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The role of breast tomosynthesis in a predominantly dense breast population at a tertiary breast centre: breast density assessment and diagnostic performance in comparison with MRI

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Abstract

Objectives To compare breast density measured on digital breast tomosynthesis (DBT) (BI-RADS-based breast composition and fully-automatic estimation) and magnetic resonance imaging (MRI) (BI-RADS amount of fibroglandular tissue), and to evaluate the diagnostic performance in terms of sensitivity and specificity of DBT and MRI in a predominantly dense breast population.

Methods Between 2015 and 2016, 152 women with 103 breast malignancies, who underwent 3-T breast MRI and DBT within 2 months’ time, were enrolled in this study. Breast composition/fibroglandular tissue and findings on DBT (two readers) and MRI were reported using BI-RADS 5th edition. Digital mammography images were analysed for breast percent density (PD) using the Libra software tool.

Results A majority of women had dense breasts as categorised by breast composition c (heterogeneously dense) (68%) and d (extremely dense) (15%). The mean PD was 44% (range, 18-89%) and the correlation between breast composition and PD was $r = 0.6$. The diagnostic performance of MRI was significantly higher compared to DBT for one reader as described by the area under the receiver operating characteristic (ROC) curve ($p = 0.004$) and of borderline significance for the other reader ($p = 0.052$).

Conclusions MRI had higher diagnostic performance than DBT in a dense breast population in the tertiary setting.

Key Points

- MRI had higher diagnostic performance than DBT in a dense breast population
- Diagnostic performance of DBT was comparable to MRI in women with fatty breasts
- MRI was superior to DBT in preoperative breast cancer size assessment

Keywords Breast cancer · Breast density · Digital breast tomosynthesis · Magnetic resonance imaging · Diagnostic techniques and procedures

Abbreviations

DBT Digital breast tomosynthesis
PD Percent density

DM Digital mammography
FGT Fibroglandular tissue
BPE Background parenchymal enhancement
FN False negative
NST No special type
DCIS Ductal carcinoma in situ
ILC Invasive lobular carcinoma
FP False positive

Introduction

Digital breast tomosynthesis (DBT) has gained increasing interest in the clinic for routine everyday use, with several early clinical studies showing superior accuracy compared to that of mammography [1–3]. Large multicentre studies have shown
general benefits of DBT over digital mammography (DM), but the role of tomosynthesis in women with dense breasts has not yet been fully established [4–6]. Prospective trials of screening a population from Europe show a statistically significant increase in cancer detection rate with DBT (with or without combined DM) compared with two-view DM independent of vendor system [7–9]. Skaane et al. [7] stated: “Notably, the additional cancers detected with mammography plus tomosynthesis were distributed across all breast densities, including fatty breasts”. Lång et al. [9] reported that the additional cancers were detected both in women with dense and fatty breasts, drawing the conclusion that in mammography even a moderate amount of breast tissue can conceal a small lesion.

Breast cancer risk increases with increasing breast density [10, 11]. The relationship holds true also for Japanese women [12, 13], who are known to have dense breasts [14] and highest age-specific breast cancer incidence at age 45–49 [15]. Breast density has until recently been classified qualitatively by use of the four categories in the ACR Breast Imaging Reporting and Data System (BI-RADS) coding system: (1) almost entirely fat (<25% fibroglandular tissue), (2) scattered fibroglandular densities (25–50% fibroglandular tissue), (3) heterogeneously dense (51–75% fibroglandular tissue) and (4) extremely dense (>75% fibroglandular tissue) [16]. Recent revision of BI-RADS concerns changes to category 3: the breasts are heterogeneously dense, which may obscure small masses, and discourage the use of a cut-off based on percent density [17]. In order to more objectively assess breast density and to reduce interobserver and intraobserver variability, quantitative measurements have been developed [18, 19]. Both qualitative and quantitative methods of measuring breast density have shown an association between high breast density and breast cancer risk [11, 20].

More than half of the USA states have begun legislating mammographic breast density reporting to women, requiring that women be notified of breast density with their mammography results [21]. As a result, the U.S. Preventive Service Task Force conducted a review of supplemental screening for breast cancer in women with dense breasts and concluded that density ratings may be recategorised on serial screening mammography and, with limited evidence, that additional imaging with DBT reduces recall rates in women with dense breasts; however, it was not clear the results implied fewer overall breast biopsies [22].

Dynamic contrast-enhanced breast (DCE) MRI has excellent sensitivity independent of breast density but still moderate specificity [22, 23]. By adding diffusion-weighted imaging (DWI) sequences, there is optimism to increase the specificity [24]. Recommended indications for DCE MRI on defined subgroups include: screening of high-risk women due to cancer susceptibility genes or greater than 20% lifetime risk of developing breast cancer, assessment of occult primary breast cancer, preoperative staging and evaluation of neoadjuvant therapy [25, 26]. MRI is expensive and time-consuming, hence the use of MRI in the clinic varies mainly due to accessibility.

The purpose of this study was to compare breast density measured on DBT and MRI, and to evaluate the diagnostic performance in terms of sensitivity and specificity of DBT and MRI in a predominantly dense breast population in the tertiary setting.

Materials and methods

Patient population and lesions

Approval was obtained from the Institutional Review Board of our institution with a waiver of informed consent due to the retrospective nature of the study. Between March 2015 and March 2016, a total of 494 patients underwent 548 MRI examinations and 2,164 patients underwent 2,292 DBT examinations at the Department of Radiology, Kyoto University Hospital, Japan. The inclusion criteria included matched MRI/DBT patient examinations within 2 months’ time (n = 249) and BI-RADS diagnostic category 3, 4 or 5 on MRI, resulting in 162 patients. Of these, 10 were excluded from analysis because of previous breast surgery (n = 5), incomplete MRI examination (n = 2), insufficient follow-up (n = 2) or Paget’s disease (n = 1). In total, 152 female patients (mean age, 57.1 ± 13.8 years; range, 28–83 years), of which 56 were healthy and 96 presented with 103 breast malignancies including 4 bilateral and 3 multicentric, were enrolled in this study. All breast cancer diagnoses were verified by surgery and/or biopsy. Patients were considered healthy after negative surgery and/or biopsy or at least 1-year negative imaging follow-up on digital mammography and ultrasound (US) and/or MRI. The MRI sample represented 31% (152/494) and the DBT sample 7% (152/2,164) of the total number of women being examined during the actual period of time. MRI was performed on average 8.7 ± 10.6 days (range, 46 to 50 days) after DBT.

All women referred for clinical assessment during the time period of the study had DBT as part of their examination, unless contraindicated or DBT was recently acquired at another hospital. Indications for MRI in this study population were mostly problem solving (n = 130; 85.5%) with the following details: suspicious finding by DM (n = 29; 19.1%), suspicious finding by US (n = 59; 38.8%), suspicious finding by DM and US (n = 31; 20.4%), suspicious finding by computed tomography (n = 11; 7.2%). Other indications included high-risk women (n = 2; 1.3%) and preoperative staging (n = 20; 13.2%).

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Imaging acquisition

All patients included underwent bilateral two views (craniocaudal and mediolateral oblique) using a full-field DM unit with tomosynthesis capability (Selenia Dimensions; Hologic, Bedford, MA, USA) operating in combo mode: DM and DBT images were obtained within a single breast compression for each projection. W/Rh and W/Al were used as anode/filter combinations for DM and DBT, respectively, automatic exposure control was employed and 15 low-dose images were acquired during the X-ray movement over an angular range of 15°. These low-dose images were reconstructed into millimetre-thin slices of the breast using filtered back-projection. The mean compressed breast thickness was 34.4 ± 12.6 mm (range, 8.5-73.5 mm).

The MRI examinations were performed with a 3.0-T scanner (MAGNETOM Trio, A Tim System; Siemens, Erlangen, Germany) with 16/4 channel breast coil. The parameters were as follows: T2-weighted images (whole breast; axial orientation; 2D-turbo spin echo with fat suppression; repetition time/echo time (TR/TE), 5,500/70 ms; field-of-view (FOV), 330 × 330 mm; matrix, 448 × 336; thickness, 3.0 mm), T1-weighted images (whole breast; axial orientation; 3D volumetric interpolated breath-hold examination (3D-VIBE); TR/TE, 4.95/2.46 ms; FOV, 330 × 330 mm; matrix, 480 × 398; thickness, 1.5 mm), T1-weighted DCE images scanning at pre-contrast state and at 0–1, 1–2 and 5–6 min after gadolinium injection (whole breast; axial orientation; 3D-VIBE with fat suppression; TR/TE, 3.95/1.43 ms; flip angle (FA), 15°; FOV, 330 × 330 mm; matrix, 384 × 346; thickness, 1.0 mm), contrast-enhanced T1-weighted images in high spatial resolution at 2–4.5 min after gadolinium injection (whole breast; coronal orientation; 3D-VIBE with fat suppression; TR/TE, 4.61/1.81 ms; FA, 15°; FOV, 330 × 330 mm; matrix, 512 × 461; thickness, 0.8 mm). The MR examination was performed during first half of the menstrual cycle in premenopausal women and without scheduling limitations in postmenopausal women.

Image interpretation and analysis

Two breast radiologists (M.I. and M.K. with 10 and 18 years of experience in mammography, respectively) reviewed and assessed the DBT images according to BI-RADS 5th edition on a dedicated workstation unaware of any clinical information or the histopathology diagnosis [17]. Breast composition (based on mammography projections) and findings on DBT were characterised and recorded. BI-RADS categories 1-3 were considered benign, and BI-RADS categories 4 and 5 were considered malignant. The MR images were assessed and approved at the time of clinical interpretation by one of two specialised breast radiologists (M.K. and S.K.) in accordance with BI-RADS. Fibroglandular tissue (FGT), background parenchymal enhancement (BPE), findings and enhancement characteristics, including kinetic curve assessment, were characterised and recorded. The software package LIBRA, a fully-automatic breast density estimation software solution, was used in this study to estimate the percentage breast density (PD) [27, 28]. The average PD of the four DM images was calculated and in cases containing larger (>30 mm) invasive tumours, only the average of the contralateral values was used (the size cut-off was chosen not to exceed the natural dense breast tissue variation between the two breasts, data not shown).

A preoperative tumour size assessment was performed by retrospectively measuring the largest lesion diameter to the nearest millimetre at the central core on both DBT and MR images to establish which imaging modality most accurately corresponds with the size of the pathology report [29, 30]. Tumours were classified as not measurable in cases of architectural distortion without definable borders or in cases where the tumour was partially obscured by dense tissue on DBT and in cases of multiple or diffuse enhancement patterns (mostly classified as non-mass enhancement) within the breast on MRI. Only invasive cancers were included in the analysis.

Statistical analysis

Inter-rater reliability was measured using percent agreement and weighted kappa statistics. Pearson’s r and Spearman’s ρ were used to test for correlations for normally and skewed distributed data, respectively. One-way analysis of variance with Bonferroni correction was used to compare PD means between groups. Diagnostic performance was assessed with receiver operating characteristics (ROC) curve analysis [31]. Area under the curve (AUC) was calculated from both parametric and trapezoidal curve fitting. Fisher’s exact test and logistic regression analysis were performed to determine which variables are associated with false negative (FN) outcomes. Size agreement between imaging modality and pathology measurements were analysed following the approach of Bland and Altman. All analyses were performed using the SPSS software (version 24; IBM Corp., Armonk, NY, USA) and p values <0.05 were considered statistically significant.

Results

Out of all 103 breast cancers, 64 were invasive carcinoma of no special type (NST), 26 ductal carcinoma in situ (DCIS), 6 invasive lobular carcinoma (ILC), 4 mucinous carcinoma, 1 micropapillary carcinoma, 1 tubular carcinoma and 1 apocrine carcinoma.

The average percentage distribution of breast composition (based on DBT) was: 2.3% (n = 3.5) category a, 15.5% (n = 23.5) category b, 67.8% (n = 103) category c, 14.5% (n = 22)
category d. There was substantial inter-rater agreement between the two readers (R1 and R2) with percent agreement of 85% (129/152) and weighted $\kappa = 0.74$ [95% confidence interval (CI), 0.64-0.84].

The percentage distribution of amount of FGT (based on MRI) was: 5.9% ($n = 9$) a. Almost entirely fat, 30.3% ($n = 46$) b. Scattered fibroglandular tissue, 53.3% ($n = 81$) c. Heterogeneous fibroglandular tissue, 10.5% ($n = 16$) d. Extreme fibroglandular tissue.

The percentage distribution of background parenchymal enhancement (BPE) level (based on MRI) was: 22.1% ($n = 31$) minimal, 32.1% ($n = 45$) mild, 26.4% ($n = 37$) moderate, 19.3% ($n = 27$) marked.

The mean PD was 43.5% (range, 18.3-88.6%). The average mean PD for the different breast compositions was: a = 23.4% (range, 18.3-26.4%), b = 31.7% (range, 18.8-51.5%), c = 42.4% (range, 18.8-88.6%) and d = 64.8% (range, 41.1-84.3%) (Fig. 1). All category means were significantly different ($p < 0.009$), except for a with b ($p = 1.00$).

The correlations between breast composition, FGT, BPE and PD ranged from weak to strong (Table 1).

On DBT, R1 had a sensitivity of 80.6% (FN = 20) and positive predictive value (PPV) of 76.1% (FP = 26) and R2 had a sensitivity of 82.5% (FN = 18) and PPV of 74.6% (FP = 29). The AUC of the parametric ROC curve was 0.875 (95% CI, 0.801-0.927) and 0.906 (95% CI, 0.852-0.944), and the AUC of the trapezoidal ROC was 0.872 (95% CI, 0.823-0.920) and 0.886 (95% CI, 0.841-0.930) for R1 and R2, respectively (Fig. 2). MRI had a sensitivity of 97.1% (FN = 3) and PPV of 62.5% (FP = 60). The AUC of the parametric ROC curve was 0.964 (95% CI, 0.931-0.983) and the AUC of the trapezoidal ROC was 0.922 (95% CI, 0.893-0.952). The

![Fig. 1 BI-RADS breast composition categories with corresponding breast percent density as estimated by Libra](image)

**Table 1** Bivariate correlations

| Breast composition | FGT       | BPE       | PD        |
|--------------------|-----------|-----------|-----------|
| Reader 1           | 0.53 (0.40-0.64) | 0.28 (0.11-0.45) | 0.57 (0.46-0.67) |
| Breast composition | 0.54 (0.41-0.65) | 0.28 (0.09-0.44) | 0.63 (0.53-0.72) |
| Reader 2           | 0.37 (0.22-0.53) | 0.51 (0.38-0.63) | 0.37 (0.22-0.51) |

*Numbers in parenthesis are 95% CIs*

*FGT* fibroglandular tissue, *BPE* background parenchymal enhancement, *PD* percent density
diagnostic performance of MRI was significantly higher compared to DBT for R1 using the parametric model ($p = 0.004$) and borderline higher compared to DBT for R2 ($p = 0.052$). R2 rated one false positive (FP) as BI-RADS 5 on DBT and one case of FP was rated as BI-RADS 5 on MRI. Twenty-five percent ($n = 5$) and 33% ($n = 6$) of the FN on DBT were rated as BI-RADS 3 by R1 and R2, respectively, and 33% ($n = 1$) of the FN on MRI was rated as BI-RADS 3.

By substituting MRI with DBT for women with mammography breast composition category a or b, the MRI FP could be reduced on average by 6% (R1, $n = 4$, R2, $n = 3$) at the cost of a 1% average reduced sensitivity (R1, $n = 0$; R2, $n = 2$). By applying PD threshold (<23.5%), a FP decrease of 3% (R1, $n = 2$; R2, $n = 2$) can be achieved at no sensitivity loss.

**Patient characteristics**

In general, FP were highly correlated on the two systems, 77% (20/26, R1) and 76% (22/29, R2).

No patient characteristics, including PD, were associated with FN on DBT (Table 2). When dichotomised as fatty (category a + b) or dense (category c + d), breast composition suggested an association with FN on DBT, although not reaching statistical significance, for R1 ($p = 0.080$). There was no association with FN on DBT for R2 ($p = 0.366$). No FN on DBT was found in category a or b for R1 and two FNs in category b (rated as c by R1) for R2. The lowest PD encountered for an FN was 23.5% for both readers.

The three FN cases on MRI were as follows: 45-year-old woman with heterogeneous fibroglandular tissue and marked BPE presenting DCIS, BI-RADS 3 (BI-RADS 1 on DBT by both readers), 56-year-old woman with scattered fibroglandular tissue and mild BPE, presenting DCIS, BI-RADS 2 (BI-RADS 4 on DBT by both readers) and 37-year-old woman with heterogeneous fibroglandular tissue and marked BPE presenting a 15-mm, grade 1, luminal a, mucinous carcinoma, BI-RADS 3 (BI-RADS 1 on DBT by both readers) (Fig. 3). Due to the low number of FNs on MRI no statistical tests were performed.

**Lesion characteristics**

The mean size 13.2 mm and 11.8 mm of FN was significantly smaller than the mean size 20.6 mm and 21.2 mm of TP on DBT for one reader (R1, $p = 0.115$ and R2, $p = 0.041$). No other cancer characteristics were associated with FN by any reader ($p > 0.05$); however, there was a trend for lobular histological type being missed (Table 2).

Of the 77 invasive masses, 12 underwent neoadjuvant chemotherapy and 1 case metastasised, leaving 64 masses with a pathological mean size of 18.3 mm (median, 15 mm; range, 1-65 mm). DBT could measure the invasive part in 57.8% (37/64) and MRI in 89.1% (57/64) of the cases with size correlation $r = 0.73$ and $r = 0.78$ versus pathology, respectively. One reason for the low DBT percentage was because of the high proportion of masses identified only by the calcification component ($n = 11$) without size comparison to pathology. Figure 4 shows the size deviations of the tumours measured with DBT and MRI compared with the averages of the pathology and said modality, respectively.

**Discussion**

The diagnostic performance of MRI was significantly higher compared to DBT for one reader ($p = 0.004$) and of borderline significance for the other reader ($p = 0.052$). Only a few studies have compared the diagnostic performance of DBT versus MRI in specific settings [32–35]. Kim et al. [35] reported higher diagnostic performance (sensitivity, 97.8%) but lower PPV (89.6%) for MRI versus DBT (sensitivity, 88.2%; PPV, 93.3%), which is in line with the results of this study. Clauser
Table 2  Univariate associations of patient and lesion characteristics with false negatives on DBT

| Patient characteristics | True positive | False negative | Significancea |
|-------------------------|---------------|----------------|---------------|
|                         | R1 R2 | R1 R2 | R1 R2 |
| Age                     | OR = 1.02 95% CI = 0.98-1.05 | OR = 1.01 95% CI = 0.98-1.05 | p = 0.334 p = 0.503 |
| Breast composition      |     |     |     |
| a (fatty)               | 3 (2) | 4 (3) | 0 (0) | 0 (0) |
| b                       | 19 (14) | 26 (19) | 0 (0) | 2 (11) |
| c                       | 93 (71) | 86 (64) | 15 (75) | 12 (67) |
| d (dense)               | 17 (13) | 18 (13) | 5 (25) | 4 (22) |
| Breast composition      |     |     |     |
| a + b (fatty)           | 22 (17) | 30 (22) | 0 (0) | 2 (11) |
| c + d (dense)           | 110 (83) | 104 (78) | 20 (100) | 16 (89) |
| FGT                     |     |     |     |
| a (fatty)               | 9 (7) | 9 (7) | 0 (0) | 0 (0) |
| b                       | 41 (31) | 39 (29) | 5 (25) | 7 (39) |
| c                       | 68 (52) | 72 (54) | 13 (65) | 9 (50) |
| d (extreme)             | 14 (11) | 14 (10) | 2 (10) | 2 (11) |
| BPE                     |     |     |     |
| Minimal                 | 28 (23) | 28 (23) | 3 (17) | 3 (19) |
| Mild                    | 37 (30) | 39 (32) | 8 (44) | 6 (38) |
| Moderate                | 34 (28) | 35 (28) | 3 (17) | 2 (13) |
| Marked                  | 23 (19) | 22 (18) | 4 (22) | 5 (31) |
| PD                      |     |     |     |
| Breast thickness        | OR = 1.02 95% CI = 0.99-1.05 | OR = 1.01 95% CI = 0.98-1.04 | p = 0.277 p = 0.553 |
| Lesion characteristics  | True positive | False negative |     |
| Size                    | OR = 0.95 95% CI = 0.90-1.02 | OR = 0.93 95% CI = 0.87-1.00 | p = 0.115 p = 0.041 |
| Type of finding         |     |     |     |
| Invasive                | 64 (77) | 63 (74) | 13 (65) | 14 (78) |
| In-situ                 | 19 (23) | 22 (26) | 7 (35) | 4 (22) |
| Histological type       |     |     |     |
| Ductal                  | 55 (86) | 54 (86) | 9 (69) | 10 (71) |
| Lobular                 | 4 (6) | 4 (6) | 2 (15) | 2 (14) |
| Other                   | 5 (8) | 5 (8) | 2 (15) | 2 (14) |
| Histological grade      |     |     |     |
| Grade 1                 | 21 (33) | 20 (32) | 3 (23) | 4 (29) |
| Grade 2                 | 24 (38) | 25 (40) | 5 (39) | 4 (29) |
| Grade 3                 | 19 (30) | 18 (29) | 5 (39) | 6 (43) |
| Subtypeb                |     |     |     |
| Luminal a               | 36 (56) | 36 (57) | 9 (69) | 9 (64) |
| Luminal b HER2-         | 13 (20) | 13 (21) | 2 (15) | 2 (14) |
| Luminal b HER2+         | 6 (9) | 6 (10) | 0 (0) | 0 (0) |
| HER2+ non-luminal       | 2 (3) | 2 (3) | 0 (0) | 0 (0) |
| Triple negative         | 7 (11) | 6 (10) | 2 (15) | 3 (21) |

Numbers in parentheses are percentages

R reader, FGT fibroglandular tissue, BPE background parenchymal enhancement, PD percent density, OR odds ratio, CI confidence interval

\[a\] Logistic regression and Fisher’s exact test were used for continuous and discrete data, respectively

\[b\] St Gallen 2013 classification
et al. [33] and Mariscotti et al. [34] investigated the role of DBT as a so-called second-look modality for additional findings at preoperative MRI, observing an increased detection rate when adding DBT to US, although, still inferior to MRI.

Breast composition, although not statistically significant, tend to be associated with FN on DBT (R1, \( p = 0.080 \)) in contrast to PD. Thus, a masking effect (particularly the use of category c) seems to be prominent regardless of specific percentage density thresholds. The moderate to strong correlation between breast composition and PD also suggests this finding. Since 68% of the women in this study were assigned breast composition c, it would be of great interest to explore any sub categorisation based on, for instance, parenchymal distribution or texture.

The prospective screening trials have not shown a significant cancer detection increase for DBT in category d, owing to the low sample size; however, there was no significant detection increase in breast composition d for DBT compared with DM alone in a multicentre study by Rafferty et al. [36], despite a larger sample size [7–9]. This lack of evidence is the main reason that DBT is still on hold as a complementary tool for the screening of women with dense breasts [22].

Eighty-two percent of the study population had dense breasts as categorised by breast compositions c and d and the density distribution was also significantly different from the anticipated 10%/40%/40%/10% by ACR BI-RADS in a screening setting [17]. As a comparison, data from the Breast Cancer Surveillance Consortium [37] showed that 43.3% of the US women are considered dense. The distribution of PD, as estimated by Libra within the radiologist-provided breast composition categories (Fig. 1.), resembled those in the work of Keller et al. [27], except for category c demonstrating a broader range. This is likely a result of the adoption of the new ACR BI-RADS category c definition, taking into account the masking effect. Comparing this Japanese study population (mean age, 57 years) with black and white Pennsylvanian women (n = 9,498; mean age, 57 years) regarding PD estimations using Libra, the population in this study had more than double PD, 44% versus 12.3% (black) and 17.1% (white) [38]. Despite not being a screening population, this result supports the view that Japanese women have dense breasts [12].

The literature is sparse regarding correlation of qualitative imaging assessments of FGT, BPE and mammographic breast composition. Hansen et al. [39] found a correlation of 0.36 between BPE and mammographic breast composition compared to 0.28 in this study. King et al. [40] found a non-significant correlation for one reader and a significant correlation of 0.4 for another between FGT and BPE, compared to 0.37 in this study. No studies were found correlating qualitative FGT and mammographic breast composition, which were considered moderate in this study (\( r = 0.53 \) and \( r = 0.54 \) for R1 and R2, respectively).

The preoperative size assessment was superior for MRI compared to DBT, showing a stronger correlation and a better size agreement with pathology (Fig. 4.). The DBT correlation coefficient of 0.73 was a little lower compared with other published studies: 0.86 by Förnvik et al. [29] and 0.86 by Luparia et al. [30], which could be explained by the predominantly dense breast population or the use of a narrow-angle DBT system. Chudgar et al. [41] assessed the use of preoperative MRI for disease extent in breast cancer detected at DBT versus DM alone and found that women with fatty breasts screened with DBT may benefit less from preoperative MRI than women with more dense breasts. By applying a PD threshold based on the FN, about 10% (13/137) of the study population...
population could potentially undergo DBT instead of MRI without any loss in sensitivity. Using breast composition groups a and b would increase specificity at a slight cost of sensitivity.

A limitation of the current study was the exclusion of BI-RADS 1 and 2 cases because of the sporadic follow-up in these groups, especially in the tertiary setting. Consequently, this could bias PPV in favour of DBT and overestimate MRI sensitivity. The relatively short minimum follow-up of 1 year for negative cases not undergoing biopsy/surgery is also a limitation of the current study. This study has followed the recommendations of BI-RADS 5th edition; thus, care should be taken when comparing the results with older BI-RADS editions, particularly regarding the breast density classification. The study population may also differ between a tertiary setting and a non-tertiary setting with regard to breast cancer profile and breast density. Another limitation was the final decision-making by one breast specialist for the MRI interpretation limiting the generalisation of the results. In addition, priors were not read during DBT interpretation, which could

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**Fig. 4** Bland-Altman plot of invasive tumour size measured by pathology subtracted from that measured by DBT (a) and MRI (b) compared with the mean of the two results. Middle dashed line is the mean difference and top and bottom dashed lines are the 95% limits of agreement (±2 standard deviations)
lower the diagnostic performance. It was not within the scope of this paper to compare the added value of DBT to DM alone.

In conclusion, DBT can be performed instead of MRI in women with non-dense breasts as determined by either a PD threshold or breast compositions a and b or in the case of MRI contraindications. MRI has higher diagnostic performance than DBT in a dense breast population in the tertiary setting.

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Compliance with ethical standards

Guarantor The scientific guarantor of this publication is Kaori Togashi, Kyoto University.

Conflict of interest The authors of this manuscript declare no relationships with any companies, whose products or services may be related to the subject matter of the article.

Statistics and biometry One of the authors has significant statistical expertise.

Informed consent Written informed consent was waived by the Institutional Review Board.

Ethical approval Institutional Review Board approval was obtained.

Methodology

• prospective
• diagnostic or prognostic study
• performed at one institution

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References

1. Poplack SP, Tosteson TD, Kogel CA et al (2007) Digital breast tomosynthesis: initial experience in 98 women with abnormal digital screening mammography. AJR Am J Roentgenol 189:616–623

2. Andersson I, Ikeda DM, Zackrisson S et al (2008) Breast tomosynthesis and digital mammography: a comparison of breast cancer visibility and BI-RADS classification in a population of cancers with subtle mammographic findings. Eur Radiol 18:2817–2825

3. Gur D, Abrams GS, Chough DM et al (2009) Digital breast tomosynthesis: observer performance study. AJR Am J Roentgenol 193:586–591

4. Rafferty EA, Park JM, Philpotts LE et al (2013) Assessing radiologist performance using combined digital mammography and breast tomosynthesis compared with digital mammography alone: results of a multicenter, multireader trial. Radiology 266:104–113

5. Friedewald SM, Rafferty EA, Rose SL et al (2014) Breast cancer screening using tomosynthesis in combination with digital mammography. JAMA 311:2499–2507

6. Gilbert FJ, Tucker L, Gillan MG et al (2015) Accuracy of digital breast tomosynthesis for depicting breast cancer subgroups in a UK retrospective reading study (TOMMY Trial). Radiology 277:697–706

7. Skaane P, Bandos AI, Gullien R et al (2013) Comparison of digital mammography alone and digital mammography plus tomosynthesis in a population-based screening program. Radiology 267:47–56

8. Ciatto S, Houssami N, Bernardi D et al (2013) Integration of 3D digital mammography with tomosynthesis for population breast-cancer screening (STORM): a prospective comparison study. Lancet Oncol 14(7):583–590

9. Lång K, Andersson I, Rosso A et al (2016) Performance of one-view breast tomosynthesis as a stand-alone breast cancer screening modality: results from the Malmö Breast Tomosynthesis Screening Trial, a population-based study. Eur Radiol 26:184–190

10. Boyd NF, Guo H, Martin LJ et al (2007) Mammographic density and the risk and detection of breast cancer. N Engl J Med 356:227–236

11. Eng A, Gallant Z, Shepherd J et al (2014) Digital mammographic density and breast cancer risk: a case-control study of six alternative density assessment methods. Breast Cancer Res 16:439

12. Nagata C, Matsubara T, Fujita H et al (2005) Mammographic density and the risk of breast cancer in Japanese women. Br J Cancer 92:2102–2106

13. Bae JM, Kim EH (2016) Breast density and risk of breast cancer in Asian women: a meta-analysis of observational studies. J Prev Med Public Health 49:367–375

14. Maskarinec G, Pagano I, Chen Z et al (2007) Ethnic and geographic differences in mammographic density and their association with breast cancer incidence. Breast Cancer Res Treat 104:47–56

15. Hori M, Matsuda T, Shibata A et al (2015) Cancer incidence and incidence rates in Japan in 2009; a study of 32 population-based cancer registries for the Monitoring of Cancer Incidence in Japan (MCIJ) project. Jpn J Clin Oncol 45:884–891

16. D’Orsi CJ, Mendelson EB, Ikeda DM (2003) ACR breast imaging and reporting data system: breast imaging atlas. American College of Radiology, Reston

17. D’Orsi CJ, Sickles EA, Mendelson EB et al (2013) ACR BI-RADS® Atlas, Breast Imaging Reporting and Data System. American College of Radiology, Reston

18. Gard CC, Aiello Bowles EJ, Miglioretti DL et al (2015) Misclassification of Breast Imaging Reporting and Data System (BI-RADS) mammographic density and implications for breast density reporting legislation. Breast J 21:481–489

19. He W, Juette A, Denton ER et al (2015) A review on automatic mammographic density and parenchymal segmentation. Int J Breast Cancer 2015:276217

20. McCormack VA, dos Santos Silva I (2006) Breast density and parenchymal patterns as markers of breast cancer risk: a meta-analysis. Cancer Epidemiol Biomarkers Prev 15:1159–1169

21. Are You Dense Advocacy, D.E.N.S.E. state efforts. http://AreYouDenseAdvocacy.org/dense/. Accessed 19 June 2017.

22. Melnikow J, Fenton JJ, Whitlock EP et al (2016) Supplemental screening for breast cancer in women with dense breasts: a systematic review for the U.S. Preventive Services Task Force. Ann Intern Med 164:268–278

23. Heywang-Köbrunner SH, Hacker A, Sledlack S (2013) Magnetic resonance imaging: the evolution of breast imaging. Breast 22(Suppl 2):S77–S82

24. Lima M, Le Bihan D (2016) Clinical intravoxel incoherent motion and diffusion MR imaging: past, present, and future. Radiology 278:13–32
25. ACR practice parameter for the performance of contrast-enhanced magnetic resonance imaging (MRI) of the breast. The American College of Radiology, Revised 2013 (Resolution 12). https://www.acr.org/~/media/20eb28eb59041e2825179af72ef624.pdf

26. Sardanelli F, Boetes C, Borisch B et al (2010) Magnetic resonance imaging of the breast: recommendations from the EUSOMA working group. Eur J Cancer 46:1296–1316

27. Keller BM, Nathan DL, Wang Y et al (2012) Estimation of breast percent density in raw and processed full field digital mammography images via adaptive fuzzy c-means clustering and support vector machine segmentation. Med Phys 39:4903–4917

28. Keller BM, Chen J, Daye D et al (2015) Preliminary evaluation of the publicly available Laboratory for Breast Radiodensity Assessment (LIBRA) software tool: comparison of fully automated area and volumetric density measures in a case-control study with digital mammography. Breast Cancer Res 17:117

29. Förnvik D, Zackrisson S, Ljungberg O et al (2010) Breast tomosynthesis: accuracy of tumor measurement compared with digital mammography and ultrasonography. Acta Radiol 51:240–247

30. Luparia A, Mariscotti G, Durando M et al (2013) Accuracy of tumour size assessment in the preoperative staging of breast cancer: comparison of digital mammography, tomosynthesis, ultrasound and MRI. Radiol Med 118:1119–1136

31. The International Commission on Radiation Units and Measurements (2008) ICRU Report 79: receiver operating characteristic analysis in medical imaging. J ICRU 8:1–62

32. Mercier J, Kwiatkowski F, Abrial C et al (2015) The role of tomosynthesis in breast cancer staging in 75 patients. Diagn Interv Imaging 96:27–35

33. Clauer P, Carbonaro LA, Pancot M et al (2015) Additional findings at preoperative breast MRI: the value of second-look digital breast tomosynthesis. Eur Radiol 25:2830–2839

34. Mariscotti G, Houssami N, Durando M et al (2015) Digital breast tomosynthesis (DBT) to characterize MRI-detected additional lesions unidentified at targeted ultrasound in newly diagnosed breast cancer patients. Eur Radiol 25:2673–2681

35. Kim WH, Chang JM, Moon HG et al (2016) Comparison of the diagnostic performance of digital breast tomosynthesis and magnetic resonance imaging added to digital mammography in women with known breast cancers. Eur Radiol 26:1556–1564

36. Rafferty EA, Durand MA, Conant EF et al (2016) Breast cancer screening using tomosynthesis and digital mammography in dense and nondense breasts. JAMA 315:1784–1786

37. Sprague BL, Gangnon RE, Burt V et al (2015) Prevalence of mammographically dense breasts in the United States. J Natl Cancer Inst 106(10). https://doi.org/10.1093/jnci/dju255

38. McCarthy AM, Keller BM, Pantalone LM et al (2016) Racial differences in quantitative measures of area and volumetric breast density. J Natl Cancer Inst 108(10). https://doi.org/10.1093/jnci/djw104

39. Hansen NL, Kuhl CK, Barabasch A et al (2014) Does MRI breast “density” (degree of background enhancement) correlate with mammographic breast density? J Magn Reson Imaging 40:483–489

40. King V, Brooks JD, Bernstein JL et al (2011) Background parenchymal enhancement at breast MR imaging and breast cancer risk. Radiology 260:50–60

41. Chudgar AV, Conant EF, Weinstein SP et al (2017) Assessment of disease extent on contrast-enhanced MRI in breast cancer detected at digital breast tomosynthesis versus digital mammography alone. Clin Radiol 72:573–579