FULL PAPER

Improving the image quality of DWI in breast cancer: comparison of multi-shot DWI using multiplexed sensitivity encoding to conventional single-shot echo-planar imaging DWI

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Objective: To compare diffusion-weighted images (DWI) acquired using single-shot echo-planar imaging (ss-EPI) and multiplexed sensitivity encoding (MUSE) in breast cancer.

Methods 20 females with pathologically confirmed breast cancer (age 51 ± 12 years) were imaged with ss-EPI-DWI and MUSE-DWI. ADC, normalised ADC (nADC), blur and distortion metrics and qualitative image quality scores were compared. The Créte-Roffet and Mattes mutual information metrics were used to evaluate blurring and distortion, respectively. In a breast phantom, six permutations of MUSE-DWI with varying parallel acceleration factor and number of shots were compared. Differences in ADC and nADC were compared using the coefficient of variation in the phantom and a paired t-test in patients. Differences in blur, distortion and qualitative metrics were analysed using a Wilcoxon signed-rank test.

Results: There was a low coefficient of variation (<2%) in ADC between ss-EPI-DWI and all MUSE-DWI permutations acquired using the phantom. 22 malignant and three benign lesions were identified in 20 patients. ADC values measured using MUSE were significantly lower compared to ss-EPI for malignant but not benign lesions (p < 0.001, p = 0.21). nADC values were not significantly different (p = 0.62, p = 0.28). Blurring and distortion improved with number of shots and acceleration factor, and significantly improved with MUSE in patients (p < 0.001, p = 0.002). Qualitatively, image quality improved using MUSE.

Conclusion: MUSE improves the image quality of breast DWI compared to ss-EPI.

Advances in knowledge: MUSE-DWI has superior image quality and reduced blurring and distortion compared to ss-EPI-DWI in breast cancer.

INTRODUCTION

Diffusion-weighted imaging (DWI) is a promising non-contrast MRI technique in the detection of breast cancer. However, DWIs acquired using single-shot echo-planar imaging (ss-EPI) are limited by image artefacts and suffer from blurring due to T2* decay during readout. The relatively low spatial resolution results in the averaging of tumour volumes with nearby breast tissues, particularly for small lesions and non-mass enhancements which are prone to volume-averaging affects due to the large voxel sizes of DWI compared to dynamic contrast-enhanced MRI (DCE-MRI). In order for diffusion MRI to be clinically relevant, it must be able to detect and characterise all lesions, even small lesions.

Parallel imaging techniques (such as SENSE) aim to reduce artefacts in images acquired using EPI by reducing the length of the echo train and therefore reducing the T2* blurring. However, the extent to which parallel imaging can improve image quality is limited by the coil hardware. Multi-shot techniques also attempt to reduce the length of the echo train by acquiring k-space in a number of segments. These techniques must account for motion between shots, often using a ‘navigator’ pulse to acquire
Multi-shot diffusion-weighted imaging using MUSE in breast cancer

Multi-shot sensitivity encoding (MUSE), a multi-shot segmented EPI technique, expands on existing sensitivity-encoding techniques by acquiring $k$-space with an interleaved trajectory with the aim of achieving better spatial resolution and high signal-to-noise ratio (SNR) without the need for navigator pulses between interleaves.\(^5\) Benefits of a navigator-free method include an increase in efficiency of acquisition as navigators require 30–40 ms per excitation.\(^6\) Similar to parallel acceleration, MUSE is limited by the design of the receiver coil and the number of coil components. While MUSE has not yet been investigated in the breast, a number of studies have investigated the improvement in image quality and diagnostic performance of RESOLVE, a readout-segmented EPI technique.\(^3\)\(^,\)\(^7\)\(^–\)\(^10\)

This study aimed to compare the image quality of DWI acquired using ss-EPI and MUSE, and investigate the effect of MUSE, used in conjunction with parallel acceleration, on the quantification of the apparent diffusion coefficient (ADC) in a breast phantom and in malignant, benign and normal breast tissue in a small cohort of patients. In addition, an optimised parallel acceleration factor and number of shots for MUSE-DWI were determined.

**METHODS AND MATERIALS**

**Phantom study**

**Phantom study design**

The effect of varying parallel acceleration factor and number of shots was first evaluated in a phantom. The phantom used in this study was manufactured by QMD (High Precision Devices, Inc., Boulder, CO), and consisted of two silicon breast inserts designed to investigate diffusion, distortion and $T_1$ relaxation. The diffusion insert contained 16 vials of solutions arranged in two concentric rings with diameter and length 15×110 mm (inner ring) and 15×80mm (outer ring) (Figure 1). 12 vials contained varying concentrations of polyvinylpyrrolidone (PVP) in water (0, 10, 14, 18, 25, and 40%) to mimic the apparent diffusion coefficients of benign and malignant tumours found in the literature and four vials contained a fat mimic. The interstitial space was filled with a solution of 35% corn syrup in water to mimic fibroglandular tissue. The phantom was kept in the scanner room at room temperature (18°C).

**Table 1. Scan parameters**

|                      | Phantom studies | Patient studies |
|----------------------|-----------------|----------------|
|                      | **ss-EPI-DWI**  | **MUSE-DWI**   |
|                      | **ss-EPI-DWI**  | **MUSE-DWI**   | **DCE-MRI**   |
| Slice Orientation    | Axial           | Axial          | Axial         |
| $b$-values (s/mm$^2$) | 0, 800          | 0, 800         | 0, 800        |
| TR/TE (ms)           | 4000/51         | 4000/54–55     | 3714-14853/60.7 | 8203-16715/71.3 | 5.1/2.4 |
| Matrix size          | 160 × 160       | 160 × 160      | 96 × 128      | 160 × 160      | 384 × 384 |
| Field-of-view (mm$^2$) | 360 × 360       | 360 × 360      | 360 × 360     | 360 × 360      | 360 × 360 |
| Slice thickness (mm) | 3.5             | 3.5            | 3.5           | 3.5            | 2.0        |
| Acceleration factor  | 2               | 1, 1.5, 2      | 2             | 1.5            | 2.5        |
| Number of shots      | 1               | 2, 3           | 1             | 2              | 2          |
| Acquisition Time     | 1 min 8 s       | 1 min 52 s, 2 min 48 s | 6–8 min | 8 min | 9 min |

ss-EPI: single-shot echo-planar imaging, DWI: diffusion-weighted imaging, MUSE: multiplexed sensitivity encoding, DCE: dynamic contrast-enhanced, TR: repetition time, TE: echo time.
All phantom acquisitions were performed on a 3T system (Discovery MR 750, GE Healthcare, Waukesha, WI, USA) using a dedicated bilateral eight-channel phased-array breast coil. DWI was performed using ss-EPI and six permutations of MUSE-DWI with varying parallel acceleration factors and numbers of shots (Table 1). A T₁W image (TR/TE = 659/15.3 ms, matrix = 512x512, slice thickness = 3.5 mm) was acquired using a fast spin echo (FSE) sequence as a geometric reference.

A measurement of blurring was obtained using the Crété-Roffet blur metric, previously adapted from the field of computer vision, for a central slice through the phantom for all acquisitions. The metric is quantified by comparing the perceptible levels of blurring between the input image and a version of the image which has been further blurred using a low-pass filter. Values of the blur metric range from 0 (sharp) to 1 (blurry). A measurement of distortion was obtained by calculating the Mattes Mutual Information (MI) metric between the ss-EPI and MUSE images, resampled to the same matrix size as the DWI images. Values of the MI metric range from 0 (distorted) to 1 (not distorted). Metrics were calculated for each slice and averaged over the whole image volume.

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For each of the 16 vials, regions of interest (ROIs) were drawn on axial images for each of the diffusion gels using Osirix (v.8.0.1, Pixmeo, Switzerland). Each ROI consisted of three rectangles drawn on contiguous slices, avoiding the edges of the vials to minimise partial volume effects. Generation of ADC maps and diffusion analyses were carried out using in-house software developed in MATLAB (The Mathworks, version 2019a). The mean ADC was measured for each ROI and the coefficient of variation calculated between all acquisitions.

**Patient study design**

Between October 2018 and October 2019, 20 females were scanned under an existing research ethics protocol. The local institutional review boards and ethics committees approved this study and written informed consent was obtained from all patients. Patients were recruited if they had pathologically confirmed invasive breast cancer and had no previous treatment (such as breast surgery or neoadjuvant chemotherapy).

All clinical acquisitions were performed on a 1.5T system (MR 450W, GE Healthcare, Waukesha, WI) using an 8-channel breast coil. ss-EPI-DWI and MUSE-DWI were performed. High-resolution T₁W dynamic contrast-enhanced (DCE) images were also acquired. Scan parameters are given in Table 1.

**Image analysis**

**Qualitative analysis**

Two breast radiologists, each of whom had at least 5 years of experience, independently reviewed all the images. Both ss-EPI and MUSE images were scored based on three qualitative image criteria: lesion conspicuity, contrast between lesion and tissue and diagnostic confidence. All criteria were scored from 1 (poor) to 5 (excellent). Lesions were also assessed for multi-focality and multi-centricity.

**Quantitative analysis**

The Crété-Roffet blur metric was calculated for ss-EPI and MUSE images, and the MI distortion metric was calculated between ss-EPI and MUSE images and the corresponding DCE images, resampled to the same matrix size as the DWI images. Metrics were calculated for each slice and averaged over the whole image volume.

| PVP concentration (%) | Apparent Diffusion Coefficient \(10^{-3} \text{mm}^2/\text{s}\) | Coefficient of variation (%) |
|-----------------------|--------------------------------------------------|-----------------------------|
|                       | ss-EPI MUSE two shots MUSE three shots            |                             |
|                       | Acceleration Factor Acceleration Factor Acceleration Factor |
| 0                     | 2.40 2.39 2.43 2.41 2.42 2.43 2.41               | 1.2                         |
| 10                    | 1.74 1.75 1.79 1.79 1.79 1.81 1.82               | 1.5                         |
| 14                    | 1.58 1.48 1.48 1.48 1.48 1.50 1.51               | 1.9                         |
| 18                    | 1.37 1.29 1.31 1.31 1.31 1.32 1.33               | 1.8                         |
| 25                    | 1.19 1.09 1.10 1.11 1.11 1.12 1.13               | 1.6                         |
| 40                    | 0.67 0.60 0.70 0.69 0.69 0.66 0.67               | 1.6                         |

PVP: polyvinylpyrrolidone, ss-EPI: single-shot echo-planar imaging, MUSE: multiplexed sensitivity encoding
were calculated for each slice and averaged over the whole image volume.

ADC maps were generated using in-house software developed in MATLAB (v. 2019a). ROIs were manually drawn on the ss-EPI and MUSE b800 images by two breast radiologists in consensus with reference to DCE images to aid tumour delineation. As previously described by Wisner et al., to obtain a measurement of signal in normal tissue the ROI for each lesion was copied and placed on contralateral fibroglandular tissue. These ROIs were copied to the ADC maps and mean ADC values were measured for each lesion (ADClesion) and normal fibroglandular tissue (ADCtissue) ROI. To account for the difference in acquired voxel size, the normalised ADC (nADC) was calculated using \[ nADC = \frac{ADC_{\text{lesion}}}{ADC_{\text{tissue}}} \].

Statistical analysis
Qualitative and quantitative image quality metrics for images using ss-EPI or MUSE were compared on a per-patient basis. ROI sizes and ADC values were compared on a per-lesion basis. All comparisons used the paired t-test or a Wilcoxon signed-rank test. The interobserver agreement of the qualitative scoring criteria was assessed using Cohen’s \( \kappa \). \( p < 0.05 \) was considered statistically significant for all statistical tests. All statistical analyses were carried out using R (R v.3.1.3; R Foundation for Statistical Computing, Vienna, Austria).

RESULTS
Phantom results
Images acquired using ss-EPI and all permutations of MUSE are shown in Figure 2. ADC values measured using the phantom were consistent with values from the literature. The variation in ADC for each acquisition is shown in Table 2. There was a low coefficient of variation for measured ADC value (<2%) for all PVP concentrations. The variation in MI metric and Créte-Roffet blur metric between acquisitions is shown in Figure 3. Both distortion and blurring improved (an increase in the MI metric and a decrease in the Créte-Roffet blur metric) with an increasing number of shots and acceleration factor. Increasing the acceleration factor when using three shots did not improve distortion. For clinical implementation, an acceleration factor of 1.5 and 2 shots were chosen as a pragmatic compromise between acquisition time and image quality.

Patient results
20 patients (median age 51 years, range 28–81 years) were imaged. A total of 22 malignant lesions and three benign lesions were identified (median size 16 mm, range 10–56 mm). There were five patients who had more than one lesion. Two patients had a second lesion located in the contralateral breast. Three patients had a second lesion located in the same breast. Details of the patient and lesion characteristics are given in Table 3. Comparisons of image quality for an invasive ductal carcinoma, an invasive lobular carcinoma, and a patient with an invasive carcinoma with mixed ductal and lobular growth patterns and columnar and fibrocystic changes with benign calcification are shown in Figure 4.
Qualitative results
Results of qualitative comparisons are given in Table 4. MUSE was superior to ss-EPI for contrast between lesion and tissue, and significantly superior for lesion conspicuity and diagnostic confidence criteria. Inter-reader agreement as measured by Cohen’s κ was higher for the scoring of MUSE images compared to ss-EPI.

Quantitative results
Results of quantitative comparisons are given in Table 5. The distributions of blur and distortion metrics using ss-EPI and MUSE are shown in Figure 5. The Créte-Roffet blur metric was significantly lower for MUSE-DWI than for ss-EPI-DWI (p < 0.001), indicating less blurring. The MI metric was significantly higher for MUSE-DWI than for ss-EPI-DWI (p = 0.002), indicating more similarity to DCE images and therefore less distortion.

Comparison of ADC values: ss-EPI versus MUSE
The distributions of ADC values measured using ss-EPI and MUSE are shown in Figure 6A. The mean ADC of malignant lesions was 1.49 ± 0.30 mm²/s using ss-EPI and 1.23 ± 0.27 mm²/s using MUSE. The mean ADC of benign lesions was 1.73 ± 0.27 mm²/s using ss-EPI and 1.42 ± 0.25 mm²/s using MUSE. ADC values measured using MUSE were significantly lower than those measured using ss-EPI for malignant lesions but not benign lesions (p < 0.001 and p = 0.21, respectively). ADC values measured were in agreement with those in the literature, although the mean ADC for malignant lesions measured using ss-EPI was high.

The distributions of nADC values measured using ss-EPI and MUSE are shown in Figure 6B. The mean nADC of malignant lesions was 0.69 ± 0.24 using ss-EPI and 0.72 ± 0.18 using MUSE. The mean nADC of benign lesions was 0.79 ± 0.21 using ss-EPI and 0.97 ± 0.23 using MUSE. There was no significant difference in nADC values of malignant and benign lesions measured using ss-EPI or MUSE (p = 0.62 and p = 0.28, respectively). The separation of the mean nADC values for malignant and benign lesions was greater using MUSE (0.25) than using ss-EPI (0.10).

DISCUSSION
In this study, we demonstrated through phantom and clinical experiments that the quality of DWI can be improved using MUSE by significantly reducing blurring and distortion. ADC values measured in the phantom and in patients were in agreement with those in the literature.

Geometric distortion is prevalent in DWI due to the slow traversal through k-space along the phase-encoding direction when using ss-EPI. Accelerating the k-space trajectory through the use of a segmented-EPI technique such as MUSE can improve distortion compared to ss-EPI, as has been shown in previous studies implementing MUSE in the brain. Distortion has been evaluated in breast DWI by comparing lesion lengths in the anteroposterior and left-right directions or by using anatomic landmarks. However, a more robust technique using the MI metric, a similarity measure often used in the field of image registration, was used by Teruel et al when investigating the reduction of distortion after correction of B₀ inhomogeneities. Distortion as measured by the MI metric in our study significantly improved using MUSE. In phantom studies, the best reduction in distortion was achieved using MUSE with three shots, however this was not clinically feasible due to the increase in acquisition time. A limitation of the MUSE technique is that the scan time increases proportional to the number of shots acquired.

MUSE-DWI using simultaneous multi-slice excitation could be employed to reduce the acquisition time. This technique has been investigated in readout-segmented DWI in the breast.

Blurring is also a limiting factor in the clinical use of ss-EPI DWI, affecting accurate measurement of the ADC. Low-resolution images result in ROIs that are averaged with adjacent normal fibroglandular tissue, leading to higher measured ADC values. While ADC

Table 4. Qualitative comparisons

| Criteria            | Score range | ss-EPI | MUSE | p-value | κ ss-EPI | κ MUSE |
|---------------------|-------------|--------|------|---------|----------|--------|
| Lesion conspicuity  | 1–5         | 3.5 ± 1.1 | 4.1 ± 0.9 | <0.001 | 0.05 | 0.4 |
| Lesion-tissue contrast | 1–5     | 3.8 ± 1.2 | 3.9 ± 1.0 | 0.86 | -0.007 | 0.3 |
| Diagnostic confidence | 1–5      | 3.7 ± 1.1 | 4.2 ± 1.1 | <0.001 | 0.08 | 0.2 |

κ: Cohen’s κ; MUSE: multiplexed sensitivity encoding, ss-EPI: single-shot echo-planar imaging. p-values were determined using a Wilcoxon signed-rank test.
values measured in the breast phantom did not vary substantially using ss-EPI or MUSE when using the same set of ROIs, ADC values measured in patients were lower using MUSE, suggesting that the difference in ADC may be a result of improved resolution and therefore improved ROI delineation using MUSE. More accurate quantification of the ADC should improve the separation of ADC values between malignant, benign and normal fibroglandular tissue, as is seen by the greater separation of nADC values between malignant and benign breast lesions. Wisner et al similarly reported an improved separation of malignant and benign lesions using RESOLVE.3 In our study, there were not enough lesions to do an analysis of diagnostic performance. However, given that readout-segmented techniques have been shown to have a better diagnostic performance compared to ss-EPI,8,9 we expect that MUSE will achieve a better diagnostic performance than ss-EPI.

Less blurring and higher spatial resolution will allow DWI to better detect smaller lesions and satellite lesions. Adhering to the EUSOBI guidelines,19 the patients included in this study were referred for MRI for preoperative local staging of ipsilateral and contralateral newly diagnosed breast cancers where patients had dense breasts or lobular cancers (resulting in a high number of lobular cancers in this study which does not reflect the natural prevalence of this type of cancer) as well as for problem-solving (equivocal findings at mammography and ultrasound). In these cases, the improved image quality of MUSE-DWI can be used to better assess multifocality and multi-centricity, which impacts surgical planning.20 Reduced distortion may also allow for DWI to be used for morphological analysis, increasing the clinical utility of DWI.

There are limitations to this study. Firstly, low numbers of patients and lesions limit the conclusions that can be drawn. More benign lesions are required to evaluate diagnostic performance. Secondly, mostly mass type tumours were included in this study. Further work with a larger range of histopathological subtypes are required to prove the robustness of MUSE-DWI. Thirdly, we were not able to acquire our current clinical protocol with a matrix size of greater than 96 × 128 to match the matrix size used for MUSE as this resulted in a high level of distortion. This limits the conclusions that can be drawn from the improvement in blurring. However, phantom images acquired with ss-EPI and MUSE at the same resolution allowed for more direct comparisons of blurring and distortion. Finally, SNR and contrast-to-noise ratio (CNR) were not compared due to the varying parallel acceleration factors and SENSE reconstructions used among acquisitions, resulting in different noise characteristics.

CONCLUSION
In conclusion, the image quality of MUSE-DWI was superior to that of ss-EPI-DWI and geometric distortion and blurring were significantly reduced, although there was an increase in acquisition time. MUSE-DWI is a promising technique for the detection and

Table 5. Quantitative measures

| Parameter                      | ss-EPI        | MUSE         | p-value |
|-------------------------------|---------------|--------------|---------|
| Mean ADC (x 10^-3 mm^2/s)     |               |              |         |
| Fibroglandular tissue         | 2.24 ± 0.45   | 1.76 ± 0.32  | <0.001  |
| Benign Lesions                | 1.73 ± 0.27   | 1.42 ± 0.25  | 0.21    |
| Malignant lesions             | 1.49 ± 0.30   | 1.23 ± 0.27  | <0.001  |
| Mean normalised ADC           |               |              |         |
| Benign Lesions                | 0.79 ± 0.21   | 0.97 ± 0.23  | 0.28    |
| Malignant lesions             | 0.69 ± 0.24   | 0.72 ± 0.18  | 0.62    |
| Crété-Roffet blur metric      | 0.10 ± 0.02   | 0.08 ± 0.02  | <0.001  |
| Mattes Mutual Information metric | 0.19 ± 0.12 | 0.23 ± 0.10  | 0.002   |

ADC: apparent diffusion coefficient; MUSE: multiplexed sensitivity encoding, ss-EPI: single-shot echo-planar imaging. p-values were determined using a paired t-test.
characterisation of breast lesions; however further work is required to evaluate the diagnostic performance of this technique using a larger range of histopathological subtypes of breast cancer.

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