Readout-Segmented Echo-Planar Diffusion-Weighted MR Imaging Improves the Differentiation of Breast Cancer Receptor Statuses Compared With Conventional Diffusion-Weighted Imaging

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Background: Readout-segmented echo-planar diffusion-weighted imaging (RS-EPI) can improve image quality and signal-to-noise ratio, the resulting apparent diffusion coefficient (ADC) value acts as a more sensitive biomarker to characterize tumors. However, data regarding the differentiation of breast cancer (BC) receptor statuses using RS-EPI are limited.

Purpose: To determine whether RS-EPI improves the differentiation of receptor statuses compared with conventional single-shot (SS) EPI in breast MRI.

Study Type: Retrospective.

Population: A total of 151 BC women with the mean age of 50.6 years.

Field strength/Sequence: A 3 T/ RS-EPI and SS-EPI.

Assessment: The ADCs of the lesion and normal background tissue from the two sequences were collected by two radiologists with 15 years of experience working of breast MRI (M.H.Z. and X.F.C.), and a normalized ADC was calculated by dividing the mean ADC value of the lesion by the mean ADC value of the normal background tissue.

Statistical Tests: Agreement between the ADC measurements from the two sequences was assessed using the Pearson correlation coefficient and Bland–Altman plots. One-way analysis of variance, Kruskal–Wallis test, and median difference were used to compare the ADC measurements for all lesions and different receptor statuses. A P value less than 0.05 indicated a significant result.

Results: The ADC measurements of all lesions and normal background tissues were significantly higher on RS-EPI than on SS-EPI (1.82 ± 0.33 vs. 1.55 ± 0.30 and 0.83 ± 0.11 vs. 0.79 ± 0.10). The normalized ADC was lower on RS-EPI than on SS-EPI (0.47 ± 0.11 vs. 0.53 ± 0.12, a median difference of −0.04 [95% CI: −0.256 to 0.111]). For both diffusion methods, only the ADC measurement of RS-EPI was higher for human epidermal growth factor receptor-2 (HER-2)-positive tumors than for HER-2-negative tumors (0.87 ± 0.10 vs. 0.81 ± 0.11), and this measurement was associated with HER-2 positive status (adjusted odds ratio [OR] = 654.4); however, similar results were not observed for the ADC measurement of SS-EPI (0.80 ± 0.10 vs. 0.78 ± 0.11 with P = 0.199 and adjusted OR = 0.21 with P = 0.464, respectively).

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Diffusion-weighted imaging (DWI), which is a complementary technique for dynamic contrast-enhanced (DCE) magnetic resonance imaging (MRI) that reduces false-positive rates in characterizing tumor lesions, is widely used in breast imaging and plays a useful role in the diagnosis, treatment, and prognosis of breast cancer (BC).\textsuperscript{1–8} Apparent diffusion coefficient (ADC) maps obtained with DWI can provide information about the microscopic cellular environment that can be used for characterizing tumors.\textsuperscript{7,9,10}

A major strength of the ADC is its quantitative character, which allows it to be used as a potential imaging biomarker for characterizing prognostic factors, such as estrogen receptor (ER), progesterone receptor (PR), and human epidermal growth factor receptor-2 (HER-2) expression, as well as the proliferation rate (Ki-67).\textsuperscript{3,11} Numerous studies have shown that tumors with a high Ki-67 index and an increased cell density are expected to have lower ADC measurements than tumors with a low Ki-67 index, whereas HER-2-positive tumors with increased neovascularity are expected to have higher ADC measurements than HER2-negative tumors, and similar results have been observed when comparing hormone receptor-positive tumors with hormone receptor-negative tumors.\textsuperscript{3,12–15}

However, previous studies have reached inconsistent results in using ADC to identify these prognostic factors.\textsuperscript{16–18} These disagreements might be attributable to the use of different MRI techniques. For example, readout-segmented (RS) echo-planar diffusion-weighted imaging (EPI) based on segment sampling of the EPI sequence in the readout direction can improve image quality and signal-to-noise ratio (SNR), and the resulting ADC value acts as a more sensitive imaging biomarker to predict BC receptor statuses than the ADC from single-shot (SS) EPI.\textsuperscript{3,16–19} However, there are still limited data regarding the diagnostic performance of RS-EPI and its potential to replace SS-EPI in characterizing the receptor statuses of BC. Therefore, this study aimed to compare the sensitivity and reliability of RS-EPI in the characterization of receptor statuses in clinical practice with those of SS-EPI.

Data Conclusion: RS-EPI can improve the distinction between HER-2-positive and HER-2-negative breast cancer, complementing the clinical application of diffusion imaging.

Evidence Level: 3
Technical Efficacy: Stage 1

Clinical Data
Clinicopathological data, including age, maximum and minimum tumor diameter, ER, PR, and HER-2 statuses, and Ki-67 index, were obtained from the medical electronic record system. The tumor size was measured on the largest section of the BC in the DCE-MRI images. ER- or PR-positive tumors were defined as tumors with at least 1% of cells that were positively stained according to immunohistochemistry.\textsuperscript{2,20} HER-2-positive tumors were defined as tumors with scores of 3+ or tumors with scores of 2+ and positive fluorescence in situ hybridization results of HER-2 gene amplification.\textsuperscript{21} A Ki-67 index ≥20% indicated high expression.\textsuperscript{2,20}

MRI Techniques
All the examinations were performed on a 3 T MRI scanner (MAGNETOM Skyra, Siemens Healthcare, Erlangen, Germany) with a 16-channel bilateral breast coil using prone positioning. The scanning protocol included conventional $T_1$-weighted acquisition (repetition time [TR]/echo time [TE] 5.5/2.5 msec, field of view [FOV] $341$ mm × $341$ mm, matrix $426$ × $448$, slice thickness $1.5$ mm, slice gap $0.3$ mm), $T_2$-weighted acquisition (TR/TE $3570/74$ msec, inversion time $230$ msec, FOV $341$ mm × $341$ mm, matrix $314$ × $448$, slice thickness $4.0$ mm, slice gap $0.4$ mm), DWI, and a DCE series. The DCE series consisted of precontrast $T_1$-weighted volume interpolated breath-hold examination (VIBE) imaging (TR/TE $3.78/1.38$ msec, FOV $340$ mm × $340$ mm, matrix $205$ × $256$, slice thickness $2$ mm, voxel resolution $1.3$ mm × $1.3$ mm × $2$ mm) and multi arterial time-resolved imaging with interleaved stochastic trajectories (TWIST)-VIBE DCE scanning with 34 consecutive phases (TR/TE $6.4/3.34$ msec, slice thickness $2.0$ mm, FOV $340$ mm × $340$ mm, matrix $289$ × $303$, temporal resolution $8.9$ seconds) after intravenous bolus injection of gadopentetate dimeglumine (Bayer Pharma AG) with an injection rate of $3.0$ mL/sec.

Axial DWI images were sequentially obtained by RS-EPI and SS-EPI techniques before contrast enhancement. RS-EPI and SS-EPI sequences were designed with the same b-values ($50$ and $800$ sec/mm$^2$) in-plane resolution of $1.8$ × $1.8$ mm$^2$, a slice thickness of $4.0$ mm with an injection rate of $3.0$ mL/sec.
| Receptor Status | Age (year) | Maximum Diameter | Minimum Diameter | ADC Value of RS-EPI ($\times 10^{-3}$ cm$^2$) | ADC Value of SS-EPI ($\times 10^{-3}$ cm$^2$) | Normalized ADC Value of RS-EPI | Normalized ADC Value of SS-EPI |
|----------------|-----------|------------------|-----------------|---------------------------------------------|---------------------------------------------|--------------------------------|--------------------------------|
| **ER**         |           |                  |                 |                                             |                                             |                                |                                |
| Negative (n = 56) | 51.59 ± 9.07 | 4.01 ± 1.67      | 2.58 ± 1.12     | 0.84 ± 0.12                                 | 0.80 ± 0.10                                 | 0.46 ± 0.11                     | 0.52 ± 0.13                     |
| Positive (n = 95) | 50.01 ± 11.62 | 3.30 ± 1.69      | 2.02 ± 0.92     | 0.83 ± 0.11                                 | 0.78 ± 0.10                                 | 0.48 ± 0.11                     | 0.53 ± 0.12                     |
| **P-value**    | 0.385     | **0.002**        | **<0.001**      | 0.510                                       | 0.480                                       | 0.543                          | 0.282                          |
| **PR**         |           |                  |                 |                                             |                                             |                                |                                |
| Negative (n = 92) | 52.33 ± 10.31 | 3.86 ± 1.86      | 2.46 ± 1.10     | 0.82 ± 0.11                                 | 0.78 ± 0.11                                 | 0.46 ± 0.11                     | 0.51 ± 0.13                     |
| Positive (n = 59) | 47.90 ± 10.93 | 3.10 ± 1.35      | 1.85 ± 0.80     | 0.85 ± 0.11                                 | 0.80 ± 0.10                                 | 0.50 ± 0.11                     | 0.56 ± 0.11                     |
| **P-value**    | **0.013** | **0.006**        | **<0.001**      | 0.192                                       | 0.480                                       | **0.020**                      | **0.002**                      |
| **HER-2**      |           |                  |                 |                                             |                                             |                                |                                |
| Negative (n = 100) | 51.00 ± 11.25 | 3.43 ± 1.68      | 2.20 ± 1.08     | 0.81 ± 0.11                                 | 0.78 ± 0.11                                 | 0.46 ± 0.11                     | 0.52 ± 0.12                     |
| Positive (n = 51) | 49.90 ± 9.75 | 3.82 ± 1.76      | 2.27 ± 0.93     | 0.87 ± 0.10                                 | 0.80 ± 0.10                                 | 0.51 ± 0.11                     | 0.55 ± 0.13                     |
| **P-value**    | 0.572     | 0.204            | 0.365           | **0.001**                                   | 0.199                                       | **0.010**                      | 0.173                          |
| **Ki-67**      |           |                  |                 |                                             |                                             |                                |                                |
| <20% (n = 36)  | 51.44 ± 10.63 | 3.18 ± 1.22      | 1.80 ± 0.60     | 0.84 ± 0.11                                 | 0.80 ± 0.11                                 | 0.50 ± 0.10                     | 0.55 ± 0.12                     |
| ≥20% (n = 115) | 50.33 ± 10.81 | 3.68 ± 1.83      | 2.36 ± 1.10     | 0.83 ± 0.12                                 | 0.79 ± 0.10                                 | 0.46 ± 0.11                     | 0.52 ± 0.12                     |
| **P-value**    | 0.589     | 0.210            | **0.003**       | 0.398                                       | 0.551                                       | **0.029**                      | 0.154                          |

Bold values indicate statistically significant in P values.

ER = estrogen receptor; PR = progesterone receptor; HER-2 = human epidermal growth factor receptor-2; ADC = apparent diffusion coefficient.
a slice gap of 0.8 mm, generalized auto-calibrating partially parallel acquisitions (GRAPPA) was also used in both sequences with an acceleration factor of 2, and enough slices were acquired to cover the entire breast. In order to match the total acquisition time of the two sequences (approximately 5 minutes), the number of averages for SS-EPI and RS-EPI was set to 8 and 3, respectively. The RS-EPI sequence used five readout segments. The remainder of the parameters were as follows: RS-EPI (TR/TE 4800/56 msec, FOV 170 mm \times 340 mm, bandwidth 822 Hz, matrix 98 \times 190, echo spacing 0.36 msec) and SS-EPI (TR/TE 4200/62 msec, FOV 149 mm \times 340 mm, bandwidth 1730 Hz, matrix 100 \times 170, echo spacing 0.68 msec).

**MRI Image Analysis**

All the imaging analyses were independently carried out by two radiologists with 15 years of experience working with breast MRI (M.H. Z. and X.F.C.). All the image data were transferred to a Siemens Syngo via workstation. The BCs were identified on high b-value (800 sec/mm²) images using DCE-MRI images for reference and then evaluated on ADC maps using breast Tissue 4D software package embedded in dedicated workstation (Syngo.via).

The representative slice of the lesion that showed the largest section of the tumor was identified by the radiologist, and a free hand region of interest (ROI) with a size range of 20–30 mm² was drawn on the lowest hypointensity region of the lesion on the RS-EPI maps corresponding to a prominent area of enhancement on the DCE-MRI images. The ROI was placed to avoid visibly necrotic, cystic, bleeding, and calcification areas. Then the same ROI was copied to SS-EPI maps to measure the ADC value of tumor. For the ADC measurement of normal background tissue, the same ROI was copied to RS-EPI maps and SS-EPI maps on contralateral normal breast tissue. The corresponding ADC values of each lesion and the normal background tissue from RS-EPI and SS-EPI were recorded after motion correction. Each ADC value (\(\times 10^{-3} \text{ mm}^2/\text{s}\)) was measured three times, which were located on the largest tumor section and its adjacent sections, and then the averaged value was calculated for further analysis.

To reduce the impact of individual breast characteristics, normalized ADC (nADC) was introduced and calculated as the mean ADC value of the lesion divided by the mean ADC value of the normal background tissue.

**Statistical Analysis**

Statistical analysis was performed using R (Version 4.0.3, R Foundation for Statistical Computing, Vienna, Austria). Continuous variables are expressed as the mean \pm standard deviation. Categorical variables are expressed as counts (percentages). The normality of the variables was investigated by using the Shapiro–Wilk test. For normally distributed variables, one-way analysis of variance was used to assess the difference between the RS-EPI measurement and the SS-EPI measurement in different receptor statuses, while the Kruskal–Wallis test was used for non-normally distributed variables.

Univariate and multivariate logistic regression models were fitted and used to identify the risk factors for predictive positive receptor statuses. In addition, a paired \(t\)-test (for normally distributed variables) or a paired Wilcoxon test was used to compare the ADC measurement from RS-EPI and SS-EPI for all lesions and different receptor statuses. The median difference (assumed symmetric distribution) between RS-EPI and SS-EPI measurements, subtracting

| TABLE 2. Univariate and Multivariate Analysis of the Parameters of Different Receptor Statuses |
|---|
| **ER** | **PR** | **HER-2** |
| **Univariate Analysis** | **Multivariate Analysis** | **Cutoff OR** | **P** | **Univariate Analysis** | **Multivariate Analysis** | **Cutoff OR** | **P** | **Univariate Analysis** | **Multivariate Analysis** | **Cutoff OR** | **P** |
| Age | 0.78 0.016 | 2.55 0.92 | 2.32 0.84 | 0.03 0.394 | 6.50 0.017 | 0.96 0.19 | 0.39 0.508 | 0.86 0.478 | 0.94 0.19 | 0.93 0.24 |
| Max-diameter | 0.57 0.003 | 2.21 0.62 | 0.94 0.84 | 0.42 0.001 | 1.16 0.46 | 0.19 0.019 | 0.76 0.38 | 2.40 0.001 | 0.72 0.28 | 2.04 0.002 |
| Min-diameter | 0.78 0.016 | 2.55 0.92 | 2.32 0.84 | 0.03 0.394 | 6.50 0.017 | 0.96 0.19 | 0.39 0.508 | 0.86 0.478 | 0.94 0.19 | 0.93 0.24 |
| ADC value of RS-EPI | 0.37 0.59 | 0.86 0.59 | 0.86 0.59 | 0.42 0.001 | 1.16 0.46 | 0.19 0.019 | 0.76 0.38 | 2.40 0.001 | 0.72 0.28 | 2.04 0.002 |
| ADC value of SS-EPI | 0.37 0.59 | 0.86 0.59 | 0.86 0.59 | 0.42 0.001 | 1.16 0.46 | 0.19 0.019 | 0.76 0.38 | 2.40 0.001 | 0.72 0.28 | 2.04 0.002 |
| Bold values indicate statistically significant in \(p\) values. Max = maximum; Min = minimum; OR = odds ratio; ER = estrogen receptor; PR = progesterone receptor; HER-2 = human epidermal growth factor receptor 2-2.
the SS-EPI measurement from the RS-EPI measurement, was tracked and further compared among tumors with different receptor statuses. Interobserver agreements for the RS-EPI and SS-EPI measurements between the two radiologists were assessed using the intraclass correlation coefficient (ICC), which was categorized as good agreement (0.61–0.80) and excellent agreement (≥0.81). The agreement between the RS-EPI and SS-EPI measurements was analyzed using Pearson correlation and Bland–Altman analysis. A P value less than 0.05 indicated a significant result.

Results

Basic Clinicopathological Characteristics

A total of 151 IDC patients were analyzed, and their mean age was 50.6 ± 10.7 years. The table comparing measurements using average values of two readers are shown in Table 1, and the table comparing measurements using annotations of reader 1 and 2 are shown in Tables E1 and E2 in the supplementary materials. The maximum and minimum tumor diameters of ER-negative tumors were higher than those of ER-positive tumors (4.01 ± 1.67 vs. 3.29 ± 1.69 and 2.58 ± 1.12 vs. 2.02 ± 0.92, respectively), those of PR-negative tumors were higher than those of PR-positive tumors (3.86 ± 1.86 vs. 3.10 ± 1.35 and 2.46 ± 1.10 vs. 1.85 ± 0.80, respectively). However, the minimum diameter of low Ki-67 index tumors was smaller than that of high Ki-67 index tumors (1.80 ± 0.60 vs. 2.36 ± 1.10). In addition, the age of PR-positive patients was younger than that of PR-negative patients (47.90 ± 10.93 vs. 52.33 ± 10.31).

Comparisons of the ADC Measurement of Tumors With Different Receptor Statuses

The ADC measurements from RS-EPI were significantly higher for HER-2-negative tumors than for HER-2-positive tumors (0.81 ± 0.11 vs. 0.87 ± 0.10), but this difference was not confirmed with the ADC measurements of SS-EPI (0.78 ± 0.11 vs. 0.80 ± 0.10, P = 0.199). In addition, there was no significant difference in the ADC measurements from RS-EPI and SS-EPI with respect to the other receptor statuses or the Ki-67 index (Table 1). Paired comparisons of the ADC measurements of tumors with different receptor statuses using univariate and multivariate analysis revealed (Table 2) that a younger age (<49.5 years, adjusted odds ratio [OR] = 0.96) and a lower minimum diameter (<1.65 cm, adjusted OR = 0.46) were associated with PR-positive status, while higher ADC measurements from RS-EPI (>0.76 × 10⁻³ mm², adjusted OR = 654.4) were associated with HER2-positive status. Representative ADC images of RS-EPI and SS-EPI sequences from a breast cancer patient with HER-2-negative and HER-2-positive invasive ductal carcinoma are shown in Fig. 1.

Interobserver Agreement

The ICC values for the ADC measurements from RS-EPI and SS-EPI between the two radiologists were 0.657 (95% CI: 0.543–0.748) and 0.740 (95% CI: 0.647–0.811), respectively, indicating good agreement. The Bland–Altman plots of ADC measurement of SS-EPI and RS-EPI from two
The ICC values for the maximum diameter and minimum diameter of tumor between the two radiologists were 0.963 (95% CI: 0.949–0.973) and 0.945 (95% CI: 0.925–0.960), respectively.

Agreement Between ADC Measurements: RS-EPI vs. SS-EPI
There was a moderate positive correlation between the ADC measurements from RS-EPI and SS-EPI ($r = 0.679$; 95% CI: 0.613–0.73) for all lesions. However, the mean ADC measurements from RS-EPI for all lesions were higher than the mean ADC measurements of SS-EPI (0.83 ± 0.11 vs. 0.79 ± 0.10). These differences remained significant for ER-positive tumors (0.83 ± 0.11 vs. 0.78 ± 0.10), PR-positive tumors (0.82 ± 0.11 vs. 0.78 ± 0.11), PR-negative tumors (0.85 ± 0.11 vs. 0.80 ± 0.10), HER-2-positive tumors (0.87 ± 0.10 vs. 0.80 ± 0.10), and tumors with a high Ki-67 index (0.83 ± 0.12 vs. 0.79 ± 0.10) (Table 3).

![FIGURE 2: Bland–Altman plots showing the agreement between the RS-EPI measurement and the SS-EPI measurement for all lesions. Solid horizontal lines represent the mean bias, and the top and bottom dashed lines denote the upper and lower limits of agreement, respectively.](image)

### TABLE 3. Comparison of ADC Values From RS-EPI and SS-EPI for Tumors With Different Receptor Statuses

| Receptor Status | ADC Value of RS-EPI ($\times 10^{-3} \text{ cm}^2$) | ADC Value of SS-EPI ($\times 10^{-3} \text{ cm}^2$) | Median Difference (95% CI) | $P^a$ |
|-----------------|--------------------------------|--------------------------------|---------------------------|--------|
| ER Negative ($n = 56$) | 0.84 ± 0.12 | 0.80 ± 0.10 | 0.056 0.026 (−0.106 to 0.294) | 0.824 |
| ER Positive ($n = 95$) | 0.83 ± 0.11 | 0.78 ± 0.10 | **0.008** 0.041 (−0.135 to 0.255) | |
| PR Negative ($n = 92$) | 0.82 ± 0.11 | 0.78 ± 0.11 | **0.027** 0.031 (−0.122 to 0.244) | |
| PR Positive ($n = 59$) | 0.85 ± 0.11 | 0.80 ± 0.10 | **0.023** 0.052 (−0.13 to 0.264) | |
| HER2 Negative ($n = 100$) | 0.81 ± 0.11 | 0.78 ± 0.10 | 0.082 0.022 (−0.121 to 0.249) | |
| HER2 Positive ($n = 51$) | 0.87 ± 0.10 | 0.80 ± 0.10 | **0.001** 0.059 (−0.133 to 0.312) | |
| Ki-67 <20% ($n = 36$) | 0.84 ± 0.11 | 0.80 ± 0.11 | 0.096 0.054 (−0.132 to 0.217) | |
| Ki-67 ≥20% ($n = 115$) | 0.83 ± 0.12 | 0.79 ± 0.10 | **0.017** 0.036 (−0.126 to 0.26) | |

$P^a$ indicates the comparison of ADC measurements between RS-EPI and SS-EPI and $P^b$ indicates the comparison of ADC measurements for tumors with different receptor statuses. RS-EPI and SS-EPI indicate the ADC measurements of the RS-EPI and SS-EPI sequences, respectively. Bold values indicate statistically significant in $P$ values.

ER = estrogen receptor; PR = progesterone receptor; HER-2 = human epidermal growth factor receptor-2.
To further explore the differences between the RS-EPI and SS-EPI measurements for different receptor statuses, the median difference in the ADC measurements for tumors with different receptor statuses only revealed a significant difference between HER-2-negative and HER-2-positive tumors, with a higher difference between RS-EPI and SS-EPI measurements for HER-2-positive tumors than for HER-2-negative tumors (0.059 [95% CI: 0.133 to 0.312] vs. 0.022 [95% CI: 0.121 to 0.249]). Bland–Altman plots showed a mean bias of 0.056 [2.10 mm²/sec (lower than the upper limit of agreement, 0.12 to 0.23) for all lesions between the ADC measurements from RS-EPI and SS-EPI (Fig. 2).

Comparison of the Normalized ADC: RS-EPI vs. SS-EPI

RS-EPI yielded significantly higher ADC measurements of normal background tissue than SS-EPI (1.82 ± 0.33 vs. 1.55 ± 0.30). However, the normalized ADC was significantly lower on RS-EPI than on SS-EPI (0.47 ± 0.11 vs. 0.53 ± 0.12). Further analysis showed that the median difference in the normalized ADC between RS-EPI and SS-EPI revealed a significant difference between the two sequences, favoring a lower normalized ADC by using RS-EPI (−0.04 [95% CI: −0.256 to 0.111]). Representative ADC images of the normal background tissue and the normalized ADC in the RS-EPI and SS-EPI maps are shown in Fig. 3. In addition, the normalized ADC value of RS-EPI were lower for PR-negative tumors than for PR-positive tumors (0.46 ± 0.11 vs. 0.50 ± 0.11), and this difference was confirmed with the normalized ADC value of SS-EPI (0.51 ± 0.13 vs. 0.56 ± 0.11). The normalized ADC value of RS-EPI was lower for HER-2-negative tumors than for HER-2-positive tumors (0.46 ± 0.11 vs. 0.51 ± 0.11) and lower for high Ki-67 index tumors than for low Ki-67 index tumors (0.46 ± 0.11 vs. 0.50 ± 0.10), but those differences were not detectable with the normalized ADC value of SS-EPI (0.52 ± 0.12 vs. 0.55 ± 0.13 with \( P = 0.173 \), and 0.52 ± 0.12 vs. 0.55 ± 0.12 with \( P = 0.154 \), respectively) (Table 1).

Discussion

The ADC measurements from RS-EPI and SS-EPI of BC patients were assessed in this study. Compared to SS-EPI, RS-EPI has the potential advantage of differentiating between breast lesions with different receptor statuses. The results demonstrate that higher ADC measurements from RS-EPI are associated with HER2-positive status, which improves the diagnosis of HER-2 status in BC. In terms of hormone receptor statuses, although the ADC values were not significantly different, there were significant differences in tumor diameter.

It is well known that the ADC value correlates with tumor grade and proliferation index; however, previous research reported that there were no statistically significant associations between ADC values and breast cancer receptor statuses. This study indicates that the solution may be to use RS-EPI. This study found that the ADC value from RS-EPI could distinguish between HER-2-positive and HER-2-negative tumors, which cannot be achieved using the ADC value from traditional SS-EPI. According to the previous literature, the RS-EPI technique can better visualize anatomical details, and it results in a higher image quality than SS-EPI by reducing image distortion and improving the spatial resolution with two-dimensional navigator echoes using shortened echo spacing. In RS-EPI sequence, this shortened k-space traversal in the phase encoding direction results in less attenuation of T2* during readout, which leads to a reduction in distortion and blurring. Although these segmentations in RS-EPI increase the scanning time, the accompanying shorter echo time increases imaging signal level. Moreover, the motion-induced phase errors were corrected by 2D navigator echoes technique in RS-EPI (Fig. E2 was added to the supplementary material to demonstrate the image quality of RS-EPI and SS-EPI). Amornsiripanitch et al indicated that using RS-EPI was more sensitive than using SS-
EPI to detect BC. Bogner et al also demonstrated that ADC measurements from RS-EPI could improve the distinction of benign from malignant lesions.\(^\text{29}\)

Similar to the finding in gastric cancers, the ADC values were higher in HER-2-positive tumors.\(^\text{33}\) HER-2 is a prognostic and predictive biomarker in several human cancers.\(^\text{34}\) The over expression of HER-2 enables the activation of growth signaling pathways to promote cell proliferation and suppress apoptosis; therefore, HER-2 positivity is associated with malignant tumors. According to previous studies, malignant lesions generally exhibit lower ADC measurements than benign lesions but with a significant overlap.\(^\text{6,14,35}\) However, this study showed that HER-2-positive tumors exhibited higher ADC measurements than HER-2-negative tumors. A potential explanation for this result may be that HER-2-positive tumors have increased tumor angiogenesis and perfusion, resulting in higher ADC measurements in vivo.\(^\text{14}\)

Although ADC values are not significantly different in tumors with different hormone receptor (ER and PR) statuses, there are significant differences in tumor diameter, but of which is not a DWI-specific feature. Consistent with previous research, larger tumors were more likely to be ER- and PR-negative, indicating the potential advanced stage of the tumor.\(^\text{36,37}\) Moreover, a larger tumor size was associated with a higher risk of recurrence.\(^\text{38}\) However, the tumor diameter cannot reflect the status of HER-2 in this study.

It is worth noting that ADC measurements are confounded by multiple factors, including imaging parameters, data analysis, and pathophysiologic features.\(^\text{6,22,39}\) Therefore, special ADC measurement methods should be used to reduce the above variation to reliably assess its clinical utility. Similar to previous studies, ADC normalization in this study was introduced to reduce these variations.\(^\text{8,39}\) This study showed that the normalized ADC from RS-EPI was significantly lower than that from SS-EPI. This result is partly consistent with a previous study that showed a low normalized ADC for RS-EPI on breast MRI compared with conventional ADC, resulting in a reduction in the overlap of ADCs between benign and malignant lesions and an increase in the diagnostic performance.\(^\text{7,39}\)

**Limitations**

First, although this study was conducted with a relatively large cohort at a single institution. Findings from a larger multicenter study could make the results more reliable and provide more effective evidence of RS-EPI for diagnosis. Second, due to the time-consuming manual measurement of 3D ROIs that may better reflect the ADC measurements of tumors, 2D ROIs were only used to evaluate the tumors. 3D tumor segmentation could potentially be automated for evaluation of the entire tumor in the future. Third, the difference in ADC values between RS-EPI and SS-EPI is still unknown. Although previous research also reported this difference, further research to understand the source of this discrepancy will be meaningful.\(^\text{29}\) Fourth, this results show that ADC values of RS-EPI can only distinguish HER-2 positive and HER-2 negative tumors, which may limit its utility. In contrast, the normalized ADC values of RS-EPI have better performance in distinguishing PR, HER-2, and Ki-67 status. However, it is still difficult to determine the status of ER through diffusion imaging, which requires further exploration in the future.

**Conclusion**

This study demonstrated that the ADC value from RS-EPI can improve the distinction between HER-2-positive and HER-2-negative breast cancer, complementing the clinical application of diffusion imaging.

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**Ethics Approval and Consent to Participate**

This retrospective study was approved by the institutional review board of Meizhou People’s Hospital.

**Consent for Publication**

The institutional review board waived the need to obtain informed consent for this retrospective study.

**Data Availability**

The data cohorts used and/or analyzed in the present study are available from the corresponding authors upon reasonable request.

**Conflict of interests**

One author of the study (M.Z.W.) is a consultant for Siemens Healthcare and provided some critical technical support in the manuscript. Other authors are not employees or consultants in the industrial fields and have reported any data or information that may present a conflict of interest.

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