 | 1.26 ± 0.05 | 1.35 ± 0.04 | 1.33 ± 0.07 |
| SN    | 1.63 ± 0.09 | 1.75 ± 0.06 | 1.68 ± 0.07 | 1.44 ± 0.03 | 1.68 ± 0.06 | 1.7 ± 0.08 | 1.68 ± 0.06 | 1.7 ± 0.08 |
| STN   | 1.19 ± 0.04 | 1.22 ± 0.05 | 1.35 ± 0.04 | 1.28 ± 0.07 |
| T2* (ms) | (7) | (8) | (9) | (4) | (5) | (6) | T2* (ms) | GRE | MP2RAGEME |
| WM    | 26.8 ± 1.2 | 25.9 ± 0.8 | 24–27 | WM | 27.1 ± 0.4 | 26.7 ± 0.1 |
| CN    | 19.9 ± 2.0 | 27.4 ± 1.1 | 28.0 ± 1.9 | 25 | CN | 27.5 ± 1.5 | 27.9 ± 0.5 |
| Putamen | 16.1 ± 1.6 | 33.2 ± 1.9 | 20–26 | Putamen | 23.9 ± 1.9 | 24.1 ± 0.6 |
| Thalamus | 30.7 ± 1.9 | 37.8 ± 1.8 | 38.9 ± 1.9 | 39.0 ± 1.9 |
| GM    | 33.2 ± 1.3 | 32.20 ± 1.35 | 32.2 ± 1.45 | 23.7 ± 1.8 | 23.0 ± 1.9 | 25–33 | GM | 32.8 ± 1.7 | 32.4 ± 0.7 |
| RN    | 17.3 ± 1.5 | 18 | RN | 17.3 ± 3.0 | 16.7 ± 2.4 |
| SN    | 13.2 ± 1.0 | 15.6 ± 1.7 | 14.1 ± 0.9 |
| STN   | 15.1 ± 1.5 | 6.0–18 | SN | 14.8 ± 1.3 | 16.4 ± 1.4 |
| χ (ppb) | (10) | (11) | (12) | (13) | (5) | (6) | χ (ppb) | GRE | MP2RAGEME |
| WM    | 40 ± 15 | 44 ± 17 | 60 | CN | 32 ± 5 | 26 ± 2 |
| Putamen | 25 ± 20 | 38 ± 17 | 20–70 | Putamen | 29 ± 8 | 27 ± 1 |
| Thalamus | 8 ± 10 | 45 ± 20 | 13 ± 5 | Thalamus | 10 ± 2 |
| GM    | 65 ± 15 | 100 ± 19 | 28 ± 36 | 80–110 | RN | 74 ± 24 | 68 ± 22 |
| SN    | 30 ± 10 | 152 ± 30 | 95 ± 34 | SN | 77 ± 22 | 75 ± 11 |
| STN   | 111 ± 27 | 27 ± 43 | 80–135 | STN | 80 ± 22 | 65 ± 20 |

CN = caudate nucleus; GM = gray matter; ME-GRE = multi-echo gradient echo; RN = red nucleus; ROIs = regions of interest; SN = substantia nigra; STN = subthalamic nucleus; WM = white matter. References: 1: (Marques et al., 2010), 2: (Rooney et al., 2007), 3: (Wright et al., 2008), 4: (Metere et al., 2017), 5: (Keuken et al., 2017), (Marques & Norris, 2017), 7: (Peters et al., 2007), 8: (Cohen-Adad et al., 2012), 9: (Govindarajan et al., 2015), 10: (Sood et al., 2017), 11: (Langkammer et al., 2012), 12: (Khabipova et al., 2014), 13: (Dietz et al., 2013b).
MP2RAGE and MP2RAGEME after correction using the DREAM $B_1^+$-map (Figure 5). Good $B_1^+$-inhomogeneity correction of both MP2RAGE and MP2RAGEME data is essential for successful cortical segmentation using automated routines (Haast, Ivanov, & Uludağ, 2018).

Both readouts of the MP2RAGE and MP2RAGEME sequences are long relative to the $T_1$-relaxation, meaning that there is significant relaxation during the acquisition of especially the first inversion image. This $T_1$-decay leads to a broadening of the point spread function in the slice encoding direction (Deichmann, Hahn, & Haase, 1999), though this is only a small effect for the protocols compared here (Figure 3).

For the iron-rich nuclei of the subcortex and cerebellum, the exact colocalization of the three contrasts can help delineate more precise boundaries across the different contrasts, for example, for the globus pallidus or the STN. For the vasculature, a precise

\[
\text{FIGURE 6} \quad \text{Bland-Altman plots. Correlation and difference plots for (a) } T_1, (b) T_2^*, \text{ and (c) QSM, for white matter (WM), caudate (Caud), putamen (Put), thalamus (Thal), gray matter (GM), red nucleus (RN), substantia nigra (SN), and subthalamic nucleus (STN). The statistics Pearson's correlation coefficient ($r^2$), reproducibility coefficient (RPC), coefficient of variation (CV), and a Kolmogorov–Smirnov (KS) test of non-Gaussianity of differences are also given. Mean regions of interest values for all six subjects are plotted. Fitted (solid) and identity (dashed) lines are given in the correlation plots (left), and mean difference (solid) and 95% confidence interval (dashed) lines (right) are depicted [Color figure can be viewed at wileyonlinelibrary.com]}
\]
Colocalization of the arterial (in T1 maps) and venous (in T2* and QSM maps) vessels was crucial, as these thin and elongated structures often run side by side.

For all ROIs, the T1, T2*, and QSM values derived from the MP2RAGEME or separately acquired MP2RAGE and ME-GRE are similar. The observed variations are small compared to the spread in literature values. When comparing the quantitative values in Table 1 with the literature values, the only noticeable differences are found for the values reported in the Caudate nucleus, with somewhat longer T1 values and lower QSM values found here than in previous work. Other work also noted that the Caudate nucleus and Putamen are difficult to distinguish (Keuken et al., 2014). This might be due to partial inclusion of the CSF in the neighboring ventricles. The relatively large SD in QSM values in Table 1, both for the MP2RAGEME and ME-GRE data, are also observed in the literature references. Still, the susceptibility values obtained from the MP2RAGEME data were similar to those obtained from the separately acquired ME-GRE data.

Other ROIs were also possibly affected by imperfect segmentation: The red nucleus T2* values were shorter than expected from literature (Table 1), possibly due to segmentation errors. Since the WM mask contained subcortical structures, reported QSM values in WM are higher than in the literature. This is also visible in Figure 5f as a small “shoulder” of low-T2* values in the histogram.

The MP2RAGEME sequence is relatively time efficient, with a dead time of only 6%. Since only the second inversion was extended to a ME readout, the first inversion time, which defines the T1-contrast in the MP2RAGE image, can be chosen optimally and total scanning time is shortened compared to an ME readout of both inversions (Metere et al., 2017).

The use of less B1+ inhomogeneity sensitive radio frequency pulses for the inversion, such as the FOCI pulse, would improve the extent of brain area with homogeneous contrast (O’Brien et al., 2014; O’Brien et al., 2014). The use of SPINS pulses for the echo trains would also yield more homogeneous images, limiting the variation in

**FIGURE 7** Comparison of the cortical contrasts. Maps of mid-cortical T1, T2*, and QSM variations obtained for a single subject (as in Figure 1) with the separate MP2RAGE and GREME or the combined MP2RAGEME approach. Cortical depth was estimated with volumetric layering and values were smoothed along the mid-cortical depth with a 0.64 mm FWHM Gaussian kernel. Surface reconstructions were obtained from the underlying T1 maps. To compare local patterns all maps were z-scored parallel to the cortex based on median and interquartile range. Locations of primary motor, auditory and visual cortices are indicated in T1 maps, as well as lower QSM values in the anterior part of the cortex [Color figure can be viewed at wileyonlinelibrary.com]

**FIGURE 8** Arterial and venous vasculature. Vasculature reconstructed from the separate MP2RAGE and GREME or the MP2RAGEME-sequence data (maximum intensity projections over 20 slices in axial and sagittal directions, colored in red for structures extracted from T1 maps (arteries) and blue for structures extracted from 1/T2* maps (veins). Note the tight interaction of arteries and veins locations, making precise coregistration of the contrast particularly important. Images are for the same single subject as in Figure 1 [Color figure can be viewed at wileyonlinelibrary.com]
the ME-GRE readout due to imperfect flip angles (Malik, Keihaninejad, Hammers, & Hajnal, 2012). Nevertheless, improved B1+-homogeneity would benefit these images and, hence, the use of parallel transmission would also be advantageous. On the other hand, if sufficient B1+- can be achieved throughout the brain, variable flip angle imaging might again show improved performance with minimal point spread function (PSF) blurring.

Subject motion was limited in the experiments performed in this article and did not lead to noticeable image degradation. Only in minimum and maximum intensity projections (Figure 7), locally varying shading patterns can be seen that may be attributed to motion. However, the long, high resolution acquisitions are susceptible to motion artifacts, especially in possible applications to study disease or large populations. Interleaving the acquisition with fat image navigators (Gallichan & Marques, 2017) or using real-time field control (Özbay, Duerst, Wilm, Puessmann, & Nanz, 2017) would be candidate approaches to correct for motion artifacts either retrospectively or prospectively.

The proposed algorithm is limited by a longer TR compared to the MP2RAGE sequence, which reduced the CNR between CSF and GM (Figure 3), however, to an acceptable level comparable to the GM/WM contrast. The reduced sequence dead time was accounted for in the signal model, that is, Tc in Equation (A1), and thus does not affect T1-quantification. Because of more and longer gradient switching, more heating of the system and consequential resonance frequency drift may occur.

The specific absorption rate (SAR) and a possible raise thereof is of limited concern in MP2RAGE and MP2RAGEME sequences. Because of the low flip angle readout trains, and the inversion pulses being far apart, SAR levels are low for both sequences. If anything the SAR per unit of time is reduced in the MP2RAGEME due to the increased repetition time. This is in strong contrast to, for example, TSE readouts, where much higher flip angles (>100°) are repetitively applied or 2D imaging with simultaneous multi-slice excitation.

5 | CONCLUSION

We show that quantitative T1, T2*, and QSM information can be acquired in a single acquisition. Furthermore, we show that the resulting quantitative values are comparable to current separately acquired sequences. Our MP2RAGEME sequence has the benefit in large time savings, perfectly coregistered data and minor image quality differences.

ACKNOWLEDGMENT

M.W.A.C. is the stock owner of Nico-lab International Ltd.

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How to cite this article: Caan MWA, Bazin P-L, Marques JP, de Hollander G, Dumoulin SO, van der Zwaag W. MP2RA-GEEM: T1, T2, and QSM mapping in one sequence at 7 tesla. Hum Brain Mapp. 2019;40:1786–1798. https://doi.org/10.1002/hbm.24490
APPENDIX

The signal model for the MP2RAGEME sequence for given inversion, echo and repetition times can be written as:

\[
S(T_{I1}, T_{E1}) = M_0 \exp \left( -\frac{T_{E1}}{T_2} \right) \sin \alpha_1 \cdot \left\{ \frac{-\eta M_0^{ss} E_A + (1 - E_A)}{M_0^2} \left( E_1 \cos \alpha_1 \right)^{1 - 1} + (1 - E_1) \frac{1 - (E_1 \cos \alpha_1)^{1 - 1}}{1 - E_1 \cos \alpha_1} \right\}
\]  
(A1)

and

\[
S(T_{I2}, T_{E2}) = M_0 \exp \left( -\frac{T_{E2}}{T_2} \right) \sin \alpha_2 \cdot \left\{ \frac{M_0^{ss} E_C^{1 + 1} + (1 - E_C^{1 + 1})}{M_0^2} \left( E_2 \cos \alpha_2 \right)^{1 - 1} + (1 - E_2) \frac{1 - (E_2 \cos \alpha_2)^{1 - 1}}{1 - E_2 \cos \alpha_2} \right\}
\]  
(A2)

with TE_{2i} one of the multiple echo times at the second inversion readout only, M_0 the net magnetization, E_j ≡ \exp(-T_j/T_1) and E_A ≡ \exp(-T_A/T_1) and E_C ≡ \exp(-T_C/T_1). η = \frac{1}{2} \left[ 1 - M_0(0^{-})/M_0(0^{+}) \right] equals the inversion of the adiabatic inversion pulse, with M_0(0^{-}) and M_0(0^{+}) the longitudinal magnetization directly before and after the pulse (Mildner et al., 2014).

M_0^{ss} is the steady-state longitudinal magnetization (Marques et al., 2010). After introducing a temporary variable x

\[
x = \left( (1 - E_A) (E_1 \cos \alpha_1)^{n} + (1 - E_1) \frac{1 - (E_1 \cos \alpha_1)^{1 - 1}}{E_1 \cos \alpha_1} \right) E_B + (1 - E_B) (E_2 \cos \alpha_2)^{n}
\]  
(A3)

we can write for M_2^{ss}:

\[
M_2^{ss} = \frac{M_0 \left[ x + (1 - E_2) \frac{1 - (E_2 \cos \alpha_2)^{1 - 1}}{E_2 \cos \alpha_2} \right]}{1 + \eta \left( \cos \alpha_1 \cos \alpha_2 \right)^{n}} \exp(-TR_{\text{MP2RAGEME}}/T_1)
\]  
(A4)

The complex-valued first echo images of both inversion readouts are combined to remove B_0, M_0, and B_1^-effects, by computing the real part \(\Re[S]\) of the normatively averaged signals and \(S^*\) denoting the complex conjugate of S:

\[
S_M = \Re \left[ \frac{S^*(T_{I1}, T_{E1}) \cdot S(T_{I2}, T_{E2})}{[S(T_{I1}, T_{E1})]^2 + |S(T_{I2}, T_{E2})|^2]} \right].
\]  
(A5)

T_1 values were computed from \(S_M\) through a lookup table with sequence parameter values (Marques et al., 2010).