

Original Article

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Correspondence to:

Bong Joo Kang, M.D., Ph.D.
Department of Radiology, Seoul
St. Mary's Hospital, College of
Medicine, The Catholic University
of Korea, 222, Banpo-daero,
Seocho-gu, Seoul 06591, Korea.
Tel. +82-2-2258-6253
Fax. +82-2-599-6771
E-mail: gmlionmain@gmail.com

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Computer-Aided Detection with Automated Breast Ultrasonography for Suspicious Lesions Detected on Breast MRI

Sanghee Kim, Bong Joo Kang, Sung Hun Kim, Jeongmin Lee, Ga Eun Park

Department of Radiology, Seoul St. Mary's Hospital, The Catholic University of Korea, Seoul, Korea

Purpose: The aim of this study was to evaluate the diagnostic performance of a computer-aided detection (CAD) system used with automated breast ultrasonography (ABUS) for suspicious lesions detected on breast MRI, and CAD-false lesions.

Materials and Methods: We included a total of 40 patients diagnosed with breast cancer who underwent ABUS (ACUSON S2000) to evaluate multiple suspicious lesions found on MRI. We used CAD (QVCAD™) in all the ABUS examinations. We evaluated the diagnostic accuracy of CAD and analyzed the characteristics of CAD-detected lesions and the factors underlying false-positive and false-negative cases. We also analyzed false-positive lesions with CAD on ABUS.

Results: Of a total of 122 suspicious lesions detected on MRI in 40 patients, we excluded 51 daughter nodules near the main breast cancer within the same quadrant and included 71 lesions. We also analyzed 23 false-positive lesions using CAD with ABUS. The sensitivity, specificity, positive predictive value, and negative predictive value of CAD (for 94 lesions) with ABUS were 75.5%, 44.4%, 59.7%, and 62.5%, respectively. CAD facilitated the detection of 81.4% (35/43) of the invasive ductal cancer and 84.9% (28/33) of the invasive ductal cancer that showed a mass (excluding non-mass). CAD also revealed 90.3% (28/31) of the invasive ductal cancers measuring larger than 1 cm (excluding non-mass and those less than 1 cm). The mean sizes of the true-positive versus false-negative mass lesions were 2.08 ± 0.85 cm versus 1.6 ± 1.28 cm ($P < 0.05$). False-positive lesions included sclerosing adenosis and usual ductal hyperplasia. In a total of 23 false cases of CAD, the most common (18/23) cause was marginal or subareolar shadowing, followed by three simple cysts, a hematoma, and a skin wart.

Conclusion: CAD with ABUS showed promising sensitivity for the detection of invasive ductal cancer showing masses larger than 1 cm on MRI.

Keywords: Breast cancer; Magnetic resonance imaging; Computer-aided detection; Automated breast ultrasonography

INTRODUCTION

Automated breast ultrasonography (ABUS) has been developed in recent decades and proposed as a promising tool for overcoming the disadvantages of hand-held ultrasonography (HHUS) (1). Although HHUS technology has progressed and is still the

standard breast US method, it is operator-dependent and nonreproducible, and cannot display breast lesions in the coronal plane (2, 3).

ABUS was primarily developed as a screening tool. Unlike HHUS, it is less operator-dependent and can acquire full three-dimensional breast US volumes that are reproducible any time. To provide complete coverage of both breasts, ABUS consists of two to five acquisitions per breast, with each acquisition comprising over 300 transverse and reconstructed images in the coronal and sagittal planes. Consequently, the large ABUS volumes might induce oversight errors involving specific malignancies. Accurate computer-aided detection (CAD) software may facilitate the detection of cancers in ABUS data sets and enhance the efficiency of radiologist work flows (4).

Research using CAD for ABUS is not exhaustive, with few studies using CAD with ABUS for suspicious lesions detected on breast MRI.

The purpose of this study was to evaluate the diagnostic performance of CAD with ABUS for suspicious lesions detected on breast MRI and to analyze the nature of CAD-detected lesions and determine the factors triggering false readings.

MATERIALS AND METHODS

Patient Population

The Institutional Review Board approved the study. All patients provided informed consent for the study. Between January 1 and December 31, 2017, we prospectively enrolled 40 women (age range, 30–76 years; mean age, 50.4 ± 9.8 years) who underwent HHUS and ABUS for multiple suspicious lesions detected on preoperative MRI. The 40 patients carried a total of 122 suspicious lesions under categories 4, 5, and 6 based on the American College of Radiology Breast Imaging Reporting and Data System (ACR BI-RADS). Most lesions were ultimately diagnosed using ultrasound-guided needle biopsy or surgical results.

Of the 122 MRI-detected lesions, we excluded 51 daughter nodules located near the main breast cancer within the same quadrant, ultimately including 71 lesions: 49 identified as malignant, 15 as nonmalignant, and 7 lesions with no change or that disappeared on follow-up MRI. Among the 64 pathologically confirmed lesions, 58 were confirmed via ultrasound-guided biopsy, 4 via mammography-guided biopsy, 1 was confirmed via ultrasound-guided localization prior to surgery, and 1 was

confirmed via MRI-guided biopsy. The 49 lesions that were identified as malignant lesions included 43 cases of invasive ductal carcinoma (IDC), 2 involving ductal carcinoma *in situ* (DCIS), 3 mucinous carcinomas, and 1 case of invasive lobular carcinoma. Among the 15 nonmalignant lesions, including benign and borderline lesions, there were six cases of fibrocystic change, two of sclerosing adenosis, two of intraductal papilloma, 1 of fibroadenoma, 1 of stromal fibrosis, 1 of intramammary lymph node, 1 of ductal hyperplasia, and 1 of atypical ductal hyperplasia. We also analyzed 23 lesions showing false CAD readings: marginal or subareolar shadows manifesting as pseudo-lesions (18/23), typical benign cysts (3/23), a hematoma (1/23), and a skin wart (1/23). A final total of 94 lesions including 71 MRI-detected suspicious lesions and 23 pseudo-lesions with false CAD readings were analyzed (Table 1).

Breast MRI

All patients underwent MR imaging of the breast at our institution, performed with a 3T MR imaging system (Ingenia, Philips, Eindhoven, The Netherlands and Verio, Siemens Medical Solutions, Erlangen, Germany). Images of both breasts were obtained in the axial plane with the patient lying in the prone position. In this study, the following sequences were used: axial, turbo spin-echo T2-weighted imaging (repetition time [TR]/echo time [TE] 4530/93 ms, flip angle 80°, 34 slices with field of view [FOV] 320 mm, matrix 576 × 403, 1 number of excitations,

Table 1. Characteristics of Patients and Lesions

Total	94
Age	
Mean ± SD	50.4 ± 9.8
Median (Range)	49 (30–72)
Pathology	71
Benign	11 (11.7%)
Borderline	4 (4.3%)
Malignancy	49 (52.1%)
No change/disappeared on follow-up	7 (7.4%)
CAD marker for pseudo-lesion	23
Marginal or subareolar shadowing	18 (19.1%)
Cyst	3 (3.2%)
Hematoma	1 (1.1%)
Skin wart	1 (1.1%)

CAD = computer-aided detection; SD = standard deviation

4 mm slice thickness, acquisition time 2 minutes 28 second). Pre- and post-contrast, axial, T1-weighted flash three-dimensional (3D), volumetric interpolated breath-hold examination (TR/TE 4.4/1.7 ms, flip angle 10°, 1.2 mm slice thickness with no gap, and acquisition time 60 s). The post-contrast scans were obtained at 7, 67, 127, 187, 247, and 367 s after injection of 0.1 mmol/kg gadopentetate dimeglumine. The images were reformatted in the sagittal and coronal planes with additional maximum intensity projection representation.

ABUS and CAD Systems

ABUS was performed using the automated breast volume scanner ACUSON S2000 ABUS system (Siemens Medical Solutions, Mountain View, CA, USA) by an experienced radiographer. The radiographer selected the most suitable setting based on the patient's breast size (A-D and DD cups). With the patient in the supine position, anterior-posterior views of the bilateral breasts were initially scanned followed by scanning of the lateral and medial views, which primarily include the outer and the inner breast, in the oblique position. After the image acquisition, the 3D volume data were immediately sent to the workstation and reviewed in multiple orientations (transverse, coronal, and sagittal planes) using multi-planar reconstruction. The scan thickness was displayed at 1-mm intervals without overlap.

In this study, we used QVCAD™ CAD software (QVCAD system from QView Medical, Inc., Los Altos, CA, USA) to process and display ABUS images on a computer monitor for review with certain areas highlighted or marked for attention. It highlights suspicious regions displayed on an

intelligent minimum intensity projection image of the ABUS view.

Image Analysis

Two specialized breast radiologists with 9 and 11 years of experience, respectively, in academic breast imaging reviewed 40 patients' breast MRIs, and categorized each suspicious lesion by BI-RADS. The CAD-detected lesions that matched suspicious lesions on MRI, were identified and analyzed.

Statistical Analysis

We statistically analyzed the diagnostic accuracy of CAD, characteristics of CAD-detected lesions and the causes underlying false-positive and false-negative cases, and false readings in general. We calculated the diagnostic performance-sensitivity, specificity, positive predictive value (PPV), and negative predictive value (NPV)-using SAS version 9.2 (SAS Institute, Cary, NC, USA). We then analyzed the characteristics of the CAD-detected lesions using the chi-square test or the Fisher's exact test and calculated the P values, assuming significance at p less than 0.05.

RESULTS

The final proportions of malignant and benign lesions in this study were 52.1% (49/94) and 47.9% (45/94), respectively. CAD revealed 62 lesions (62/94, 65.9%) including 35 IDC lesions, 1 mucinous carcinoma, and 1 invasive lobular carcinoma (Table 2). Among the malignant

Table 2. Sensitivity and Specificity of Computer-Aided Detection (CAD) with Automated Breast Ultrasonography for Suspicious Lesions Detected on Breast MRI and CAD-False Lesions

	Malignancy			
	Benign	Malignancy	Sensitivity (95%)	Specificity (95%)
Total (n = 94)	45	49		
CAD (-)	20	12	75.5 (61.1-86.7)	44.4 (29.6-60.0)
CAD (+)	25	37		
	IDC only			
	Non IDC	IDC	Sensitivity (95%)	Specificity (95%)
Total (n = 94)	51	43		
CAD (-)	24	8	81.4 (66.6-91.6)	47.1 (32.9-61.5)
CAD (+)	27	35		

IDC = invasive ductal carcinoma
 Non IDC including ductal carcinoma *in situ*, mucinous carcinoma, and invasive lobular carcinoma.

lesions, we failed to detect 8 IDC lesions, 2 mucinous carcinomas, and 2 DCIS lesions using CAD.

The sensitivity, specificity, PPV, and NPV of CAD for automated breast ultrasound were 75.5% (37/49), 44.4% (20/45), 59.7% (37/62), and 62.5% (20/32), respectively. CAD facilitated the detection of 81.4% (35/43) of the invasive ductal cancers and 84.9% (28/33) of the invasive ductal cancers that showed masses (Table 3). CAD also revealed 90.3% (28/31) of the invasive ductal cancers measuring larger than 1 cm mass (excluding non-masses and lesions smaller than 1 cm). Among the 71 MRI-detected lesions, 55 were mass lesions on MRI, 15 were non-mass lesions, and 1 was missed. Masses larger than 1 cm in size were detected at a significantly higher rate ($P < 0.05$), even with invasive ductal cancers that showed masses (Table 4, Fig. 1).

False-negative lesions included 8 IDC lesions, 2 mucinous carcinomas, and 2 DCIS lesions. The mean sizes of the true-positive versus false-negative mass lesions were 2.08 ± 0.85 cm versus 1.6 ± 1.28 cm ($P < 0.05$). The false-positive lesions included a sclerosing adenosis and a case of usual ductal hyperplasia. Additionally, the false CAD marks for pseudo lesions involved 18 marginal or subareolar shadowings, 3 simple cysts, a hematoma, and a skin wart (Fig. 2). Table 4 also shows the correlation between CAD-

positive lesions based on MRI and ABUS findings.

DISCUSSION

HHUS showed promising results as a breast cancer screening tool in the past decade. However, it remains somewhat challenging and time-consuming for radiologists. ABUS may address these limitations by greatly reducing operator dependency and providing high-resolution 3D images (5). Although ABUS has many advantages over HHUS, it is still associated with a few disadvantages.

In a number of prior studies, the mean diameters of ABUS-detected lesions were larger than those of the handheld-detected lesions, with varying cut-offs (2, 6, 7). Therefore, it is possible that ABUS reflects the CAD detection sensitivity, especially in size. In this study, the mean sizes of the true-positive versus false-negative mass lesions were 2.08 ± 0.85 cm vs. 1.6 ± 1.28 cm ($P < 0.05$).

In previous studies, Chang et al. (8) reported statistically significant differences in ABUS detection rates of factors including margin, orientation, and boundary, and van Zelst et al. (9) suggested that a higher BI-RADS category indicated a higher detection rate on ABUS. In our study, the lesion shape, orientation, and margin showed statistically

Table 3. Sensitivity and Specificity of Computer-Aided Detection (CAD) for Breast Cancer According to Mass/Non-Mass Lesions and Size in MRI

	Malignancy				IDC only			
	Benign	Malignancy	Sensitivity (95%)	Specificity (95%)	Non IDC	IDC	Sensitivity (95%)	Specificity (95%)
Mass in MRI (n = 55)								
CAD (-)	16	8	78.4 (61.8-90.2)	88.9 (65.3-98.6)	19	5	84.9 (68.1-94.9)	86.4 (65.1-97.1)
CAD (+)	2	29			3	28		
Non-mass in MRI (n = 15)								
CAD (-)	4	4	63.6 (30.8-89.1)	100.0 (39.8-100.0)	5	3	66.7 (29.9-92.5)	83.3 (35.9-99.6)
CAD (+)	0	7			1	6		
Size in MRI: < 1 (n = 22)								
CAD (-)	16	4	20.0 (0.5-71.6)	94.1 (71.3-99.9)	17	3	25.0 (0.6-80.6)	94.4 (72.7-99.9)
CAD (+)	1	1			1	1		
Size in MRI: ≥ 1 (n = 49)								
CAD (-)	4	8	81.8 (67.3-91.8)	80.0 (28.4-99.5)	7	5	87.2 (72.6-95.7)	70.0 (34.8-93.3)
CAD (+)	1	36			3	34		

IDC = invasive ductal carcinoma

Non-IDC including ductal carcinoma *in situ*, mucinous carcinoma, and invasive lobular carcinoma

A single missing record involved mass/non-mass in MRI, and the sum was 70 (55+15) instead of 71.

significant differences in CAD detection rate ($P < 0.01$) on ABUS.

ABUS provides volumetric images with many slices and large FOV, which can lead to fatigue and missed radiological findings involving the large FOV, leading to interpretation errors, and ultimately decreasing the accuracy of detection. Skilled and accurate interpretation of ABUS requires considerable radiological training and experience (10). Therefore, a CAD system with ABUS may be more feasible for radiologists in practice. Previous studies have shown positive results supporting the diagnostic utility of CAD with ABUS (4, 11). As shown in our study, the sensitivity, specificity, PPV, and NPV of CAD with automated breast ultrasound were 75.5%, 44.4%, 59.7%, and 62.5%, respectively. The sensitivity of this study was relatively low (75.5%), because we used the CAD with ABUS for suspicious lesions detected on breast MRI. Breast MRI has proven to be the most sensitive tool for detecting breast carcinoma (12).

Therefore, we enrolled patients with multiple suspicious lesions on preoperative MRI, and included many small and subtle lesions in this study.

Our study analyzed the diagnostic performance of ABUS with CAD for lesions detected using MRI. MR-directed ultrasound, also referred to as second-look ultrasound and targeted ultrasound, has been shown to be useful in identifying lesions detected initially on MRI. In one previous study, the role of MRI was indispensable for preoperative assessment, although the combination of HHUS and ABUS showed the highest sensitivity. In addition, ABUS imaging was better than HHUS for preoperative evaluation (6).

We analyzed MRI lesions that correlated with detection rates on CAD with ABUS, and the detection rate was slightly higher if a lesion on MRI was defined as a mass (31/55, 56.4%) rather than a non-mass (7/15, 46.7%) lesion. This result suggests the same trend, albeit weaker, as that of a previous study in which the HHUS detection rates were

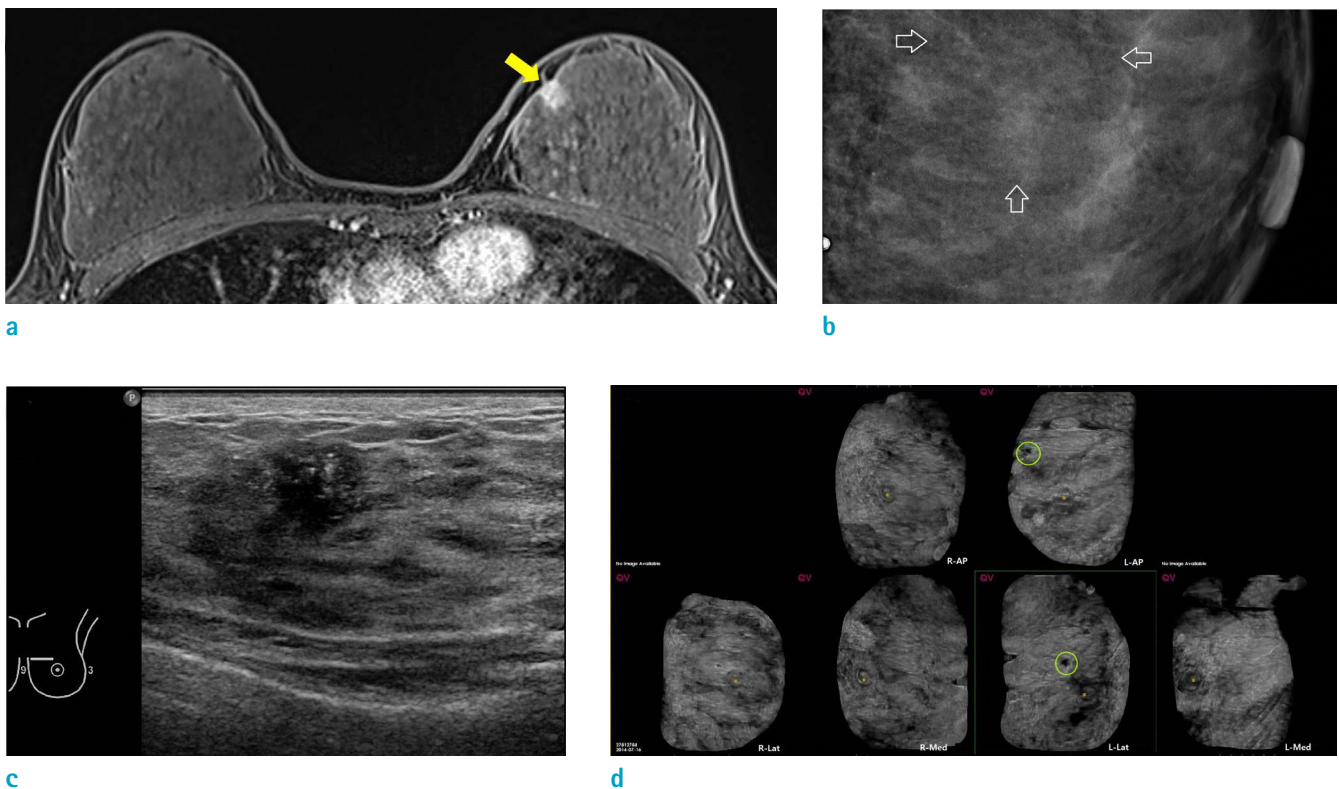


Fig. 1. Images from a 32-year-old woman with a suspicious lesion detected on MRI and investigated subsequently with automated breast ultrasound (ABUS). (a) MRI image showed an approximately 1.1 cm enhancing mass at the 10 o'clock position on the left breast (arrow). (b) Mammography showed microcalcifications with suspicious architectural distortion involving the upper left inner quadrant (arrows). (c) Handheld ultrasonography revealed about 1.1-cm irregular mass with microcalcifications in the same direction. (d) 3D ABUS revealed correlating suspicious lesion on CAD in the left AP and medial views, later confirmed as invasive ductal carcinoma.

58% for mass versus 29% for non-mass lesions (13, 14), and the pathologically invasion by malignant lesions with space-occupying mass formation showed a higher detection

rate. In our study, 81.4% (35/43) of the invasive ductal cancer was detected using CAD. CAD was used to detect 84.9% (28/33) of the invasive ductal mass cancers (exclude

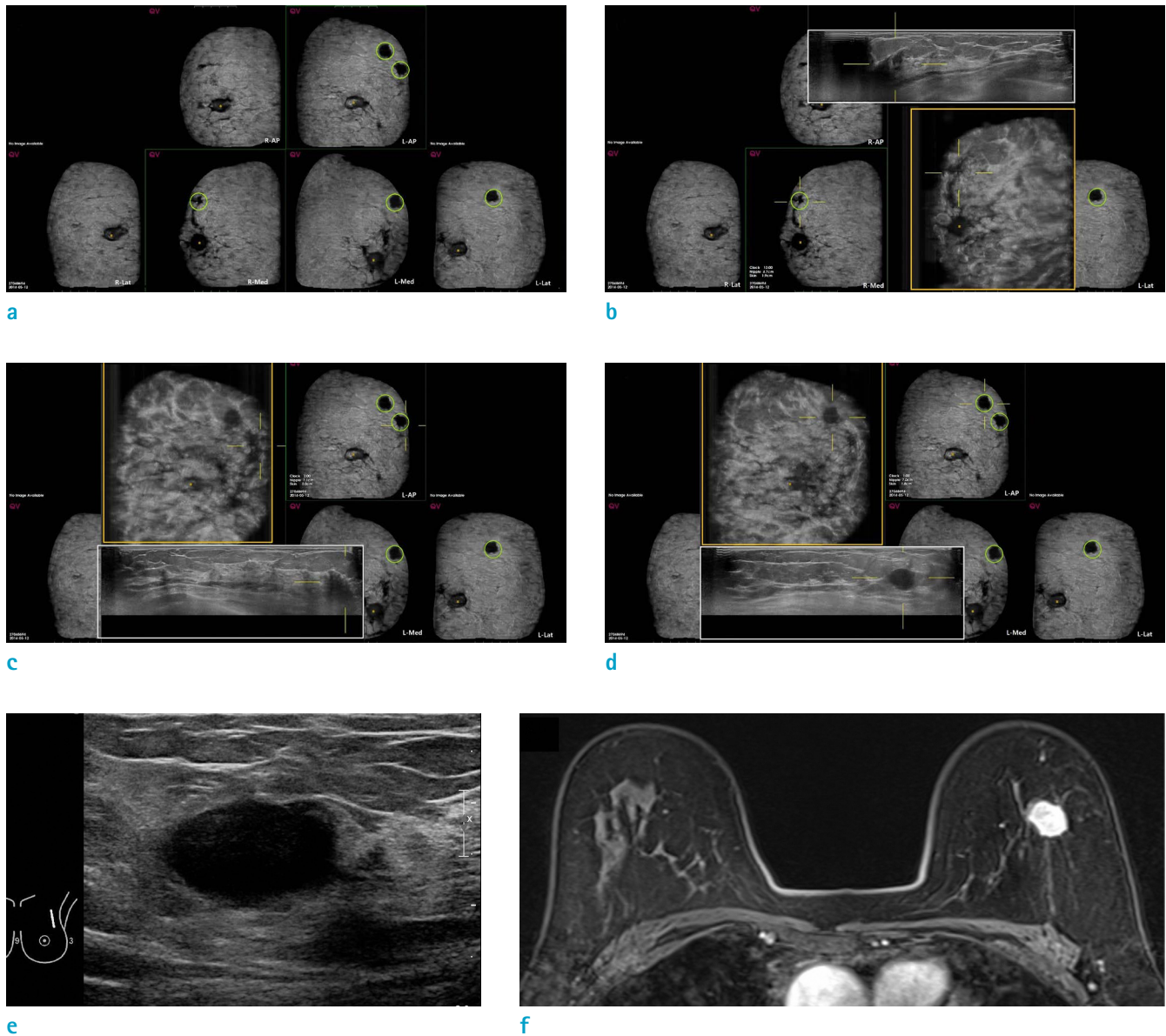


Fig. 2. A 62-year-old woman who showed three CAD-detected lesions in both breasts. (a) CAD revealed a suspicious lesion in the right breast and only one marked lesion in the right medial view. The other two marked lesions involved the left breast: one was only marked in the left AP view, and the other one was marked in the whole 3D views. (b) Axial (white box) and maximal intensity projection (MIP) reconstruction image (yellow box) shows a right breast lesion that was confirmed as a pseudo lesion based on marginal shadowing. (c) Axial (white box) and MIP reconstruction image (yellow box) of one of the left breast lesions, which was marked by CAD only on AP view; it was also a pseudo lesion due to marginal shadowing. (d) Axial (white box) and MIP reconstruction image (yellow box) of the other left breast lesion, which was entirely marked by CAD in 3D view, and confirmed as IDC. (e) HHUS image of a biopsy-proven IDC lesion involving left breast shows a 1.8-cm marked hypoechoic mass with microlobulation in the left 2-h direction. (f) MRI of biopsy-proven IDC lesion shows 1.7-cm markedly enhanced mass in the left breast at the 1 o'clock position.

Table 4. Correlation between Computer-Aided Detection (CAD)-Positive Lesions Based on MRI and Automated Breast Ultrasonography (ABUS)

	N (%)	Missing	Total	CAD (-)	CAD (+)	P value
Size in MRI						
< 1		0 (0)	22 (31.0)	20 (90.9)	2 (9.1)	< 0.001
≥ 1			49 (69.0)	12 (24.5)	37 (75.5)	
Mass/non-mass in MRI						
mass		1 (1.4)	55 (77.5)	24 (43.6)	31 (56.4)	0.504
non-mass			15 (21.1)	8 (53.3)	7 (46.7)	
Mass shape in MRI						
oval		16 (22.5)	25 (35.2)	15 (60.0)	10 (40.0)	0.0255
round, irregular			30 (42.3)	9 (30.0)	21 (70.0)	
Mass margin in MRI						
circumscribed		16 (22.5)	17 (23.9)	14 (82.4)	3 (17.6)	0.0001
not circumscribed			38 (53.5)	10 (26.3)	28 (73.7)	
Internal pattern in MRI						
homogeneous		1 (1.4)	20 (28.2)	15 (75.0)	5 (25.0)	0.0019
not homogeneous			50 (70.4)	17 (34.0)	33 (66.0)	
Non-mass distribution in MRI						
not segmental, linear		56 (78.9)	6 (8.5)	0 (0.0)	6 (100.0)	0.0014
segmental, linear			9 (12.7)	8 (88.9)	1 (11.1)	
Kinetic-initial in MRI						
not fast		1 (1.4)	23 (32.4)	11 (47.8)	12 (52.2)	0.804
fast			47 (66.2)	21 (44.7)	26 (55.3)	
Kinetic-delayed in MRI						
not washout		1 (1.4)	34 (47.9)	12 (35.3)	22 (64.7)	0.089
washout			36 (50.7)	20 (55.6)	16 (44.4)	
Shape in ABUS						
oval		4 (5.6)	15 (21.1)	13 (86.7)	2 (13.3)	0.0001
round, irregular			52 (73.2)	16 (30.8)	36 (69.2)	
Orientation in ABUS						
parallel		4 (5.6)	48 (67.6)	25 (52.1)	23 (47.9)	0.0209
not parallel			19 (26.8)	4 (21.1)	15 (78.9)	
Margin in ABUS						
circumscribed		4 (5.6)	16 (22.5)	15 (93.8)	1 (6.3)	<0.001
not circumscribed			51 (71.8)	14 (27.5)	37 (72.5)	
Echo pattern in ABUS						
an, iso, hyper		4 (5.6)	7 (9.9)	5 (71.4)	2 (28.6)	0.2251
hypo, heterogenous, complex			60 (84.5)	24 (40.0)	36 (60.0)	
Posterior feature in ABUS						
no, enhance		4 (5.6)	55 (77.5)	26 (47.3)	29 (52.7)	0.1583
shadow, complex			12 (16.9)	3 (25.0)	9 (75.0)	

non-mass). We found a statistically significant correlation between CAD with ABUS detection rate and the size of the lesions characterized as masses on MRI. Masses measuring greater than 1 cm in size were more likely to be detected on CAD, although we did not find this association between size and the likelihood of non-mass lesions detected on MRI. The discrepancies between mass and non-mass lesions may be because mass lesions, defined as three-dimensional space-occupying lesions, are more readily defined with CAD on ABUS.

In this study, we found 23 false CAD marks. The false CAD marks for pseudo lesions were easily identified as benign rather than suspicious. The false readings on CAD were attributed to typical marginal or subareolar shadowing, simple cysts, a cystic hematoma, and a skin wart, not mimicking a true breast lesion.

Our study has a few limitations. First, the study comprised a relatively small sample that lacked representativeness. Second, in some cases biopsy was not confirmed, and this might have affected outcomes. Third, two radiologists independently reviewed the MRI and HHUS and MRI and ABUS; however, this study did not provide the radiologists' performance analysis such as inter-observer agreement. In this study, we focused on the diagnostic performance of CAD with ABUS and interpreted the lesions associated with errors in the CAD system. The last limitation of this study is that we were not always confident about the precise MRI-ultrasound correlations. However, we stipulated stringent criteria to determine the sonographic correlation, such as lesion position, size, and similarity in shape. As such, we can guarantee true MRI-ultrasound correlations.

In this study, we determined the diagnostic performance of CAD with ABUS for suspicious lesions detected on breast MRI, the characteristics of CAD-detected lesions, and the causes of false positives, false negatives, and false CAD marks. In conclusion, it was not difficult for operators familiar with CAD combined with ABUS to identify false CAD marks for pseudo lesions. We demonstrated that CAD with ABUS showed a reliable diagnostic performance for invasive ductal cancers involving masses larger than 1 cm.

Acknowledgments

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