Breast dynamic contrast enhanced MRI: fibrocystic changes presenting as a non-mass enhancement mimicking malignancy

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Background. We aimed to analyse the morphokinetic features of breast fibrocystic changes (nonproliferative lesions, proliferative lesions without atypia and proliferative lesions with atypia) presenting as a non-mass enhancement (NME) in dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) examination.

Patients and methods. Forty-six patients with histologically proven fibrocystic changes (FCCs) were retrospectively reviewed, according to Breast Imaging Reporting and Data System (BI-RADS) lexicon. Prior to DCE-MRI examination, a unilateral breast lesion suspicious of malignancy was detected clinically, on mammography or breast ultrasonography.

Results. The predominant features of FCCs presenting as NME in DCE-MRI examination were: unilateral regional or diffuse distribution (in 35 patients or 76.1%), heterogeneous or clumped internal pattern of enhancement (in 36 patients or 78.3%), plateau time-intensity curve (in 25 patients or 54.3%), moderate or fast wash-in (in 31 patients or 67.4%). Nonproliferative lesions were found in 11 patients (24%), proliferative lesions without atypia in 29 patients (63%) and lesions with atypia in six patients (13%), without statistically significant difference of morphokinetic features, except of the association of clustered microcysts with proliferative dysplasia without atypia.

Conclusions. FCCs presenting as NME in DCE-MRI examination have several morphokinetic features suspicious of malignancy, therefore requiring biopsy (BI-RADS 4). Nonproliferative lesions, proliferative lesions without atypia and proliferative lesions with atypia predominantly share the same predefined DCE-MRI morphokinetic features.

Key words: breast; magnetic resonance imaging; fibrocystic changes

Introduction

Fibrocystic changes (FCCs) are the most frequent benign conditions of breast, diagnosed in 50% of women examined clinically and in 90% of women in histopathological studies. These benign disorders have two important implications from the point of view of breast cancer diagnosis and management. First, FCCs can mimic breast cancer on clinical examination, mammography and breast ultrasonography, leading to unnecessary breast biopsies and patient anxiety. Second, some types of FCCs represent a risk factor for the subsequent development of breast cancer. Based on a classification system of FCCs proposed by Dupont and Page and according to other studies, women with
Histologically confirmed nonproliferative lesions have no increased breast cancer risk. On the contrary, women whose breast biopsies show proliferative lesions with or without atypia are at risk of developing cancer, with relative risk ranging from 3.9–13.0 and 1.3–1.9 respectively.2,4

Combining morphological and enhancement kinetics features of the breast lesions, dynamic contrast-enhanced MRI (DCE-MRI) shows the highest sensitivity of all imaging methods in detecting breast diseases, up to 100%. Nevertheless, the specificity in the differentiation between benign and malignant lesions is lower, up to 75%.1 The major cause of false positive findings in DCE-MRI examination and consecutive unnecessary biopsies are the lesions with non-mass enhancement (NME). NME refers to the lesion that is seen only on post-contrast DCE-MRI sequences and does not have space-occupying effect. The enhancement pattern of NME is distinct from normal surrounding breast parenchyma and may contain interspersed fat. On the contrary, a mass enhancement is a three-dimensional space-occupying lesion.6,7 The causes of NME include FCCs, inflammatory benign lesions, in situ ductal carcinoma (DCIS), invasive lobular carcinoma and some cases of oestrogen receptor-negative invasive ductal carcinoma. The mass enhancement is usually confined to malignant or benign tumors.8,11

Our study of pathologically confirmed FCCs presenting as NME in DCE-MRI examination has two goals: 1) to analyse morphological and enhancement kinetics features of FCCs, 2) to compare these features between nonproliferative lesions, proliferative lesions without atypia and proliferative lesions with atypia.

Patients and methods

Patients

In the period of two years (January 2010 to January 2012) a total of 947 patients were examined by two radiologists (MZC and NMM) using the standardized breast DCE-MRI full diagnostic protocol. From this group, 46 patients with FCCs presenting as NME were selected and retrospectively reviewed. The study was performed in accordance with the Declaration of Helsinki and was approved by the Institutional Review Board Committee (No. 4502-01/2011). All patients gave written informed consent to participate in DCE-MRI examination. The age of the patients was 50.78 ± 8.99 years. In all premenopausal women DCE-MRI was performed in the second and third week of the menstrual cycles. None of the patients had previous breast biopsy, breast surgery or current hormone replacement therapy. All patients initially had unilateral breast lesion suspicious of malignancy either on clinical examination, mammography or ultrasonography. The lesions were presented as calcifications or asymmetric tissue on mammography and hypoechoic non-mass lesions on ultrasonography (BI-RADS 4a to BI-RADS 4c categories according to the ACR BI-RADS lexicon).6,12 The indication for breast DCE-MRI prior to biopsy was to evaluate the local extent of the lesion in the dense breast on mammography (32 patients) or suspected multifocal lesions on mammography and/or ultrasonography (14 patients).13 After the pathological confirmation of FCCs, the patients were examined biannually by physical examination and mammography. During the predefined follow-up period, no ipsilateral or contralateral breast cancer was detected.

Methods

The DCE-MRI examinations were performed with a 1.5 Tesla MRI unit (Magnetom Avanto, Siemens Medical Solutions, Erlangen, Germany) with dedicated bilateral breast coil and the patient in the prone position. The standard protocol was used for the axial-plane images with the slice thickness of 2 mm (Table 1).14 The contrast medium was gadopentetate dimeglumine (Magnevist, Bayer Schering Pharma, Berlin, Germany) applied as the bolus injection of 0.1 mmol/kg body weight injected with the automatic injector (Mississippi, Ulrich Medical, TABLE 1. Standard dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) protocol for axial-plane images for T1-weighted FLASH 3D precontrast and five postcontrast series

| Parameter                  | T1-weighted | T2-weighted | T2-weighted | T1-weighted | T1-weighted |
|----------------------------|-------------|-------------|-------------|-------------|-------------|
| MRI sequence/parameters    | TIRM        | TSE         | TSE         | FLASH 3D    |
| Echo time [ms]             | 60          | 70          | 12          | 4.8         |
| Repetition time [ms]       | 7690        | 5900        | 910         | 9.1         |
| Inversion time [ms]        | 180         |             |             |             |
| Flip angle (°)             | 150         | 180         | 90          | 25          |
| Field of view (mm×mm)      | 340×340     | 340×340     | 340×340     | 340×340     |
| Image matrix               | 320×256     | 384×319     | 320×234     | 576×564     |

3D = three-dimensional; DCE-MRI = dynamic contrast-enhanced magnetic resonance imaging; FLASH = fast low-angle shot pulse sequence; TIRM = turbo inversion recovery magnitude; TSE = turbo spin-echo.
Ulm, Germany) at the rate of 2 mL/s, followed by the flush of 20 mL saline. Contrast-enhanced dynamic sequences were acquired five times every 1 min 23 s. The postprocessing methods, including a creation of time intensity curves (TICs), i.e. enhancement (%) against time (s), were performed on the workstation Leonardo, using the image processing software Syngo (Syngo, Siemens Medical Solutions).

Two experienced radiologists in breast MRI, ZCM, 20 yr. and MMN, 7 yr., reviewed the DCE-MRI examinations. Based on the fifth edition of the BI-RADS lexicon, published in 2013, the following morphokinetic features of NME were analysed: distribution of enhancement (focal, linear, segmental, regional, multiple regions, diffuse), internal enhancement pattern (homogeneous, heterogeneous, clumped, clustered ring enhancement, stippled), type of TICs (persistent curve or type 1, plateau curve or type 2, wash out curve or type 3), wash-in i.e. the enhancement rate 90 s after intravenous contrast application (slow, medium, rapid), and signal intensity on T2W images.

A surgical biopsy was performed in all patients. Among them, 21 patients with nonpalpable lesions, prior to the biopsy underwent a radiogu- ided occult lesion localization (ROLL) procedure with Technetium-99m, under stereotactic or ultrasonographic guidance. Haematoxylin and eosin stained slides of formalin-fixed and paraffin-embedded tissue blocks were assessed by a pathologist, experienced in breast pathology. According to the classification system of Dupont and Page, the lesions were classified as nonproliferative lesions, proliferative lesions without atypia, and proliferative lesions with atypia (atypical ductal and lobular hyperplasia).

**Statistical analysis**

Following DCE-MRI features of NME were analysed: distribution of postcontrast enhancement, internal enhancement pattern, type of TICs, wash-in, and signal intensity on T2W images. These distinctive DCE-MRI features of NME were compared for three groups of FCCs: nonproliferative lesions, proliferative lesions without atypia, and proliferative lesions with atypia. Frequencies were used to describe the distribution of categorical variables. The difference between investigated variables in three groups of FCCs was analysed using χ² test. P values less than 0.05 were considered statistically significant. SPSS for Windows, Statistics version 16.0. (SPSS Inc., Chicago, USA) was used to perform statistical analyses.

**Results**

The lesions were detected clinically in 25 patients (54.3%), on mammography in 43 patients (93.5%) and by breast ultrasonography in 30 patients (65.2%).

Out of 46 biopsies, nonproliferative lesions were found in 11 patients (24%), proliferative lesions without atypia in 29 patients (63%) and proliferative lesions with atypia in six patients (13%). On DCE-MRI examination, all cases were presented as unilateral non-mass enhancement of BI-RADS 4 category (Figure 1).

Forty-three lesions (93.5%) were equal to or larger than 1 cm in size and three lesions were smaller than 1 cm and larger than 0.5 cm in size (6.5%). The distribution of the postcontrast enhancement was segmental in 11 patients (23.9%) and regional or diffuse in 35 patients (76.1%). The other types of distribution (focal, linear enhancement and multiple regions) were not found. The internal enhancement pattern was homogeneous in two patients (4.3%), heterogeneous and clumped, clustered ring enhancement, stippled. Clustered ring enhancement was not detected. The types of time-intensity curves were persistent (type 1) in 17 patients (37.0%), plateau (type 2) in 25 patients (54.3%), and wash out (type 3) in four patients (8.7%). The initial postcontrast signal in-
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Intensity enhancement (enhancement rate, wash-in) was slow in 15 patients (32.6%), moderate in 14 patients (30.4%), and fast in 17 patients (37.0%). On T2W images FCCs were associated with clustered microcysts in 25 cases (54.3%).

As shown in Table 2, the features of nonproliferative lesions, proliferative lesions without atypia and proliferative lesions with atypia on DCE-MRI examination did not show statistically significant difference in the term of the size (p = 0.454), the distribution of postcontrast enhancement (p = 0.168), the internal enhancement pattern (p = 0.722), the types of time-intensity curves (p = 0.097) and the initial postcontrast signal intensity enhancement (p = 0.752). Presence of microcysts on T2W images in these three groups of FCCs was statistically significant compared to the lack of the feature (p = 0.014).

**Discussion**

Numerous studies have demonstrated the clinical importance of FCCs, related to the high prevalence of the condition, the considerable impact on quality of life and the increased breast cancer risk for women with proliferative lesions.2-4,16-18 Nevertheless, only few studies analysed features of FCCs on DCE-MRI, based on the limited number of cases. Chen et al. reported two studies with 31 patients and 11 patients, analysing FCCs in DCE-MRI examination and MR spectroscopy for choline detection.19,20 Van den Bosch et al. and Kiyak et al. reported morphological and kinetic features of FCCs on DCE-MRI in the group of 14 patients and 27 patients, respectively.21,22

Our study included 46 symptomatic patients with pathologically confirmed FCCs presenting as NME, mimicking malignancy in DCE-MRI examination. Chen et al. reported that FCCs on DCE-MRI had characteristics of NME in 39% of cases, mass enhancement in 35% of cases, while 26% of cases were nonenhancing lesions.19 According to the published data, breast DCE-MRI has the potential to distinguish benign from malignant mass lesions effectively. Nevertheless, DCE-MRI is inferior in discriminating benign from malignant NME lesions.23 Hence, a meticulous analysis of the multiple, standardized parameters from BI-RADS lexicon is crucial to achieve a higher diagnostic performance of DCE-MRI in the case of FCCs presenting as NME, including the distribution of enhancement, the internal enhancement pattern, the

| Statistical difference | Proliferative lesions without atypia (N = 6) | Proliferative lesions with atypia (N = 29) | Non proliferative lesions (N = 11) | The DCE-MRI features of fibrocystic changes |
|------------------------|---------------------------------------------|------------------------------------------|-----------------------------------|------------------------------------------|
| p = 0.454              | 1                                           | 1                                        | 1                                 | 0.5 < d < 1 cm                           |
|                        | 5                                           | 28                                       | 10                                | d ≥ 1 cm                                 |
| p = 0.168              | 3                                           | 7                                        | 1                                 | Segmental NME distribution               |
|                        | 3                                           | 22                                       | 10                                | Regional or diffuse                       |
| p = 0.722              | 0                                           | 2                                        | 0                                 | Homogeneous NME internal enhancement     |
|                        | 1                                           | 4                                        | 3                                 | Stippled                                  |
|                        | 5                                           | 23                                       | 8                                 | Heterogeneous or clumped                 |
| p = 0.097              | 3                                           | 8                                        | 6                                 | Persistent TIC                           |
|                        | 3                                           | 19                                       | 3                                 | Plateau                                   |
|                        | 0                                           | 2                                        | 2                                 | Wash out                                 |
| p = 0.752              | 2                                           | 8                                        | 5                                 | Slow Wash-in                             |
|                        | 1                                           | 10                                       | 3                                 | Moderate                                 |
|                        | 3                                           | 11                                       | 3                                 | Fast                                     |
| p = 0.014              | 1                                           | 18                                       | 2                                 | Present Microcysts (T2W images)          |
|                        | 5                                           | 11                                       | 9                                 | Absent                                   |

\(d = \) longest diameter; DCE-MRI = dynamic contrast-enhanced magnetic resonance imaging; NME = non-mass enhancement; TIC = time-intensity curve; T2W = T2-weighted
type of TICs, the wash-in, and the signal intensity on T2W images.

In our study the postcontrast enhancement of FCCs was unilateral in all patients. The most frequent types of the distribution of NME were regional - involving more than 25% of a breast quadrant, and diffuse - involving the entire breast (35 patients or 76.1%). Segmental distribution, reflecting the ductal distribution, was detected in 11 patients (23.9%). Thomassin-Naggara et al. emphasize that „the more extensive distribution of enhancement is, the less suspicious for malignancy it is”.8 Agrawal et al. showed that the diffuse enhancement was highly suggestive of benign lesions.24 On the other hand, segmental NME was considered the most suspicious, with the positive predictive value (PPV) for malignancy from 76% to 100%, confirmed with the study by Tozaki and Fukuda.25 Our results show that FCCs mainly show the more extensive distribution - regional and diffuse, suggesting benign aetiology of the condition.

The internal pattern of enhancement in our study was heterogeneous and clumped in 36 cases (78.3%). The stippled enhancement was noted in eight patients (17.4%) and homogeneous in two patients (4.3%). According to the previous studies (Thomassin-Naggara et al., Tozaki and Fukuda), when the enhancement is heterogeneous or clumped the risk of malignancy is considered high, with the PPV of 53–58%, while the homogeneous enhancement has low PPV for malignancy, up to 5%. The stippled enhancement represents the normal breast parenchymal enhancement.8,25 Since almost 80% of the cases in our study showed heterogeneous or clumped enhancement on DCE-MRI, the internal pattern can be one of the most important reasons for mimicry of the breast cancer by FCCs.

Kinetic curve enhancement reflects functional aspects of blood vessel permeability in normal and pathological breast tissue after intravenous gadolinium contrast application. According to the published data, the permanent enhancement (type 1 time-intensity curve) is seen in benign lesions in 85% of cases, the plateau (type 2) curve is seen in 36% of malignant cases and the washout (type 3) curve in 57% of malignant cases.26 In our study the majority of patients have type 2 curve (25 or 54.3%), followed by type 1 curve in 17 patients (37%) and type 3 in four patients (8.7%). Generally, the published data about the kinetic curve enhancement of NME significantly differ. In the study of FCCs by Chen et al. 90% of NME lesions had the type 1 curve.29 Bartella et al. reported predominantly the type 2 curve in cases of malignant NME.27 Goto et al. did not find any differences in the kinetic curve enhancement between benign and malignant NME lesions.28 These differences may be partly due to the fact that the time-intensity curves are based on the semiquantitative analysis of gadolinium contrast uptake with the free-hand selected region of interest (ROI) on the heterogeneous areas of NME. Some recent studies suggested that the quantitative analysis using automated computer-aided diagnosis (CAD) can overcome subjectivity of the free-hand selection of ROI.29 The postcontrast enhancement rate types in our study were slow in 15 patients (32.6%), moderate in 14 patients (30.4%) and fast in 17 patients (37.0%). Based on the published data, slow wash-in indicates benign lesions, medium wash-in may indicate lesions like mastitis, fresh scar, FCCs or DCIS, while the fast initial enhancement is suggestive of malignancy, especially when combined with the type 2 or the type 3 enhancement curve.30

NME lesions in our study of FCCs were associated with microcysts on T2W images in 54.3% of cases, which was suggestive of benign conditions.8

Additionally, we investigated DCE-MRI morphokinetic features of three distinct histological types of FCCs: nonproliferative lesions (11 cases or 24%), proliferative lesions without atypia (29 cases or 63%) and proliferative lesions with atypia (6 cases or 13%). Our results show that nonproliferative lesions, proliferative lesions without atypia and proliferative lesions with atypia have similar morphokinetic features (distribution of NME, internal pattern of enhancement, type of TICs and wash-in) with the exception of the significant association of microcysts with proliferative hyperplasia without atypia. To our knowledge, our study is the first, which analyses the morphokinetic features of distinct histological types of FCCs on MRI examination. Our study was prompted by the report from Hartman et al. published in 2015, which anticipated a potentially important role of DCE-MRI in case of atypical hyperplasia.31 Hartman et al. showed that in women with atypical hyperplasia a lifetime risk of breast cancer approaches 30% at 25 years. This high cumulative incidence is not widely recognized, and thus screening DCE-MRI is not routinely recommended for these patients.13,32 Hartman et al. suggested more intensive screening of women with atypical hyperplasia, with DCE-MRI added to mammography, as well as the use of selective oestrogen-receptor modulators and aromatase inhibitors to prevent breast cancer in women with atypical hyperplasia. We hypothesized that the differentiation of atypical hyperplasia by DCE-MRI
examination from other types of FCCs could be helpful in DCE-MRI screening. Our study was limited by the low number of proliferative lesions with atypia, thus necessitating further investigation.

Our study has some limitations. We did not analyse incidental findings of FCCs on DCE-MRI. Our analysis was conducted in the selected group of symptomatic patients: the lesions were suspicious of malignancy on clinical examination, mammography or ultrasonography and larger than 1 cm in DCE-MRI examination in 93.5% of cases. These facts may have influenced the interpretation and results of DCE-MRI examination.

In conclusion, the profile of FCCs presented as NME in DCE-MRI examination predominantly includes: unilateral regional or diffuse distribution (76.1% of cases), heterogeneous or clumped internal pattern of enhancement (78.3% of cases), plateau (type 2) time-intensity curve (54.3% of cases) with moderate or fast wash-in (67.4% of cases), and associated clustered microcysts (54.3% of cases). Although these findings do not have the classic appearance of malignancy, they are sufficiently suspicious to recommend the biopsy, as final BI-RADS 4 category. In case of NME, proliferative lesions without atypia and proliferative lesions with atypia (13%), DCE-MRI cannot show the subtle histological differences between nonproliferative lesions, proliferative lesions without atypia and proliferative lesions with atypia. According to some novel data, MR diffusion-weighted imaging (DWI), related to tissue cellularity and thermal motion of water molecules instead of permeability of blood vessels after the contrast uptake, may be more specific than DCE-MRI to define the benign nature of FCCs.33,34

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