Preoperative assessment of breast cancer: Multireader comparison of contrast-enhanced MRI versus the combination of unenhanced MRI and digital breast tomosynthesis

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A B S T R A C T
Purpose: To compare the sensitivity for breast cancer (BC) and BC size estimation of preoperative contrast-enhanced magnetic resonance imaging (CEMRI) versus combined unenhanced magnetic resonance imaging (UMRI) and digital breast tomosynthesis (DBT).

Patients and methods: We retrospectively included 56 women who underwent DBT and preoperative 1.5 T CEMRI between January 2016–February 2017. Three readers with 2–10 years of experience in CEMRI and DBT, blinded to pathology, independently reviewed CEMRI (diffusion-weighted imaging [DWI], T2-weighted imaging, pre- and post-contrast T1-weighted imaging) and a combination of UMRI (DWI and pre-contrast T1-weighted imaging) and DBT. We calculated per-lesion sensitivity of CEMRI and UMRI + DBT, and the agreement between CEMRI, UMRI and DBT versus pathology in assessing cancer size (Bland-Altman analysis). Logistic regression was performed to assess features predictive of cancer missing.

Results: We included 70 lesions (64% invasive BC, 36% ductal carcinoma in situ or invasive BC with in situ component). UMRI + DBT showed lower sensitivity (86–89%) than CEMRI (94–100%), with a significant difference for the most experienced reader only ($p = 0.008$). False-positives were fewer with UMRI + DBT (4–5) than with CEMRI (18–25), regardless of the reader ($p = 0.001–0.005$). For lesion size, UMRI showed closer limits of agreement with pathology than CEMRI or DBT. Cancer size $<1$ cm was the only independent predictor for cancer missing for both imaging strategies (Odds ratio 8.62 for CEMRI and 19.16 for UMRI + DBT).

Conclusions: UMRI + DBT showed comparable sensitivity and less false-positives than CEMRI in the preoperative assessment of BC. UMRI was the most accurate tool to assess cancer size.

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1. Introduction

Contrast-enhanced magnetic resonance imaging (CEMRI) has emerged as the most reliable tool for assessing loco-regional extent of breast cancer (BC). Based on superior contrast resolution and the capability of exploiting tumoral neoangiogenesis [1], CEMRI is highly sensitive in detecting multicentric, multifocal and bilateral cancer foci [2,3], as well as in predicting BC size at pathology [4–6]. Despite this, the impact on surgical outcomes of preoperative CEMRI is disputed [7,8]. On the other hand, this technique is increasingly used for tailoring the surgical approach [9].

There are several disadvantages limiting the widespread adoption of CEMRI. First, examination specificity is generally regarded as suboptimal for the purpose of minimizing overdiagnosis and overtreatment of false-positive (FP) findings, even though second-
We identify each technique, i.e. high contrast resolution for UMRI, and high retrospective design.

2.1. Study population and standard of reference

Patients and methods

In this light, the main purpose of this study was to compare CEMRI and UMRI + DBT in the preoperative assessment of BC in terms of cancer detection (index lesions and multicentric/multifocal cancer foci) and cancer size estimation.

2. Patients and methods

2.1. Study population and standard of reference

The referring Institutional Review Board approved this study, waiving for the acquisition of the informed consent because of the retrospective design.

By performing a search between January 2016—February 2017, we identified all the women referred to surgery for biopsy-proven BC who underwent preoperative CEMRI, as required by the policy of our tertiary referral institution. Of 514 subjects, we excluded those addressed to neoadjuvant therapy after MRI (n = 7) or showing post-biopsy changes such as clip-related susceptibility artefacts, seroma or haemorrhage (n = 128). Other 323 women were excluded because they did not undergo DBT as prior first-line imaging, or underwent it monolaterally. Final population included 56 women with bilateral CEMRI and DBT, presenting a total of 58 index lesions (i.e., lesions initially prompting CEMRI), and 12 additional lesions confirmed by subsequent ultrasound-guided biopsy and/or final pathological examination after surgery. Of 56 women, 2 showed bilateral index lesions. Among additional lesions, 9 were found in the same breast quadrant of the index lesion, while 3 were in a different quadrant. Women showed a mean age of 58.8 years (±12.3 standard deviation, range 33—81 years).

Pathological analysis was the standard of reference for imaging findings included in the surgical specimen. Analysis was performed by a pool of three pathologists with 5—25 years of experience in breast pathology, who processed surgical specimens and measured cancer according to the College of American Pathologists guidelines valid at the time in which surgery was performed (January 2016—February 2017) [28,29]. Imaging follow-up of at least 2 years served as surrogate standard of reference for those findings not included in the surgical specimen and assessed as benign (see below). Imaging included variable combinations of mammography, ultrasound, DBT and CEMRI.

2.2. Imaging protocols

DBT was acquired on one of two digital mammography systems: Giotto TOMO (Internazionale Medico Scientifica, Bologna, Italy); AW5000 3D Selenia Dimensions (Hologic, Bedford, MA, USA). Each breast was investigated with standard cranio-caudal and mediolateral oblique views, with both systems generating synthetic two-dimensional (2D) images (s2D) for each view. Table 1 in Appendix A shows the acquisition parameters of DBT.

CEMRI examinations were performed in the prone position on one of two 1.5 T magnets: Magnetom Avanto or Magnetom Aera (Siemens Medical Solutions, Erlangen, Germany), using a bilateral breast 4-channel or 16-channel coil, respectively. The CEMRI protocol, which included DWI, pre- and post-contrast T1-weighted dynamic contrast enhanced imaging (DCE), and T2-weighted imaging, is detailed in Table 2 of Appendix A. Using a remote control injector (OptiStar Elite injector, Mallinckrodt, UK), we administered gadobenate dimeglumine (Multihance, Bracco, Milan, Italy) intravenously, at a dose of 0.1 mmol/kg and an injection rate of 2 ml/s.

2.3. Image analysis

A study coordinator with 4 years of experience in breast imaging, not involved in image analysis, organized reading sessions on our picture archiving and communication system (PACS) workstation (SuitEstensa, Ebit AET, Esaote, Genoa, Italy). ADC maps were generated by the vendor’s software (Leonardo Syngo, Siemens Medical Systems, Erlangen, Germany). Cases were prompted during independent and separate reading sessions to three different readers, including a senior radiologist (R1), and two residents (R2 and R3). R1, R2 and R3 showed 10, 3 and 3 years of experience in CEMRI and DBT, respectively. Readers were aware of interpreting preoperative examinations, though they were blinded to prior imaging and clinical information (including lesions’ number and type). They accessed two types of examinations: CEMRI alone (including all the sequences reported in Table 2 of Appendix A) or UMRI + DBT. While DBT included native and s2D images for each view, UMRI was composed by the unenhanced T1-weighted sequence and DWI. DWI included the following sets of images: 1) images obtained with low diffusion-weighting (so called low b-
measurement was performed on the slice and image (T1-weighted, measurement on either CEMRI or DBT [31,32]. On UMRI, lesion better visibility in the case of DBT. Spiculae were excluded from in the case of CEMRI, and on the view in which the lesion showed formed in the subtracted series showing better lesion conspicuity two imaging modalities.

Lignant if assessed as category 4 or larger with at least one of the When interpreting UMRI readers referred to the categorization score illustrated in Fig. 1. Since the BI-RADS is inapplicable to UMRI, were de on CEMRI [30]. For the purpose of analysis, breast density and BPE were on DBT, and the degree of breast parenchymal enhancement (BPE) RADS categorization included also the assessment of breast density and size (cm) of imaging imaging sets.

Table 1

| Type of lesions                  | Histological type       | Number of lesions (% on the total of lesions) | Median size (cm) | IQR         |
|---------------------------------|-------------------------|-----------------------------------------------|------------------|-------------|
| Index cancers (n = 58)          | IDC<sup>a</sup>         | 30 (42.6%)                                     | 2.2              | 13.5–30.0   |
|                                 | IDC + DCIS<sup>b</sup>  | 18 (25.7%)                                     | 1.1              | 9.0–15.0    |
|                                 | IDLC                    | 4 (5.7%)                                       | 2.8              | 16.0–41.0   |
|                                 | ILC                     | 1 (1.4%)                                       | 1.3              | –           |
|                                 | ILC + LCIS              | 1 (1.4%)                                       | 2.1              | –           |
|                                 | DCIS<sup>c</sup>        | 4 (5.7%)                                       | 6.0              | 32.0–92.5   |
| Additional cancers (n = 12)     | IDC<sup>d</sup>         | 8 (11.4%)                                      | 0.9              | 8.5–12.0    |
| Multifocal cancers              | IDC<sup>e</sup>         | 1 (1.4%)                                       | 0.6              | –           |
| Multicentric cancers            | IDC<sup>e</sup>         | 2 (2.9%)                                       | 1.0              | –           |
|                                 | DCIS<sup>e</sup>        | 1 (1.4%)                                       | 0.4              | –           |

<sup>a</sup> DCIS: grade 1 (n = 1), grade 2 (n = 2), and grade 3 (n = 2).
<sup>b</sup> DCIS component associated to invasive cancer: grade 1 (n = 3), grade 2 (n = 7), and grade 3 (n = 6).
<sup>c</sup> IDC: grade 1 (n = 4), grade 2 (n = 20), grade 3 (n = 16).
<sup>d</sup> IDC with associated in situ component: grade 1 (n = 4), grade 2 (n = 13), grade 3 (n = 2).

Table 2

Per-reader distribution of the sensitivity, true positive (TP), false negative (FN), and false positive (FP) cases achieved with contrast-enhanced magnetic resonance imaging (CEMRI) versus the combination of unenhanced magnetic resonance imaging (UMRI) and digital breast tomosynthesis (DBT). Denominators of the ratios provided in the rows represent the total number of TP + FN + FP cases.

| Reader 1 | Reader 2 | Reader 3 |
|----------|----------|----------|
|          | CEMRI    | UMRI + DBT | p<sup>a</sup> | CEMRI    | UMRI + DBT | p<sup>a</sup> | CEMRI    | UMRI + DBT | p<sup>a</sup> |
| Sensitivity % (95%CI) | 100 (0.95–1.00) | 88.6 (0.79–0.94) | 0.008 | 94.3 (0.87–0.98) | 87.1 (0.77–0.93) | 0.180 | 94.3 (0.87–0.98) | 85.7 (0.75–0.92) | 0.180 |
| TP       | 70/92    | 62/75     | 0.342 | 66/95    | 61/75     | 0.109 | 66/88    | 60/74     | 0.448 |
| FN       | 0/92     | 8/75      | 0.001 | 4/95     | 9/75      | 0.008 | 4/88     | 10/74     | 0.052 |
| FP       | 22/92    | 5/75      | 0.002 | 25/95    | 5/75      | 0.001 | 18/88    | 4/74      | 0.005 |

<sup>a</sup> Significance of the intra-reader comparison between CEMRI versus UMRI + DBT.

value images), which are indispensable to build the ADC map; 2) images with strong diffusion-weighting (so called high b-value images), which show the highest tumor-to-normal tissue contrast; 3) the ADC map, obtained from low b-value and high b-value images, representing the spatial distribution of the ADC as a measure of water diffusion in the tissue (the term “apparent” accounts for several tissue effects making this index different from pure diffusion coefficient [24]). CEMRI or UMRI + DBT dataset were presented randomly by separating the readings performed on the same women by an interval of time of ≥3 weeks between the two imaging sets.

For each patient, R1-R3 recorded the site (side and quadrant) and size (cm) of imaging findings, categorizing them with the Breast Imaging Reporting and Data System (BI-RADS) for CEMRI and DBT [24]. Findings were assessed as malignant if scored ≥4. BI-RADS categorization included also the assessment of breast density on DBT, and the degree of breast parenchymal enhancement (BPE) on CEMRI [30]. For the purpose of analysis, breast density and BPE were defined as “high” when categorized ≥C and moderate-to-marked, respectively. Since the BI-RADS is not applicable to UMRI, readers referred to the categorization score illustrated in Fig. 1. When interpreting UMRI + DBT, a finding was considered as malignant if assessed as category 4 or larger with at least one of the two imaging modalities.

Cancer size was measured on the slice showing the largest diameter when reading CEMRI or DBT. Measurement was performed in the subtracted series showing better lesion conspicuity in the case of CEMRI, and on the view in which the lesion showed better visibility in the case of DBT. Spiculae were excluded from measurement on either CEMRI or DBT [31,32]. On UMRI, lesion measurement was performed on the slice and image (T1-weighted, high b-value or ADC map) where the observation showed maximum conspicuity and diameter.

2.4. Statistical analysis

After assessing data normality with the Shapiro-Wilk test, we reported continuous variables as median and interquartile range (IQR) values. Proportions were reported together with 95% confidence intervals (95%CI).

We calculated the sensitivity for each reader, defined as the percent ratio between true positive (TP) cases and the sum of TP and false negative (FN) cases. On an intra-reader basis, we assessed whether there was any significant difference between CEMRI and UMRI + DBT in terms of sensitivity (using the McNemar test), as well as in the number of TP or FP or FN cases over the total number of TP + FP + FN cases (using the Fisher exact test).

Concerning cancer size, we evaluated CEMRI, DBT and UMRI separately and on an intra-reader basis. Using the Friedman test, we assessed whether there was any significant difference in estimating the size of TP findings that there were detected by all imaging modalities. On the same set of findings, we used Bland-Altman analysis to exploit the agreement of each imaging modality with pathological size. Data underwent logarithmic transformation to match the prerequisites of analysis [33,34], so that mean bias and 95% limits of agreement between measurements were presented as dimensionless values. Bland-Altman analysis was integrated with the intraclass correlation coefficient (ICC), using the following reference values for the agreement [35]: ≤0.40 = poor; 0.40–0.59 = fair; 0.60–0.74 = good; 0.75–1.00 = excellent.

Finally, we built two stepwise logistic regression models for determining which clinical or imaging variable was predictive of.
missing cancer by at least one of the three readers when using CEMRI or UMRI + DBT.

Analysis was performed on a commercially available software (MedCalc Software bvba, version 18.11.6, Ostend, Belgium). Alfa level was set 0.05, using the Bonferroni correction in the case of multiple pairwise comparisons.

3. Results

3.1. Cancers characteristics

Included women (n = 56) underwent unilateral mastectomy in 33 cases, bilateral mastectomy in 1 case, unilateral breast-conserving surgery in 21 cases, and bilateral breast-conserving surgery in 1 case, respectively.

Table 1 shows the histological type and size of 70 breast cancers found at pathological examination. Median size of all lesions, index lesions alone, and additional lesions alone was 1.5 cm (IQR 0.9–3.0 cm), 1.8 cm (IQR 1.1–3.0 cm) and 0.7 cm (IQR 0.5–1.1 cm), respectively. Overall, cancers size was ≤1.0 cm in 22/70 lesions (31.4%), corresponding to 14/58 index lesions, and 8/12 additional lesions.

3.2. Sensitivity for breast cancer

As shown in Table 2, CEMRI showed higher sensitivity for BC than UMRI + DBT (range 94.3–100% versus 85.7–88.6%). The difference was statistically significant (p = 0.008) in the case of R1 only. FNs induced by UMRI + DBT were significantly (p = 0.001–0.008) or nearly significantly (p = 0.052) larger in number than those induced by CEMRI for R1-R2 and R3, respectively.

Higher sensitivity of CEMRI was counterbalanced by a significantly larger number of FPs compared to UMRI + DBT, regardless of the reader (p ranging 0.001–0.005). There was no significant difference in the number of TPs achieved by both imaging modalities for each reader. The distribution and characteristics of FN and FP findings are detailed in Table 3 Appendix A. FP cases were confirmed by core needle biopsy (CNB), pathological examination if included in the surgical specimen, or imaging follow-up, as detailed in Table 3 in Appendix A. All the FP cases included in the surgical specimen were operated at the time of primary surgery, except for one B3 lesion found by core needle biopsy (identified by R2 only in the study), which underwent lumpectomy separately (no atypia at pathological examination).

Most of CEMRI-induced FPs were avoided by UMRI + DBT, i.e. 21/22 (95.4%) for R1, 24/25 (96.0%) for R2, and 17/18 (94.4%) for R3. Exemplificative cases are shown in Figs. 2–3.

Overall, CEMRI and UMRI + DBT agreed in categorizing cancers with a score ≥4 in 62/70 cases for R1 (88.6%; 95% CI 78.2–94.6), 59/70 cases for R2 (84.3%; 95% CI 73.2–91.5), and 56/70 for R3 (80.0%; 95% CI 68.4–88.3). Agreement in categorizing a finding as suspicious was not investigated with Cohen’s kappa statistic because of the low number of <3 assignments attributed in common by the two imaging strategies (0 for R1 and R3, and 2 for R2).

The number of benign findings identified by R1, R2 and R3 was 20 in 18 women, 28 in 15 women, and 23 in 14 women using CEMRI, and 19 in 17 women, 20 in 18 women and 22 in 16 women using UMRI + DBT, respectively. None of those findings was shown to be malignant over time.

3.3. Cancer size

Median (IQR) size of category ≥4 findings on CEMRI, DBT and UMRI was 1.5 cm (1.0–3.2), 2.1 cm (1.0–3.0) and 2.0–05 cm (1.3–3.0) for R1; 1.7 cm (1.1–3.0), 2.0 cm (1.1–3.0) and 2.0 cm (1.3–3.0) for R2; 1.7 cm (1.1–3.0), 1.8 cm (1.1–2.8) and 2.0 cm (1.0–3.0) for R3, respectively.

Lesions detected by all imaging modalities were 51/70 for R1, 53/70 for R2, and 43/70 for R3. Among them, there was no significant difference in size between CEMRI, DBT and UMRI (p = 0.328–0.826) as interpreted by R1 (median/IQR 2.3/1.3–3.6 cm, 2.2/1.6–3.0 cm, and 2.1/1.4–3.1 cm, respectively) and R2 (median/IQR 2.2/1.3–3.1 cm, 2.0/1.1–3.0 cm and 2.1/1.3–3.3 cm, respectively).
respectively). In the case of R3, CEMRI provided significantly larger size (median/IQR 2.2/1.3–3.1 cm) than DBT (median/IQR 1.9/1.3–2.9 cm) and UMRI (median/IQR 2.0/1.4–3.1 cm).

The results from agreement analysis (Fig. 4) showed that, regardless of the reader, UMRI alone showed narrower limits of agreement and higher ICC values compared, in the descending order, to CEMRI and DBT alone. ICCs showed a similar trend (Fig. 4),

Table 3
Main characteristics of false negative (FN) and false-positive (FP) cancers achieved by contrast-enhanced magnetic resonance imaging (CEMRI) and unenhanced magnetic resonance imaging plus digital breast tomosynthesis (UMRI + DBT). IQR = interquartile range; IDC = invasive ductal carcinoma; DCIS = ductal carcinoma in situ; IDLC = mixed invasive ductal-lobular carcinoma; G1 = grade 1; G2 = grade 2; G3 = grade 3.

| Reader | Number | Histological type | Median size (IQR) (cm) | FN cancers | FP cancers | FN cancers | FP cancers | FN cancers | FP cancers |
|--------|--------|-------------------|-----------------------|------------|------------|------------|------------|------------|------------|
| 1      |        |                   |                       | 0          | 22         | 8          | 5          | 0          | 1          |
|        |        | IDC G2, 1 IDC G3, 1 IDC G1 + DCIS G2, 1 IDC G2 + DCIS G2 | 1.5 (0.7) | 0.7 (0.5–0.7) | 0.8 (0.8) | -2.3       | -          | -          | -          |
| 2      | 4      | IDC G2, 1 IDC G3, 1 IDC G2 + DCIS G1, 1 IDLC G2 | 0.8 (0.6–0.9) | 1.1 (0.9) | 0.7 (0.5–0.8) | 1.0 (0.9) | -1.9       | -          | -          |
|        |        |                   |                       | 18         | 25         | 9          | 4          | 0          | 1          |
| 3      |        | IDC G2, 1 IDC G3, 1 DCIS G3 | 2.5 (0.8–5.1) | 0.9 (0.7) | 0.7 (0.6–1.4) | 1.2 (0.9) | -1.8       | -          | -          |
with R1–R3 values in the range of excellent agreement for UMRI (0.82–0.87) versus good agreement for CEMRI (0.71–0.77), and fair-to-good agreement for DBT (0.45–0.65). Mean bias at Bland–Altman analysis was minimal, regardless of the imaging method.

3.4. Cancer missing

The number and main characteristic of cancers missed by each reader are reported in Table 3. Cancers missed by at least one reader were 8/70 for CEMRI, and 12/70 for UMRI + DBT, respectively.

All readers categorized breast density as BI-RADS B, C and D in 8/56, 34/56, and 14/56 patients, respectively. Readers agreed also in assessing BPE as minimal in 32/56 women, mild in 10/56 women, moderate in 11/56 women, and marked in 3/56 women, respectively.

At multivariate analysis (Table 4), smaller lesions size (<1.0 cm) was the only independent predictor of missing cancer for both CEMRI and UMRI + DBT, though odds ratio (OR) values showed wide 95% CIs.

4. Discussion

In our study, the sensitivity of UMRI + DBT in the preoperative assessment of BC was high (range 85.7–88.6%), even if inferior to that of CEMRI (range 94.3–100%). The difference was statistically significant in the case of the most experienced readers (R1) only (p = 0.008), possibly because less experienced readers missed more cancers even when using the standard imaging of CEMRI, thus reducing the gap in sensitivity as compared to UMRI + DBT. However, CEMRI provided a nearly significant-to-significant (p = 0.001–0.052) lower number of FN cases, regardless of the reader. On the bright side, all readers achieved significantly (p = 0.001–0.005) less FP findings than CEMRI when using UMRI + DBT. Our results suggest that UMRI + DBT can be a reliable diagnostic alternative to CEMRI in the preoperative assessment of BC, while carrying some specific advantages of UMRI. Indeed, UMRI is a mean to reduce the acquisition time and costs related to contrast medium administration [36]. Its use might potentially increase the accessibility to preoperative breast MRI, and in turn expand current indications [37]. Moreover, avoiding contrast administration would eliminate safety concerns, as exemplified at best in those patients in which gadolinium chelates should be used with caution because of impaired renal function, risk for adverse reaction, or risk for accumulation in the central nervous system [14,15].

Results on sensitivity were expected, given the well-known superiority of CEMRI in detecting loci of tumoral neoangiogenesis [1–3]. Not surprisingly, they are in line with previous studies comparing UMRI alone versus CEMRI, either using different unenhanced protocols (range of sensitivity 50.0–93.8% versus 86.0–98.0%, respectively) [17,18,20,22,23], or investigating a similar combination of a T1-weighted sequence and DWI (range of sensitivity 78.4–91.6%) [16,23]. However, we found that the absolute concordance rate between the two imaging strategies was comparably high across the readers (range 80.0–88.6%), suggesting a substantial and reproducible agreement in categorizing imaging findings, regardless of the radiologist’s experience.

We did not calculate specificity, given the ambiguity of this index in a context with 100% cancer prevalence. However, our results on FP findings are reasonably in contrast with previous results on specificity. While DBT was shown to be more specific than CEMRI (88.2% versus 74.2%) [38], UMRI alone showed overlapping or even lower specificity in studies of direct comparison with CEMRI (range 71.4–89.3% versus 76.8–92.8%). A possible explanation for the discrepancy is that, differently from those studies, readers were aware of interpreting preoperative examinations, and in turn of the presence of BC. This scenario was affected by the unavoidable detection bias related to the higher likelihood that an imaging finding was malignant [39]. In this scenario, which reflects clinical practice, UMRI + DBT avoided most FPs induced by CEMRI, at a comparable extent in all readers. It should be pointed that the median size of CEMRI-induced FPs was small (range 0.9–1.5 cm for R1–R3). This finding matches with the results of logistic regression, showing that smaller BC size was the only independent predictor of cancer missing, at a larger extent for UMRI + DBT than CEMRI (OR 19.16 versus 8.62, respectively). Our results suggest that UMRI + DBT has the potential of avoiding small FPs induced by CEMRI, and in turn reducing additional “second-look” imaging or biopsy needed for ruling out malignancy [10–12]. This would further limit costs, patient anxiety, and the overall time from diagnosis to surgery, though the role for UMRI + DBT in this setting should be confirmed in further studies.

Finally, our study prevalently included invasive ductal carcinomas, which is usually available as a pre-biopsy examination, assuming that main role for UMRI was cancer detection, given high contrast resolution inherent to the technique (especially DWI), while DBT better exploited lesions’ morphology. Of note, higher breast density did not impair cancer detection according to multivariate analysis, suggesting that lesions potentially missed by DBT can be complementary diagnosed by UMRI, whose image contrast is inherently unaffected by the amount of fibroglandular tissue. Finally, our study prevalently included invasive ductal carcinomas,
alone or in combination with non-invasive components, suggesting that the generalizability of our results should be proven in series encompassing a large number of different histological types, especially invasive lobular carcinoma, as well as difficult-to-diagnose lesions with benign-like appearance (e.g., mucinous tumors).

5. Conclusions

Combined UMRI + DBT showed high sensitivity when performing preoperative assessment of BC. Compared to CEMRI, sensitivity was expectedly lower, though UMRI + DBT avoided the large majority of the FPs induced by the former technique. UMRI showed the highest agreement with pathology in terms of cancer size, even with regard to CEMRI, thus appearing as the reference tool to measure BC in the UMRI + DBT approach. BC size $\leq 1.0$ cm was the only independent predictor of missing cancer for both imaging strategies. Based on our results, one might propose UMRI + DBT as an alternative to CEMRI having the potential of increasing the accessibility to preoperative breast MRI, facilitating patients’ workup, and reducing contrast-related risks.

Ethical approval

The study was approved by the Institutional Review Board. The acquisition of written informed consent from patients was waived because of the retrospective design.

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Declaration of competing interest

Francesco Sardanelli declares to have received grants from or being member of the speakers’ bureau/advisory board for Bayer-Schering, Bracco, and General Electric. All the other authors have no conflict of interest to declare.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.breast.2019.11.013.

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| Lesions missed by CEMRI | Lesions missed by UMRI + DBT |
|------------------------|-----------------------------|
| **Univariate analysis** | **Multivariate analysis** |
| Age                    |                             |
| $<$49 years 2/16 (12.5%) | $<$49 years 4/16 (25%)      |
| 50–69 years 5/32 (15.6%) | 50–69 years 5/32 (15.6%)    |
| $>$70 years 1/22 (4.5%)  | $>$70 years 3/22 (13.6%)    |
| $p = 0.386$             | $p = 0.625$                 |
| High breast density ($\geq$BI-RADS C) 8/48 (16.7%) $p = 0.043$ | 9/48 (18.7%) $p = 0.600$ |
| High BPE (moderate-to-marked) 2/19 (10.5%) $p = 0.885$ | Not included Non included |
| Invasive histology 7/55 (10.7%) $p = 0.534$ | 10/65 (15.4%) $p = 0.162$ |
| Size $\leq 1.0$ cm 6/22 (27.3%) $p = 0.005$ | 10/22 (45.5%) $p = 0.0001$ |
| $p = 0.012$ or 8.62 (95%CI 1.58–47.13) | $p = 0.0004$ or 19.16 (95%CI 3.69–99.38) |
| Being an additional lesion 1/12 (8.3%) $p = 0.713$ | 5/12 (41.6%) $p = 0.013$ |

* Chi-square test.
especially the dentate nucleus. Investig Radiol 2016;51:273–9. https://doi.org/10.1097/IR.0000000000000273.

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