Harmonization of Quantitative Parenchymal Enhancement in T1-Weighted Breast MRI

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Background: Differences in imaging parameters influence computer-extracted parenchymal enhancement measures from breast MRI.

Purpose: To investigate the effect of differences in dynamic contrast-enhanced MRI acquisition parameter settings on quantitative parenchymal enhancement of the breast, and to evaluate harmonization of contrast-enhancement values with respect to flip angle and repetition time.

Study Type: Retrospective.

Phantom/Populations: We modeled parenchymal enhancement using simulations, a phantom, and two cohorts (N = 398 and N = 302) from independent cancer centers.

Sequence Field/Strength: 1.5T dynamic contrast-enhanced T1-weighted spoiled gradient echo MRI. Vendors: Philips, Siemens, General Electric Medical Systems.

Assessment: We assessed harmonization of parenchymal enhancement in simulations and phantom by varying the MR parameters that influence the amount of T1-weighting: flip angle (8°–25°) and repetition time (4–12 msec). We calculated the median and interquartile range (IQR) of the enhancement values before and after harmonization. In vivo, we assessed overlap of quantitative parenchymal enhancement in the cohorts before and after harmonization using kernel density estimations. Cohort 1 was scanned with flip angle 20° and repetition time 8 msec; cohort 2 with flip angle 10° and repetition time 6 msec.

Statistical Tests: Paired Wilcoxon signed-rank test of bootstrapped kernel density estimations.

Results: Before harmonization, simulated enhancement values had a median (IQR) of 0.46 (0.34–0.49). After harmonization, the IQR was reduced: median (IQR): 0.44 (0.44–0.45). In the phantom, the IQR also decreased, median (IQR): 0.96 (0.59–1.22) before harmonization, 0.96 (0.91–1.02) after harmonization. Harmonization yielded significantly (P < 0.001) better overlap in parenchymal enhancement between the cohorts: median (IQR) was 0.46 (0.37–0.58) for cohort 1 vs. 0.37 (0.30–0.44) for cohort 2 before harmonization (57% overlap); and 0.35 (0.28–0.43) vs. 0.37 (0.30–0.44) after harmonization (85% overlap).

Data Conclusion: The proposed practical harmonization method enables an accurate comparison between patients scanned with differences in imaging parameters.

Level of Evidence: 3
Technical Efficacy Stage: 4

Dynamic contrast-enhanced magnetic resonance imaging (MRI) is the principal sequence in breast MRI for assessment of breast cancer. Healthy breast parenchymal tissue also demonstrates contrast-enhancement on MRI. This background parenchymal enhancement (BPE) is increasingly used in the context of cancer diagnosis, prognosis, and risk assessment.2,3
Assessment of breast MRI follows guidelines from the Breast Imaging Reporting and Data System (BI-RADS). These guidelines address standardization of image acquisition parameters such as the timing of the postcontrast images. They do not, however, address imaging parameter settings such as the values of the flip angle and repetition time (TR). These parameters influence the signal intensity in spoiled gradient echo imaging, the most commonly used imaging technique in breast dynamic contrast-enhanced MRI. Therefore, differences in parameters will influence computer-extracted measures from these images.

Recently, several researchers suggested computer-extracted measures to quantify BPE. These measurements are inherently sensitive to differences in imaging parameter settings, complicating pooling of results across studies. For example, two studies reporting an association between parenchymal enhancement of the contralateral breast and patient survival used cohort-specific data-driven cutoffs. In a recent prospective multicenter study, computer-extracted biomarkers based on contrast enhancement needed to be manually adjusted to account for variability in the MRI system and scan protocol. Since it is difficult to prospectively account for all variation in multicenter studies, post-hoc harmonization of enhancement biomarkers is of interest (ie, the process of achieving a consistent value regardless of imaging protocol).

Taking the above into account, we aimed to investigate the effect of differences in MRI acquisition parameter settings on quantitative parenchymal enhancement. Furthermore, we evaluated a harmonization method to adjust the differences in enhancement caused by differences in parameters.

**Material and Methods**

**Simulation**

Contralateral parenchymal enhancement (CPE) is defined as the top 10% relative signal-enhancement between the first and the last postcontrast scan in dynamic contrast-enhanced MRI (Fig. 1). We simulated the image intensities of breast parenchymal tissue at three timepoints in a contrast-enhanced series: precontrast, first postcontrast (eg, 90 seconds after contrast injection), and last postcontrast (eg, 360 seconds after contrast injection) (Fig. 1).

We simulated the parenchymal tissue at known quantitative T1 values at 1.5T. We modeled the signal of the precontrast parenchyma using 1000 voxel values with a T1 value of 1246 msec. We used a range of T1 values in the postcontrast simulations since the parenchyma typically shows heterogeneity of signal enhancement in response to contrast influx.

We chose a T1 ranging from 959–1149 msec for the first postcontrast series, and 597–892 msec for the last postcontrast series. The ranges decrease over time because the concentration of gadolinium after contrast typically continues to increase in parenchymal tissue (Fig. 1).

We simulated the signal using the steady-state spoiled gradient echo signal equation, and added Gaussian noise to the signal with a signal-to-noise ratio (SNR) comparable to our phantom measurements (ie, 25, 28, and 45 for pre-, first post-, and last postcontrast, respectively). We simulated this signal using a range of imaging parameter settings resembling those used in clinical protocols: TRs between 4 and 12 msec and flip angles between 8° and 25°. Proton density and TE were kept constant.

We assessed the effect of SNR on the ability to accurately correct parenchymal enhancement. For this, we simulated the precontrast and postcontrast signals with increasing noise levels, and recorded at which noise level the error between the harmonized CPE and the reference CPE (ie, flip angle 10°, TR 4 msec) became higher than 10%.

Furthermore, we assessed the effect of B1+ field inhomogeneities using a similar method, by recording at what field inhomogeneity the error between the harmonized CPE and the reference CPE (ie, flip angle 10°, TR 4 msec) became higher than 10%. We simulated the inhomogeneities using both a bias and deviations from the nominal flip angle. The bias of the simulated B1+ field inhomogeneities was set at 120%, while ranging the absolute deviations from this bias between 0% and 100%. Hence, a B1+ inhomogeneity of 40% refers to a B1+ field ranging between 80% and 160%, with the average at 120%.

**Phantom**

We measured parenchymal enhancement using a calibrated phantom containing a series of materials with known relaxation times (Test object 5, Eurospin II Test System, Diagnostic Sonar, Edinburgh, UK).
The phantom included a set of 18 calibrated test tubes. Each tube is filled with doped polysaccharide gel and provides specific relaxation times at specified temperatures and magnetic field strengths. We defined three signals using these tubes: one signal consisted of a “precontrast tube” mimicking parenchymal tissue without contrast, and two consisted of “postcontrast tubes” mimicking the gradual decrease of T₁ (Fig. 1). We chose the tube calibrated at a T₁ of 1246 msec as the precontrast tube. We chose the tubes calibrated at a T₁ of 1149, 1023, and 959 msec as the first postcontrast tubes, and the tubes at a T₁ of 892, 745, and 597 as the last postcontrast tubes.

We approximated dynamic contrast-enhanced MRI in the phantom using different tubes at known T₁ values. Hence, our timepoints (ie, precontrast, first postcontrast, and last postcontrast) were actually different locations in the MR image. These spatial variations may influence the measurements because of differences in the B₁⁺ field and may therefore influence the adjustment of parenchymal enhancement. We showed the effect of these spatial variations by using the actual flip angle in our harmonization, which was the flip angle multiplied by the scaling of the B₁⁺ field at that location.

**MRI**

We imaged the phantom using a 1.5T MR unit (Achieva, Philips, Best, The Netherlands). We used a dedicated breast MR coil (SENSE 7 Breast coil, Philips) and performed spoiled gradient echo imaging using the same range of clinically used imaging parameter as in the simulation.¹⁸,²³–²⁵ Other imaging parameters were: echo time (TE) 1.7 msec, voxel size 1 × 1 × 1 mm³, and image dimension 224 × 224 × 50 voxels. We assessed the SNR of the MR image by measuring the noise of the phantom using a scan without gradients and radiofrequency pulses at TR of 4 msec and flip angle of 10°.²⁷ We acquired a B₁⁺-map to assess the actual flip angle (actual flip angle imaging, TRs 30 and 150 msec, TE 2.4 msec, flip angle 60°, resolution 2 × 2 × 4 mm³).²⁸

**Image Analysis**

We automatically extracted the calibrated tubes from the MR image using a threshold based on Otsu’s method²⁹ and labeled them with a connected component analysis, yielding a binary mask per tube. We eroded the binary masks (radius of two voxels) to account for partial volume effects along the borders.

To assess parenchymal enhancement, we used the previously defined signal-tubes: the precontrast tube, the first postcontrast tubes, and the last postcontrast tubes. From the latter tubes, we randomly sampled the same number of voxels (ie, 5201 voxels) yielding a total of 15,603 voxel values from which we calculated the CPE.

**Harmonization of Parenchymal Enhancement**

We harmonized parenchymal enhancement for differences in TR and flip angle based on the method proposed by Haacke et al.²² First, we assessed the T₁ value of each voxel in the parenchyma at the postcontrast image²²:

\[
T₁(t) = \frac{-TR}{\ln(x - y)}
\]

with:

\[
\begin{align*}
  y &= \frac{S(t)}{S(0)} \\
  x &= 1 - e^{-\frac{TR}{T₁(0)}}
\end{align*}
\]

where S(0) is the signal of the precontrast image, S(t) the signal at the postcontrast image at timepoint t, T₁(0) the T₁ value in without contrast agent, TR the repetition time, and α the flip angle.

After obtaining T₁(t), we calculated the simulated signal \(\hat{S}\) in each voxel of the parenchymal tissue as if they were imaged with different repetition time (TRnew) and flip angle (αnew):

\[
\hat{S}(t, α_{new}, TR_{new}) = \frac{\sinα_{new} \left( 1 - e^{-\frac{TR_{new}}{T₁(0)}} \right)}{1 - \cosα_{new} \cdot e^{-\frac{TR_{new}}{T₁(0)}}}
\]

We chose a TR of 4 msec and flip angle 10°.

We performed these steps for both the first postcontrast and the last postcontrast signal. From these new adjusted postcontrast signals, the adjusted parenchymal enhancement was calculated using the same CPE definition.

All harmonization steps were performed in Python 3.7.4.

**Clinical Data**

We evaluated CPE using two previously reported cohorts with a total of 696 patients.¹⁸,¹⁹ Institutional Review Board (IRB) approvals were obtained for the analyses of patient data¹⁸,¹⁹; written informed consent was obtained¹⁸ or waived by the IRB.¹⁹ Both cohorts solely included patients with unilateral estrogen receptor-positive/HER2-negative breast cancer who received a preoperative dynamic contrast-enhanced T₁-weighted MRI of both breasts. All patients were eligible for breast-conserving surgery based on conventional imaging and clinical examination (Table 1). CPE was calculated in parenchymal tissue voxels on dynamic contrast-enhanced MRI, which were automatically segmented in 3D (Fig. 2, see¹⁸,¹⁹ for implementation details) (Table 1).
Notable differences in MRI acquisition between these two patient cohorts were the discrepancies in vendor, flip angle, and TR. Cohort 1 was scanned on a Siemens MR unit with TR of 8.0 msec and flip angle of 20°, whereas cohort 2 was scanned on a General Electric Medical Systems (Milwaukee, WI) MR unit with TR of 6.0 msec and flip angle 10° (Table 2).

To assess whether our clinical data have the required SNR for harmonization, we calculated the SNR in the parenchymal tissue using manual annotations in 10 patients (Table 3). The signal was assessed in the parenchymal tissue of the first postcontrast image; the noise was assessed as a Rician distribution in the air.30

**Statistical Analysis**

We assessed the effect of harmonization on the simulations and phantom using the median and interquartile range (IQR). We tested the similarity of the CPE distributions in the two cohorts using the overlap of the kernel density estimations,31 and tested the improvement of overlap after harmonization using the paired Wilcoxon signed rank test after bootstrapping. We considered the results statistically significant when the two-sided P-value was under 0.05.

Statistical analyses were performed using R 3.6.1. (R Foundation for Statistical Computing, Vienna, Austria).

**Results**

**Simulation**

As shown in Fig. 3, the simulated parenchymal enhancement increased with increasing flip angle and decreased with increasing TR. Before harmonization, simulated parenchymal enhancement values had a median (IQR) value of 0.46 (0.34–0.49). After harmonization, the IQR reduced: median (IQR): 0.44 (0.44–0.45) (Fig. 3).

The error to the reference parenchymal enhancement value (flip angle 10°, TR 4 msec) after harmonization was lower than 10% in most clinically used parameter settings for commonly used SNRs (Fig. 4) and for clinically expected B1+ field inhomogeneities (Fig. 5).
Phantom

We observed similar behavior in the phantom as in the simulation: signal enhancement values increased with increasing flip angle and decreased with increasing TR (Fig. 6).

Before harmonization, signal enhancement values measured in the phantom had a median (IQR) of 0.96 (0.59–1.22). After harmonization, the IQR decreased: median (IQR): 0.91 (0.86–0.98) without mitigating $B_1^+$ inhomogeneities, median (IQR): 0.96 (0.91–1.02) with mitigating $B_1^+$ inhomogeneities.

The SNR was 25 in the precontrast tube, 28 in the first postcontrast tubes, and 46 in the last postcontrast tubes. The median actual flip angle was above 90% and below 110% in all tubes.

Clinical Data

Consistent with the simulation and phantom, CPE was higher at a flip angle of $20^\circ$ and TR of 8 msec (cohort 1, median [IQR]: 0.46 [0.37–0.58]) compared to using flip angle of $10^\circ$ and TR of 6 msec (cohort 2, median [IQR]: 0.37 [0.30–0.44], Fig. 7). After harmonization of cohort 1 to the parameter settings of cohort 2, the distribution in cohort 1 better resembled cohort 2 (median [IQR]: 0.35 [0.28–0.43], Fig. 7). The overlap of the kernel density estimations significantly increased from 57% (95% confidence interval [CI]: 49–63%) to 85% (95% CI: 73–89%) after harmonization ($P < 0.001$). The mean SNR that we measured in the parenchymal tissue of patients was 15.9 with a standard deviation (SD) of 2.7 (Table 3).

Discussion

We demonstrated that harmonization of parenchymal enhancement values is needed to ensure an accurate comparison between patients scanned with different dynamic contrast-enhanced MRI acquisition parameter settings. We proposed a harmonization method to adjust differences in parenchymal enhancement caused by differences in these parameters. Additionally, we showed that the CPE observed

### TABLE 2. MRI Characteristics

| Characteristic       | Cohort 1 ($N=398$) | Cohort 2 ($N=302$) |
|----------------------|--------------------|--------------------|
| Field strength (T)   | 1.5                | 1.5                |
| Vendor               | Siemens            | General Electric Medical Systems |
| Sequence             | Spoiled gradient echo | Spoiled gradient echo |
| Repetition time (msec)| 8.0                | 6.0                |
| Echo time (msec)     | 4.0                | 4.2                |
| Flip angle ($^\circ$)| 20                 | 10                 |
| Contrast agent       | Prohance (Bracco-Byk Gulden) | Magnevist (Bayer Health Care Pharmaceuticals) |
| Injected dose (mmol/kg)| 0.1              | 0.1                |
| Duration of dynamic contrast-enhanced series (s) | 360 | 360 |
| Voxel size (mm$^3$)  | $1.35 \times 1.35 \times 1.35$ | $0.7 \times 0.7 \times 3.0$ |
| Matrix size          | $256 \times 256 \times 100$ | $256 \times 256 \times 100$ |
| Fat suppression      | No                 | Yes                |

### TABLE 3. Signal-to-Noise Ratios in 10 Patients From Cohort 1

| Patient Study ID | SNR  |
|-----------------|------|
| 1               | 15.4 |
| 2               | 11.2 |
| 3               | 13.5 |
| 4               | 15.1 |
| 5               | 13.8 |
| 6               | 18.3 |
| 7               | 21.4 |
| 8               | 15.0 |
| 9               | 17.1 |
| 10              | 17.7 |
| Mean (SD)       | 15.9 (2.7) |
in two patient cohorts was equivalent after harmonization. These results suggest that pooling of data in multi-institutional studies is feasible after correction of enhancement values.

Differences in image acquisition parameter settings influence computer-extracted measures of dynamic contrast-enhanced MRI. In addition to complicating pooling of data across studies, it also complicates correctly classifying prospectively acquired subject data, hindering research into personalized healthcare. Therefore, it is desirable to produce the same value for such biomarkers regardless of acquisition parameters. One way to achieve such harmonization is by strict guidelines or community-wide harmonization efforts. While these guidelines are important for established biomarkers, they are unfeasible for relatively novel biomarkers. Only more established biomarkers are likely to receive the required support from the scientific and clinical community to a priori harmonize relevant parameters between centers. Therefore, post-hoc harmonization of relatively novel biomarkers is of interest.
We provided such harmonization by adjusting the CPE biomarker.\textsuperscript{18,19,32,33} We showed that we were able to adjust this biomarker in our simulation with realistic parameter settings. Compared to the simulation, our phantom study showed slightly more variation after harmonization. This may be because the noise levels in our simulations were based on the SNR measurement from our phantom, which was assessed once at a flip angle of $10^\circ$ and TR of 4 msec. As a result, the noise varies between acquisition settings in our phantom study, where it does not in our simulation. Another reason may be $T_2^*$-effects, which were not addressed by our simulations. To minimize this potential reason, we chose tubes in our phantom with similar $T_2$-values and we chose our TE as short as possible on our MR system.

In our clinical data, the improved similarity between CPE values in the cohorts after harmonization was in agreement with the simulations for the parameter settings we were adjusting given the SNR in our images. Therefore, even though the SNR of the MR acquisition is often not optimized for parenchymal tissue, harmonization with real patient cohort data still appears feasible.

The proposed harmonization method is relatively straightforward to implement using the provided description and the article from Haacke et al.\textsuperscript{22} Although we used seemingly straightforward MR physics, there are several aspects that potentially complicate harmonization of patient data. These aspects include postprocessing on the scanner by vendor software, noise, patient motion, and biological patient

FIGURE 6: Parenchymal enhancement measured in the phantom at various flip angles and TRs before (left) and after harmonization (right). The reference situation was at flip angle of $10^\circ$ and TR of 4 msec. The median (IQR) of the measurements was 0.96 (0.59–1.22) before harmonization (left figure) and 0.96 (0.91–1.02) after harmonization and mitigation of $B_1^+$ inhomogeneities (right figure). The colors are based on the percentages compared to the reference comparable to Fig. 3.

FIGURE 7: The overlap of the kernel density estimations between contralateral parenchymal enhancement in cohort 1 ($n=394$, blue) and cohort 2 ($n=302$, orange) significantly increased after harmonization from 57% (left) to 85% (right) ($P < 0.001$).
variation. It is promising that, in spite of these potential complications, our results show a good harmonization between patient cohorts.

The duration of the dynamic contrast-enhanced series was comparable in the cohorts. Hence, our study did not investigate the effect of temporal resolution on enhancement features. This is subject to future research. International guidelines prescribe, however, limits to the range of timing in dynamic contrast-enhanced series.5,34

Different contrast agents have different relaxivity properties that might influence enhancement characteristics.35 The contrast agents used in our study (Prohance and Magnevist) have, however, similar relaxivity properties.35

We harmonized the parenchymal enhancement in the contralateral breast. Although we focused on the contralateral breast to exclude cancer-induced enhancement, the same methodology can be applied to the ipsilateral breast.

The effect of harmonization on other enhancement biomarkers, such as radiomics features extracted from the tumor,36–38 may also lend themselves to harmonization. In future research, we will investigate the effect of harmonization on these biomarkers.

We chose a model-based harmonization method. The alternative would be a data-driven approach. Such approaches exist; for example, by harmonizing given feature distributions,39 or by harmonizing MR images using deep learning, from which features can then be calculated.40 Our model-based approach may have several advantages over data-driven approaches. First, by using a model-based approach, one can anticipate the efficacy of the harmonization under different conditions. Second, data-driven approaches typically require much data. This can be a problem with the prospective use of such methods. For example, in the case of a new MR unit, one needs to gather enough data first in order to perform accurate harmonization.

Limitations

Our study has some limitations. First, we performed our analyses only at 1.5T. Our results may slightly vary at 3T, due to the slightly higher $T_1$ of parenchymal tissue at 3T and the decreased effect of contrast agent at 3T.20,35 However, since the overall behavior of spoiled gradient echo is similar at both field strengths, we expect our results to hold for 3T. Our study shows promising results at $B_1^+$ field inhomogeneities that are expected at 1.5T.26 These inhomogeneities are higher at higher field strengths.41 Vendors, however, have acknowledged this and updated their systems accordingly.42

We only gave indirect evidence for the efficacy of the harmonization method in patients. A more direct assessment of the harmonization method in vivo may be a test–retest measurement: scanning the same patient twice or more often with different flip angles and TRs. To fully elucidate the parenchymal enhancement, this would mean scanning the same patient on different days, since clearance of the contrast agent from the body takes ~1 day.43,44 Furthermore, even if all these steps would be taken, there are considerable physiological variations in breast parenchyma that occur from day to day,45,46 rendering it difficult to control such an experiment.

Conclusion

We showed promising results that harmonization of parenchymal enhancement values can enable accurate comparisons between patients scanned with differences in imaging parameters.

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