A multiparametric $[{^{18}}F]$FDG PET/MRI diagnostic model including imaging biomarkers of the tumor and contralateral healthy breast tissue aids breast cancer diagnosis

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Abstract
Purpose To develop a multiparametric $[{^{18}}F]$FDG positron emission tomography/magnetic resonance imaging (PET/MRI) model for breast cancer diagnosis incorporating imaging biomarkers of breast tumors and contralateral healthy breast tissue.

Methods In this prospective study and retrospective data analysis, 141 patients (mean 57 years) with an imaging abnormality detected on mammography and/or ultrasound (BI-RADS 4/5) underwent combined multiparametric $[{^{18}}F]$FDG PET/MRI with PET/computed tomography and multiparametric MRI of the breast at 3 T. Images were evaluated and the following were recorded: for the tumor, BI-RADS descriptors on dynamic contrast-enhanced (DCE)-MRI, mean apparent diffusion coefficient (ADCmean) on diffusion-weighted imaging (DWI), and maximum standard uptake value (SUVmax) on $[{^{18}}F]$FDG-PET; and for the contralateral healthy breast, background parenchymal enhancement (BPE) and amount of fibroglandular tissue (FGT) on DCE-MRI, ADCmean on DWI, and SUVmax.

Histopathology served as standard of reference. Uni-, bi-, and multivariate logistic regression analyses were performed to assess the relationships between malignancy and imaging features. Predictive discrimination of benign and malignant breast lesions was examined using area under the receiver operating characteristic curve (AUC).

Results There were 100 malignant and 41 benign lesions (size: median 1.9, range 0.5–10 cm). The multivariate regression model incorporating significant univariate predictors identified tumor enhancement kinetics ($P=0.0003$), tumor ADCmean ($P<0.001$), and BPE of the contralateral healthy breast ($P=0.0019$) as independent predictors for breast cancer diagnosis. Other biomarkers did not reach significance. Combination of the three significant biomarkers achieved an AUC value of 0.98 for breast cancer diagnosis.

Conclusion A multiparametric $[{^{18}}F]$FDG PET/MRI diagnostic model incorporating both qualitative and quantitative parameters of the tumor and the healthy contralateral tissue aids breast cancer diagnosis.

Keywords Breast cancer · Positron emission tomography · Fluorodeoxyglucose · Magnetic resonance imaging · Imaging biomarker
Introduction

In recent years, positron emission tomography (PET) and multiparametric magnetic resonance imaging (mpMRI) of the breast have been established as promising tools [1–3] that provide morphologic and functional data and are of complementary value for the differentiation of benign and malignant breast tumors [4]. MpMRI consisting of dynamic contrast-enhanced MRI (DCE-MRI) [5, 6] and diffusion-weighted imaging (DWI) [7, 8] provides information on tumor neoangiogenesis, microenvironment and microstructural breast tumors [4]. PET using the radiotracer $^{18}$F-fluorodeoxyglucose ($^{18}$F-FDG) provides data on lesion glucose metabolism and is of complimentary value to MRI for breast cancer diagnosis [2, 9–13]. To combine the strengths of these imaging techniques, hybrid imaging systems have been developed and introduced into the clinical routine. With PET/MRI systems, morphologic and different functional information can be merged to non-invasively characterize breast tumors as benign or malignant.

The predictive and prognostic value of imaging features derived from healthy breast tissue such as background parenchymal enhancement (BPE) in DCE-MRI have been recognized only recently. BPE is the physiological contrast enhancement of fibroglandular tissue (FGT). Previous studies have found correlations between higher levels of BPE and FGT and increased cancer risk in a high risk population [14, 15], while associations between apparent diffusion coefficient (ADC) values of normal breast tissue and cancer outcomes have not been explored. Similar to MRI, healthy breast parenchyma shows varying degrees of tracer uptake in $^{18}$F-FDG PET, reflecting its physiologic activity, referred to as breast parenchymal uptake (BPU) [16, 17] and is directly correlated with BPE and FGT. It has therefore been suggested that BPU could potentially serve as another breast cancer imaging biomarker [16–18].

With the clinical implementation of PET/MRI scanners and given the improvements in diagnosis and prognosis that can be achieved with mpMRI, it is of interest to explore the potential of combined $^{18}$F-FDG PET/MRI to improve diagnostic accuracy and obviate unnecessary breast biopsies of benign breast tumors detected with MRI. To date it is unclear which quantitative and qualitative imaging features from both tumor and healthy tissue contribute to an accurate diagnosis. To close this gap in knowledge we aimed to develop a multiparametric $^{18}$F-FDG PET/MRI model for breast cancer diagnosis incorporating imaging biomarkers of breast tumors and contralateral healthy breast tissue.

Material and methods

The local institutional review board (Ethikkommission Medizinische Universitaet Wien/Ethics Committee Medical University of Vienna) approved this prospective single-institution study (EK 510/2009) and retrospective data analysis. The research was performed in accordance with relevant guidelines/regulations and informed consent was obtained from all patients prior to $^{18}$F-FDG PET/MRI.

Patient population

Patients were examined with $^{18}$F-FDG PET/computed tomography (CT) and mpMRI in a prospective trial. For the current study that focused on model development, the prospectively populated research database was searched for patients who underwent multiparametric $^{18}$F-FDG PET/MRI with PET/CT and mpMRI of the breast at 3 T between December 2009 and September 2014 and who met the following inclusion criteria: 18 years or older; not pregnant; not breastfeeding; no previous breast cancer treatment; imaging abnormality on mammography or breast ultrasound (BI-RADS 4, suspicious abnormality; or 5, highly suggestive for malignancy); and no contraindications for MRI or MRI contrast agents. Exclusion criteria were high-risk patients (confirmed BRCA 1/2 mutation, lifetime risk >20%, incomplete examinations, severe motion or susceptibility artifacts, no histopathologic confirmation by image-guided or surgical biopsy after MRI and $^{18}$F-FDG PET/CT, and presence of another suspicious lesion in the contralateral breast. Thus, 141 patients with a unilateral breast lesion in mammography, ultrasound, MRI, and $^{18}$F-FDG PET/MRI (BI-RADS 1) were included for analysis. Electronic medical records were reviewed to record patient age and histopathology results, including tumor grade, subtype, and receptor status for malignant lesions. A number of patients included in this study have been analyzed and reported before in different contexts [4, 11, 16].

Imaging

All patients underwent combined multiparametric $^{18}$F-FDG PET/MRI with PET/CT using the tracer $^{18}$F-FDG and multiparametric MRI of the breast at 3 T. Examinations were scheduled no longer than 6 days apart (mean, 1.15; range, 0–6; same day, $n = 70$; 1 day, $n = 31$; 2 days, $n = 12$; 3 days, $n = 11$; 4 days, $n = 11$; 5 days, $n = 5$; 6 days, $n = 1$).

$^{18}$F-FDG pet/CT

A combined PET/CT in-line system (Biograph 64 TruePoint PET/CT system, Siemens, Erlangen, Germany) was used to acquire PET images. Patients fasted 6 h before they were injected with approximately 300 MBq $^{18}$F-FDG, depending on the patient’s weight, with an uptake time of 45 min. Blood glucose levels were below 150 mg/dl (8.3 mmol/l) at the time of tracer application. A prone PET dataset of the breasts was acquired and an unenhanced low-dose CT was used for
attenuation correction. PET images were reconstructed using the TrueX algorithm (Siemens, Erlangen, Germany), which includes commonly used correction factors, as well as a specific point-spread function correction. Four iterations per 21 subsets were used, with a matrix size of 168 × 168, trans-axial field of view of 605 mm (pixel size, 3.6 mm), and section thickness of 5 mm.

**Multiparametric MRI**

MRI was performed in the prone position using a 3 T MRI scanner (Tim Trio, Siemens, Erlangen, Germany) and a four-channel breast coil (InVivo, Orlando, FL, USA). In premenopausal patients, MRI was performed between the 7th and 14th day of the menstrual cycle to minimize hormonal influence on BPE [19]. All patients underwent a state-of-the-art MRI protocol as follows.

- **T2-weighted turbo spin echo (TSE)** imaging with fat suppression: repetition time/echo time (TR/TE) 4800/9 ms; field of view (FOV) 340 mm; 48 slices at 3 mm; flip angle 128°; matrix 384 × 512; time of acquisition (TA) 2 min, 16 s.
- **Axial three-acquisition trace diffusion-weighted**, double-refocused single-shot echo-planar imaging (EPI) with inversion recovery fat suppression: TR/TE/time of inversion (TI) 8000/59/210 ms; FOV 360 × 202 mm; 24 slices at 5 mm; intersection gap (%) 10; matrix 172 × 96 [50% oversampling]; b-values 50, 850 s/mm; TA 2 min, 16 s.

DCE-MRI using a split-dynamics protocol until 12/2011 [21] consisting of five alternating sections of high-spatial and high-temporal resolution T1-weighted sequences: (1) high spatial resolution unenhanced coronal T1-weighted fast-low-angle-shot-(FLASH)-3D sequences without preparation pulse, with selective water-excitation (TR/TE 877/3.82 ms; FOV 320 mm; 96 slices; 1 mm isotropic; matrix 320 × 134; one average; bandwidth 200 Hz/pixel; TA 2 min); (2) high temporal resolution coronal T1-weighted Volume Interpolated Breath-hold Examination (VIBE) (TR/TE 3.61/1.4 ms; FOV 320 mm; 72 slices; 1.7 mm isotropic; matrix 192 × 192; one average; bandwidth 400 Hz/pixel; 13.2 s per volume); (3) repeated high spatial resolution unenhanced coronal T1-weighted 3D-FLASH at peak contrast enhancement; (4) repeated high temporal resolution VIBE (25 measurements), repeated 3D-FLASH, TA 5 min, 35 s); and (5) high spatial resolution T1-weighted images 3D-FLASH for delayed phase. Total TA was 9 min 20 s.

DCE-MRI protocol from 01/2012 onwards: transversal T1-weighted time-resolved angiography with stochastic trajectories (TWIST); water excitation fat-saturation; TR/TE 6.23/2.95 ms; flip angle 15°, FOV 196 × 330 mm²; 144 slices; spatial resolution 0.9 × 0.9 × 1 mm; temporal interpolation factor 2; temporal resolution 14 s; matrix 384 × 384; one average; center k-space region, resampling rate of 23%; reacquisition density of peripheral k-space 20%; TA 6 min, 49 s.

DCE-MRI image series were acquired before and after the injection of a standard dose (0.1 mmol/kg body-weight) of Gadotareneglumine (Gd-DOTA; Dotarem®, Guerbet, France) as a bolus using a power injector (Spectris Solaris EP®, Medrad, Pittsburgh, PA, USA).

MpMRI and PET data were fused semi-automatically using the “landmark matching” tool of the TrueD fusion workstation (Siemens, Erlangen, Germany) to generate combined [18F]FDG PET/MRI data.

**Data analysis**

**[18F]FDG pet**

All measurements were performed by a radiologist (D.L., 3 years of experience) trained in hybrid imaging under the supervision of a nuclear medicine physician. A three-dimensional volume of interest (VOI) was placed around the primary breast tumor to record lesion maximum standardized uptake value (SUVmax) using the TrueD fusion workstation (Siemens, Erlangen, Germany). The VOI was established using the Region Grow function with a fixed threshold as determined by the reader to represent PET metabolic tumor volume but excluding physiologic [18F]FDG uptake in surrounding tissues. For quantification of BPU of the normal contralateral breast, a VOI was placed around the glandular tissue. Adequate distance to the nipple and areola was kept. All VOI measurements were repeated three times and averaged to ensure data consistency.

**DCE-MRI**

DCE-MRI data was evaluated by two radiologists in consensus (K.P., 12 years of experience, and D.L., 3 years of experience). Both readers were aware that the patient had a lesion but were not provided with previous imaging or histopathology results. All images were assessed using the revised ACR MRI BI-RADS lexicon [22]. Lesions were classified as mass or non-mass-enhancement (NME), and described accordingly using BI-RADS descriptors. For the evaluation of enhancement kinetics, automated semi-quantitative curve-type analysis was performed using dedicated software (DCE tool plugin v2.2 for OSIRIX software) [23]. The largest diameters measured on DCE-MRI were recorded. For the contralateral healthy breast, BPE and FGT were assessed qualitatively, as recommended by the revised ACR MRI BI-RADS lexicon [22]. BPE was evaluated using DCE-MRI images 90 s after contrast agent administration and classified as minimal, mild, moderate, or marked. FGT was evaluated using unenhanced T1-weighted fat-suppressed images and graded as ACR (a) for almost entirely fatty breasts, (b) for scattered fibroglandular tissue, (c) for heterogeneous fibroglandular tissue, and (d) for...
extreme amount of fibroglandular tissue. Both BPE and FGT were assessed in the healthy contralateral breast only.

**DWI**

DCE-MR images were used to identify the lesion on DWI with ADC mapping. The slice with the greatest diameter of the tumor was selected, and an ROI (minimum 10 mm²) was drawn on the part of the tumor with the lowest ADC by two radiologists in consensus (K.P., and D.L.). ADCmean was recorded. For quantification of ADCmean of the healthy contralateral breast, a 10 mm² standardized 2D ROI was manually drawn in normal appearing fibroglandular tissue. For each lesion and healthy breast, ROI measurements were repeated three times and results were averaged.

All tumor characteristics on [18F]FDG PET, DCE-MRI and DWI are displayed in Table 1, while parameters derived from healthy breast tissue can be found in Table 2.

**Histopathology**

Histopathology served as standard of reference. Histopathological diagnosis was established using tissue derived from image-guided needle biopsy or surgical specimens. In case of a high-risk lesion with uncertain potential for malignancy, the final diagnosis was established with surgery.

In 33 patients with benign lesions, final histology was established via core-needle biopsy, while three patients opted for elective surgery. Five high-risk lesions on breast biopsy remained benign on surgery. In 48 patients with malignant lesions, final histology was established from surgical specimen, while in 52 patients that received neoadjuvant chemotherapy, histology was obtained from core-needle biopsy specimen.

**Statistical analysis**

Statistical tests were performed using SAS, v9.4 (SAS Institute, Cary, NC, United States). All calculations were performed on a per-lesion basis. The main outcome was presence of malignancy. Using histopathology as the standard of reference, disease status was dichotomized as positive or negative for malignancy. Dichotomized variables further included mass shape (round/oval/no enhancement vs. irregular), mass margins (circumscribed vs. irregular/spiculated), mass internal enhancement characteristics (homogeneous/dark internal septation vs. heterogeneous/rim enhancement), NME distribution (region/segmental vs. multiple regions/diffuse), and NME pattern of enhancement (homogeneous vs. heterogeneous/clumped). Enhancement kinetics of the tumor (persistent, plateau, wash-out) were examined as a three-level categorical variable, while BPE (minimal, mild, moderate, marked) and FGT (ACR a–d) were assessed as four-level categorical variables. Patient age, lesion size, tumor and parenchyma ADCmean, and FDG SUVmax were kept in continuous form.

Univariate analysis, bivariate analysis and multivariate logistic regression were applied to assess the relationships between malignancy and imaging features. We analyzed the frequency distribution of demographic, clinical and imaging characteristics using the chi-square test or Fisher’s exact test for categorical variables and quantile-quantile plots (Q–Q plots), non-parametric Wilcoxon test and T-test for continuous variables, where appropriate.

Variables with an association in bivariate analysis were examined. We assessed the association between histology and the following imaging features: BPE of the healthy breast, internal enhancement characteristics, mass margin and shape, enhancement kinetics, tumor ADCmean, tumor SUVmax, and tumor size. A priori confounder, age, was also examined. Confounders were included in the final model if they altered

**Table 1** Lesion imaging characteristics assessed on [18F]FDG PET/MRI

| Imaging modality | BI-RADS descriptors/imaging characteristics | Categories/explanation                                                                 |
|------------------|--------------------------------------------|----------------------------------------------------------------------------------------|
| DCE-MRI          | Mass                                       | Round, oval, irregular                                                                  |
|                  | Shape                                      | Circumscribed, irregular, spiculated                                                   |
|                  | Margin                                     | Homogeneous, heterogenous, rim enhancement, dark internal septations                    |
|                  | Internal enhancement characteristics        | Focal, linear, segmental, regional, multiple regions, diffuse                           |
|                  | Distribution                               | Homogeneous, heterogeneous, clumped, clustered ring                                    |
|                  | Internal enhancement pattern                | Symmetric, asymmetric                                                                   |
|                  | Symmetry                                   | Slow, medium, fast                                                                     |
|                  | Kinetic curve                              | Persistent, plateau, washout                                                           |
|                  | Initial phase                              | Mean ADC value with ROI drawn over the part of the lesion with the visually lowest ADC  |
|                  | Delayed phase                              | SUVmax value with ROI drawn over the whole tumor                                        |

NME non mass enhancement, ADC apparent diffusion coefficient, SUVmax maximum standardized uptake value
Results

There were 100 malignant and 41 benign lesions in 141 patients (140 female; mean age, 57 ± 14.3 years, range 18–86 years). Among these lesions, there were 121 masses (93 malignant, 28 benign) and 20 NME lesions (13 malignant, 7 benign) as defined according to the MRI BI-RADS lexicon. The average size of malignancies (27.3 mm) was larger than the size of benign lesions (22.6 mm) (P = 0.0227).

On DCE-MRI, morphological and kinetic features predictive of breast cancer presenting as masses were irregular/spiculated margin, irregular shape, heterogeneous internal enhancement and plateau/wash-out curves (P < 0.0001). Likewise, frequent features for benign mass lesions were round/oval shape, circumscribed margins, homogeneous/dark septations internal enhancement characteristics and persistent enhancement curves (P < 0.0001). Patients with a malignancy showed decreased BPE of the contralateral healthy breast (P = 0.001), while a borderline significant difference was observed for FGT (P = 0.0564).

On DWI, malignancies showed significant lower average ADCmean (0.96 × 10⁻³ mm²/s) compared with benign lesions (1.52 × 10⁻³ mm²/s) (P < 0.0001).

On [¹⁸F]FDG PET, breast cancers showed a higher FDG avidity (mean SUVmax 5.2) than benign tumors (mean SUVmax 2.6) (P < 0.0001).

Stratified analysis showed that menopause is not a confounder to our results.

Results of univariate analysis are summarized in Table 3.

Multiparametric PET-MRI model

The previously published ADCmean cut-off of ≤1.25 × 10⁻³ mm²/s was used to dichotomize ADC values as positive or negative to differentiate benign from malignant tumors [30]. Multivariate logistic regression analysis determined that tumor ADCmean in DWI, tumor enhancement kinetics, and BPE of the contralateral healthy breast in DCE-MRI were significantly associated with breast cancer diagnosis.

Following these findings, our multiparametric [¹⁸F]FDG PET/MRI model incorporated these three significant univariate predictors. Based on this combination, the model discriminated between benign (Fig. 1) and malignant tumors (Fig. 2) with an area under the curve (AUC) value of 0.98 (Fig. 3).

The odds ratio (OR) of a lesion with ADCmean ≤1.25 × 10⁻³ mm²/s being a breast cancer is 1 (OR: 0.993, 95% confidence interval (CI): 0.99–0.996, P < 0.001). Lesions with a plateau or washout type kinetic curve had 18.7 greater odds of being malignant (OR: 18.7, 95%CI: 3.86–90.69) compared with those with persistent type enhancement (P = 0.0003). Lesions with decreased BPE of the contralateral healthy breast had 0.1 greater odds of being malignant (OR: 0.14, 95%CI: 0.04–0.49, P = 0.0019).

Discussion

With the increasing use of PET/MRI worldwide, it is of interest to clarify whether qualitative and quantitative MRI and PET imaging biomarkers can contribute to accurate breast cancer diagnosis. We analyzed the contribution of tumor [¹⁸F]FDG SUVmax, BPU of healthy breast tissue, tumor DCE-MRI BI-RADS descriptors, BPE, and FGT of the contralateral healthy breast and ADCmean of the tumor and healthy breast tissue. The results of this study show that a multiparametric [¹⁸F]FDG PET/MRI diagnostic model incorporating both qualitative and quantitative parameters of the tumor and the healthy contralateral tissue aids breast cancer diagnosis. In particular, tumor enhancement kinetics, tumor ADCmean, and BPE of the contralateral healthy breast were most strongly associated with breast cancer. While our findings indicate that imaging biomarkers of healthy breast parenchyma can contribute to breast cancer diagnosis, [¹⁸F]FDG PET parameters did not add to accuracy.

To date, in breast imaging, regression models based on MRI have already been shown to clarify which parameters...
are necessary for breast cancer detection and staging. Knuttel et al. developed a predictive model incorporating DCE-MRI parameters including FGT for prediction of the ductal carcinoma in situ (DCIS) component associated with the primary breast cancer.

Table 3 Results of univariate analysis: qualitative and quantitative variables for predicting malignancy

|                          | Total          | Malignant (n = 100) | Benign (n = 41) | P value |
|--------------------------|----------------|---------------------|-----------------|---------|
| Patient age              |                |                     |                 |         |
| Mean (SD)                | 59.85 (12.01)  | 49.85 (17.00)       |                 | .0007   |
| Median (min-max)         | 61.33 (36.88–86.11) | 50.29 (0.0–86.11)  |                 | .0227   |
| Tumor size               |                |                     |                 |         |
| Mean (SD)                | 27.26 (20.56)  | 22.56 (19.178)      |                 |         |
| Median (min-max)         | 19.5 (6.0–100.0) | 13.0 (5–80)        |                 |         |
| Mass margin              |                |                     |                 |         |
| Circumscribed            | 3 (2.5)        | 23 (19.2)           |                 | <.0001  |
| Irregular/spiculated     | 90 (75.0)      | 3 (2.5)             |                 |         |
| Mass shape               |                |                     |                 | <.0001  |
| Round/oval/no enhancement| 15 (12.4)      | 23 (19.0)           |                 |         |
| Irregular                | 78 (64.5)      | 5 (4.1)             |                 |         |
| Mass internal enhancement characteristics |                |                     |                 | <.0001  |
| Homogeneous/dark         | 15 (12.5)      | 21 (17.5)           |                 |         |
| Heterogeneous/rim        | 78 (65.0)      | 6 (5.0)             |                 |         |
| NME distribution         |                |                     |                 |         |
| Regional/segmental       | 1 (5.0)        | 3 (15.0)            |                 |         |
| Multiple/diffuse         | 6 (30.0)       | 10 (50.0)           |                 |         |
| NME enhancement pattern  |                |                     |                 | 0.2487  |
| Homogeneous              | 0 (0)          | 4 (20.0)            |                 |         |
| Heterogeneous/clumped    | 7 (65.0)       | 9 (45.0)            |                 |         |
| Enhancement kinetics     |                |                     |                 | <.0001  |
| Persistent               | 4 (2.9)        | 21 (15.2)           |                 |         |
| Plateau                  | 44 (31.9)      | 14 (10.1)           |                 |         |
| Wash-out                 | 52 (72.5)      | 3 (2.2)             |                 |         |
| BPE healthy breast       |                |                     |                 | .001    |
| Minimal                  | 53 (37.6)      | 8 (5.7)             |                 |         |
| Mild                     | 36 (25.5)      | 20 (14.2)           |                 |         |
| Moderate                 | 9 (6.4)        | 10 (7.1)            |                 |         |
| Marked                   | 2 (1.4)        | 3 (2.1)             |                 |         |
| FGT healthy breast       |                |                     |                 | 0.0564  |
| ACR A                    | 30 (21.4)      | 5 (3.6)             |                 |         |
| ACR B                    | 43 (30.7)      | 17 (12.1)           |                 |         |
| ACR C                    | 19 (13.6)      | 10 (7.1)            |                 |         |
| ACR D                    | 30 (5.7)       | 8 (3.6)             |                 |         |
| ADCmean tumor            |                |                     |                 | <.0001  |
| Mean (SD)                | 1515.17 (325.05) | 961.57 (228.61)    |                 |         |
| Median (min-max)         | 1475.0 (883–2520) | 945.5 (521–1616)  |                 |         |
| ADCmean healthy breast   |                |                     |                 | 0.1779  |
| Mean (SD)                | 1698.5 (265.09) | 1770.37 (265.29)   |                 |         |
| Median (min-max)         | 1684.0 (1121–2398) | 1806 (1329–2369) |                 |         |
| SUVmax tumor             |                |                     |                 | <.0001  |
| Mean (SD)                | 5.19 (4.41)    | 2.56 (1.45)         |                 |         |
| Median (min-max)         | 3.97 (1.1–25.62) | 2.27 (0.4–6.9)     |                 |         |
| BPU healthy breast       |                |                     |                 | 0.0816  |
| Mean (SD)                | 1.7597 (0.59)  | 1.9483 (0.646)      |                 |         |
| Median (min-max)         | 1.59 (0.88–4.55) | 1.92 (0.88–3.96) |                 |         |

BPE background parenchymal enhancement, FGT amount of fibroglandular tissue, ADC apparent diffusion coefficient, SUVmax maximum standardized uptake value, BPU breast parenchymal uptake.
invasive breast cancer [24]. DWI with ADC mapping was also demonstrated to be a useful addition to prediction models presented by Rahbar et al. for the differentiation between high grade and non-high grade DCIS [25], and by Cheeney et al. for the prediction of upgrade to malignancy at surgical excision [26]. However, to date, neither a multiparametric model approach has been applied to PET/MRI of the breast nor have imaging features of the tumor as well as the healthy contralateral breast tissue been incorporated for cancer diagnosis. In this respect this study is particularly relevant.

Prior studies showed a positive correlation between BPE, FGT with MRI and BPU with [18F]FDG PET and therefore suggested that BPU could potentially aid as another easily quantifiable imaging biomarker for breast cancer [16, 17]. In our study, both features derived from [18F]FDG PET, SUVmax of the tumor and BPU SUVmax of the contralateral healthy breast, did not significantly contribute to breast cancer diagnosis. At least with respect to tumor SUVmax, this is not entirely surprising as in prior studies the additional information provided by [18F]FDG PET was not as great as originally anticipated. The radiotracer [18F]FDG has a good sensitivity but limited specificity and there is an overlap with several types of benign breast diseases. The use of other PET tracers that specifically target cancer hallmark processes such as hypoxia ([18F]-fluoromisonidazole) [27], proliferation ([18F]-fluorocholine, [18F]-fluorodeoxythymidine) [28] and tumor and microenvironment acidity (pHLIP) [29, 30] may add more diagnostic, predictive and prognostic information [31] and therefore it can be expected that the full potential of PET/MRI is not realized yet.

**Fig. 1** A 55-year-old postmenopausal woman with a fibroadenoma in the left breast presenting as a round, circumscribed mass with homogeneous enhancement on DCE-MRI. The lesion had no hindered diffusivity on DWI with an ADCmean of 1.625 $10^{-3}$ mm$^2$/s and was mildly [18F]FDG avid on PET with an SUVmax of 3.7. Unenhanced T1-weighted imaging shows heterogeneous fibroglandular tissue (ACR C) of the healthy breast, with moderate BPE on DCE-MRI, ADCmean of the healthy contralateral breast parenchyma is 1.642 $10^{-3}$ mm$^2$/s. On [18F]FDG PET, BPU of normal breast parenchyma is relatively high with an SUVmax of 3. A unenhanced fat-saturated T1-weighted MRI; B DCE-MRI; C DWI; D [18F]FDG PET; E fused [18F]FDG PET/MRI; F [18F]FDG PET/CT.
Previous studies have suggested that larger tumor size, irregular or spiculated shape and margin, heterogeneous internal enhancement, and wash-out kinetics on DCE-MRI as described using the BI-RADS lexicon and descriptors to be most strongly indicative of malignancy [32–34], which is in good agreement with our results. According to several studies and the revised BI-RADS lexicon, the combination of morphological and functional information acquired with DCE-MRI is necessary for breast cancer diagnosis [35]. Likewise, in the present study, multivariate logistic regression analysis identified tumor enhancement kinetics to be most strongly indicative of breast cancer.

Imaging parameters of healthy breast parenchyma are currently being examined as potential non-invasive predictive and prognostic breast cancer biomarkers [14, 15, 36]. Recently, BPE has been proposed as a risk factor for breast cancer in high-risk patients who were compared with matched control groups with benign or false-positive findings [14, 15]. These initial results were confirmed by Grimm et al., who compared another 61 high risk cancer patients with matched...
The multiparametric PET/MRI model developed in this study identified tumor ADCmean as one of three parameters essential for breast cancer diagnosis. Our results confirm the growing body of evidence that DWI is a valuable complimentary technique to DCE-MRI, adding specificity as well as functional information [5, 20, 40–45], and hence should constitute an essential part in breast imaging. Tumor ADCmean has even been found to be potentially useful as a non-invasive biomarker for tumor grade, subtype, receptor status, and recurrence [46, 47]. A recent study found ADC values of 248 benign and malignant lesions to be independent from BPE, FGT, and menopausal status [48]. This study verifies the usefulness of lesion ADC as a stable, reliable imaging biomarker in breast imaging despite different patient characteristics. Meanwhile, apart from their positive correlation with mammographic breast density and independence of menstrual cycle and BPE [49–51], not much is known about ADC values of healthy breast parenchyma. ADC values of the healthy breast did not add accuracy to the model developed in this study.

This study has some limitations. First, BPE and FGT were assessed qualitatively. However, the BI-RADS lexicon recommends qualitative assessment of these parameters, and excellent inter- and intra-reader agreement between experienced readers has been demonstrated previously for DCE-MRI and DWI parameters including BPE and FGT [4, 16, 52, 53]. Second, our study was not performed using a hybrid PET/MRI scanner, but fused PET/CT and MRI instead. Potential benefits of using a hybrid scanner would be improved fusion of PET and MRI images, as well as acquisition of both imaging series on the same day. Hence, in our study, not all MRI and PET/CT examinations were performed on the same day (range 0–6), which might have influenced parameters such as BPE and BPU due to changes with menstrual cycle [18, 54]. Nevertheless, relevant changes in these biomarkers seem unlikely, as the mean time of 1.15 days between examinations was short. Third, in this study, the same data set was used to train and evaluate regression models. Typically, such models perform better on the data they were trained with than on additional data. Further studies with larger patient collectives, separate training and validation datasets, and bilateral quantitative assessment of all parameters are warranted to confirm our findings. In addition, the model developed in this study needs to be validated in an independent test group.

In conclusion, in a multiparametric [18F]FDG PET/MRI model including imaging biomarkers of the tumor and healthy breast tissue, tumor enhancement kinetics, ADCmean tumor, and BPE of the contralateral breast were most strongly associated with breast cancer diagnosis. Other imaging biomarkers did not significantly contribute to breast cancer diagnosis. Implementation of multiparametric PET/MRI imaging biomarkers of healthy breast tissue for breast cancer diagnosis should be considered.

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Compliance with ethical standards

Conflict of Interest M.S.J. has received speaker honoraria from GE Healthcare. All other authors declare that they have no conflicts of interest.

Research involving human participants and/or animals Ethical approval: All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. This article does not contain any studies with animals performed by any of the authors.

Informed consent Informed consent was obtained from all individual participants included in the study.

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References

1. Morrow M, Waters J, Morris E. MRI for breast cancer screening, diagnosis, and treatment. Lancet. 2011;378(9805):1804–11.
2. Moy L, Ponzo F, Noz ME, Maguire GQ Jr, Murphy-Walcott AD, Deans AE, et al. Improving specificity of breast MRI using prone PET and fused MRI and PET 3D volume datasets. J Nucl Med. 2007;48(4):528–37.
3. Avril N, Adler LP. F-18 fluoro-deoxyglucose-potison emission tomography for primary breast cancer and loco-regional staging. Radiol Clin N Am. 2007;45(4):645–57 vi.
4. Magometschnigg HF, Baltzer PA, Fueger B, Helbich TH, Karanikas G, Dubsky P, et al. Diagnostic accuracy of (18)F-FDG PET/CT compared with that of contrast-enhanced MRI of the breast at 3 T. Eur J Nucl Med Mol Imaging. 2015;42(11):1656–65.
5. Zhang L, Tang M, Min Z, Lu J, Lei X, Zhang X. Accuracy of combined dynamic contrast-enhanced magnetic resonance imaging and diffusion-weighted imaging for breast cancer detection: a meta-analysis. Acta Radiol. 2016;57(6):651–60.
6. Hanahan D, Weinberg RA. The hallmarks of cancer. Cell. 2000;100(1):57–70.
7. Domius MD, Dijkstra H, Oudkerk M, Sijens PE. Effect of b value and pre-admission of contrast on diagnostic accuracy of 1.5-T breast DWI: a systematic review and meta-analysis. Eur Radiol. 2014;24(11):2835–47.
8. Chen X, Li WL, Zhang YL, Wu Q, Guo YM, Bai ZL. Meta-analysis of quantitative diffusion-weighted MR imaging in the differential diagnosis of breast lesions. BMC Cancer. 2010;10:693.
9. Kirchner J, Grueneisen J, Martin O, Oehmigen M, Quick HH, Bittner AK, et al. Local and whole-body staging in patients with primary breast cancer: a comparison of one-step to two-step staging utilizing (18)F-FDG-PET/MRI. Eur J Nucl Med Mol Imaging. 2018;45(13):2328–337.
10. Melsaether A, Moy L. Breast PET/MR Imaging. Radiol Clin N Am. 2017;55(3):579–89.
11. Pinker K, Bogner W, Balzer P, Karanikas G, Magometschnigg H, Brader P, et al. Improved differentiation of benign and malignant breast tumors with multiparametric 18fluorodeoxyglucose positron emission tomography magnetic resonance imaging: a feasibility study. Clin Cancer Res. 2014;20(13):3540–9.
12. Ostenson J, Pujara AC, Miheev A, Moy L, Kim SG, Melsaether AN, et al. Voxelwise analysis of simultaneously acquired and spatially correlated (18) F-fluorodeoxyglucose (FDG)-PET and intravoxel incoherent motion metrics in breast cancer. Magn Reson Med. 2017;78(3):1147–56.
13. Margolis NE, Moy L, Sigmund EE, Freed M, McKellogg J, Melsaether AN, et al. Assessment of aggressiveness of breast cancer using simultaneous 18F-FDG-PET and DCE-MRI: preliminary observation. Clin Nucl Med. 2016;41(8):e355–61.
14. King V, Brooks JD, Bernstein JL, Reiner AS, Pike MC, Morris EA. Background parenchymal enhancement at breast MR imaging and breast cancer risk. Radiology. 2011;260(1):50–60.
15. Dontchos BN, Rahbar H, Partridge SC, Kordje L, Lam DL, Scheel JR, et al. Are qualitative assessments of background parenchymal enhancement, amount of fibroglandular tissue on MR images, and mammographic density associated with breast cancer risk? Radiology. 2015;276(2):371–80.
16. Leitlner D, Baltzer PA, Magometschnigg HF, Wengert GJ, Karanikas G, Helbich TH, et al. Quantitative assessment of breast parenchymal uptake on 18F-FDG PET/CT: correlation with age, background parenchymal enhancement, and amount of fibroglandular tissue on MRI. J Nucl Med. 2016;57(10):1518–22.
17. Mema E, Mango VL, Guo X, Karcich J, Yeh R, Wynn RF, et al. Does breast MRI background parenchymal enhancement indicate metabolic activity? Qualitative and 3D quantitative computer imaging analysis. J Magn Reson Imaging. 2018;47(3):753–9.
18. Vranjesvic D, Schiepers C, Silverman DH, Quon A, Villalpando J, Dahlbom M, et al. Relationship between 18F-FDG uptake and breast density in women with normal breast tissue. J Nucl Med. 2003;44(8):1238–42.
19. Mann RM, Balleyguier C, Baltzer PA, Bick U, Colin C, Cormford E, et al. Breast MRI: EUSOBI recommendations for women’s information. Eur Radiol. 2015;25(12):3669–78.
20. Bogner W, Pinker-Domenig K, Bickel H, Chmelik M, Weber M, Helbich TH, et al. Readout-segmented echo-planar imaging improves the diagnostic performance of diffusion-weighted MR breast examinations at 3.0 T. Radiology. 2012;263(1):64–76.
21. Pinker K, Grubner G, Bogner W, Gruber S, Szomolanyi P, Trattnig S, et al. A combined high temporal and high spatial resolution 3 tesla MR imaging protocol for the assessment of breast lesions: initial results. Invest Radiol. 2009;44(9):53–8.
22. ACR BI-RADS® Atlas. Breast Imaging Reporting and Data System. 5th ed. Reston, VA: American College of Radiology; 2013.
23. Rosset A, Spadola L, Ratib O. OsiriX: an open-source software for navigating in multidimensional DICOM images. J Digit Imaging. 2004;17(3):205–16.
24. Knuttel FM, van der Velden BH, Loo CE, Elias SG, Wesseling J, van den Bosch MA, et al. Prediction model for extensive ductal carcinoma in situ around early-stage invasive breast cancer. Invest Radiol. 2016;51(7):462–8.
25. Rahbar H, Partridge SC, Demartini WB, Gutierrez RL, Allison KH, Peacock S, et al. In vivo assessment of ductal carcinoma in situ grade: a model incorporating dynamic contrast-enhanced and diffusion-weighted breast MR imaging parameters. Radiology. 2012;263(2):374–82.
26. Cheeneey S, Rahbar H, Dontchos BN, Javid SH, Rendi MH, Partridge SC. Apparent diffusion coefficient values may help
predict which MRI-detected high-risk breast lesions will upgrade at surgical excision. M. J Magn Reson Imaging. 2017;46(4):1028–36.

27. Quintela-Fando M, Luch A, Manso L, Calvo I, Cortes J, Garcia-Saenz JA, et al. (18)F-fluorosimodizidazole PET and activity of neo-adjuvant Nintedanib in early HER2-negative breast cancer: a window-of-opportunity randomized trial. Clin Cancer Res. 2017;23(6):1432–41.

28. de Almeida Schirmer BG, de Araujo MR, Silveira MB, Pereira JM, Vieira LC, Alves CG, et al. Comparison of [(18)F]Fluorocholine and [(18)F]Fluorodeoxyglucose for assessment of progression, lung metastasis detection and therapy response in murine 4T1 breast tumor model. Appl Radiat Isot. 2018;140:278–88.

29. Wyatt LC, Lewis JS, Andreev OA, Lewis JS, et al. Targeting breast tumors with pH (low) insertion peptides. Mol Pharm. 2014;11(8):2896–905.

30. Demoin DW, Wyatt LC, Edwards KJ, Abdel-Atti D, Sarparanta M, Pourat J, et al. PET imaging of extracellular pH in tumors with (64)cu- and (18)F-labeled pHLIP peptides: a structure-activity optimization study. Bioconjug Chem. 2016;27(9):2014–22.

31. Liberman L, Mason G, Morris EA, Dershaw DD. Does size matter? Positive predictive value of MRI-detected breast lesions as a function of lesion size. AJR Am J Roentgenol. 2006;186(2):426–30.

32. Wang LC, DeMartini WB, Partridge SC, Peacock S, Lehman CD. MRI-detected suspicious breast lesions: predictive values of kinetic features measured by computer-aided evaluation. AJR Am J Roentgenol. 2009;193(3):826–31.

33. Gutierrez RL, DeMartini WB, Eby PR, Kurland BF, Peacock S, Lehman CD. BI-RADS lesion characteristics predict likelihood of malignancy in breast MRI for masses but not for nonmasslike enhancement. AJR Am J Roentgenol. 2009;193(4):994–1000.

34. Pinker-Domenig K, Bogner W, Gruber S, Bickel H, Duffy S, Schernthaner M, et al. High resolution MRI of the breast at 3 T: which BI-RADS(R) descriptors are most strongly associated with the diagnosis of breast cancer? Eur Radiol. 2012;22(2):322–30.

35. Bennani-Baiti B, Dietzel M, Baltzer PA. Correction: MRI background parenchymal enhancement is not associated with breast cancer. PLoS One. 2016;11(9):e0162936.

36. Grimm LJ, Saha A, Ghate SV, Kim C, Soo MS, Yoon SC, et al. Relationship between background parenchymal enhancement on high-risk screening MRI and future breast cancer risk. Acad Radiol. 2019;26(1):69–75.

37. Pike MC, Pearce CL. Mammographic density, MRI background parenchymal enhancement and breast cancer risk. Ann Oncol. 2013;24(Suppl 8):viii37–41.

38. Ramadan S, Arm J, Silcock J, Santamaria G, Buck J, Roy M, et al. Lipid and metabolite deregulation in the breast tissue of women carrying BRCA1 and BRCA2 genetic mutations. Radiology. 2015;275(3):675–82.

39. Partridge SC, Nissan N, Rahbar H, Kitsch AE, Sigmund EE. Diffusion-weighted breast MRI: clinical applications and emerging techniques. J Magn Reson Imaging. 2017;45(2):337–55.

40. Pinker K, Moy L, Sutton EJ, Mann RM, Weber M, Thakur SB, et al. Diffusion-weighted imaging with apparent diffusion coefficient mapping for breast Cancer detection as a stand-alone parameter: comparison with dynamic contrast-enhanced and multiparametric magnetic resonance imaging. Invest Radiol. 2018;53(10):587–95.

41. Partridge SC, DeMartini WB, Kurland BF, Eby PR, White SW, Lehman CD. Quantitative diffusion-weighted imaging as an adjunct to conventional breast MRI for improved positive predictive value. AJR Am J Roentgenol. 2009;193(6):1716–22.

42. Ei Khouli RH, Jacobs MA, Mezban SD, Huang P, Kamel IR, Macura KJ, et al. Diffusion-weighted imaging improves the diagnostic accuracy of conventional 3.0-T breast MR imaging. Radiology. 2010;256(1):64–73.

43. Baltzer A, Dietzel M, Kaiser CG, Baltzer PA. Combined reading of contrast enhanced and diffusion weighted magnetic resonance imaging by using a simple sum score. Eur Radiol. 2016;26(3):884–91.

44. Pinker K, Bickel H, Helbig TH, Gruber S, Dubskey P, Plutschog U, et al. Combined contrast-enhanced magnetic resonance and diffusion-weighted imaging reading adapted to the “breast imaging reporting and data system” for multiparametric 3-T imaging of breast lesions. Eur Radiol. 2013;23(7):1791–802.

45. Martinicich L., Deantonio V, Bertotto I, Redana S, Kubatzi F, Sarotto I, et al. Correlations between diffusion-weighted imaging and breast cancer biomarkers. Eur Radiol. 2012;22(7):1519–28.

46. Bickel H, Pinker-Domenig K, Bogner W, Spick C, Bago-Horvath Z, Weber M, et al. Quantitative apparent diffusion coefficient as a noninvasive imaging biomarker for the differentiation of invasive breast cancer and ductal carcinoma in situ. Investig Radiol. 2015;50(2):95–100.

47. Horvat JV, Durando M, Milans S, Patil S, Massler J, Gibbons G, et al. Apparent diffusion coefficient mapping using diffusion-weighted MRI: impact of background parenchymal enhancement, amount of fibroglandular tissue and menopausal status on breast cancer diagnosis. Eur Radiol. 2018;28(6):2516–24.

48. McDonald ES, Schopp JG, Peacock S, DeMartini WB, Rahbar H, Lehman CD, et al. Diffusion-weighted MRI: association between patient characteristics and apparent diffusion coefficients of normal breast fibroglandular tissue at 3 T. AJR Am J Roentgenol. 2014;202(5):W496–502.

49. Partridge SC, McKinnon GC, Henry RG, Hylton NM. Menstrual cycle variation of apparent diffusion coefficients measured in the normal breast using MRI. J Magn Reson Imaging. 2001;14(4):433–8.

50. O’Flynn EA, Morgan VA, Giles SL, deSouza NM. Diffusion weighted imaging of the normal breast: reproducibility of apparent diffusion coefficient measurements and variation with menstrual cycle and menopausal status. Eur Radiol. 2012;22(7):1512–8.

51. Bogner W, Gruber S, Pinker K, Grabner G, Stadlbauer A, Weber M, et al. Diffusion-weighted MR for differentiation of breast lesions at 3 T: how does selection of diffusion protocols affect diagnosis? Radiology. 2009;253(2):341–51.

52. Bickel H, Pinker K, Polanec S, Magometschnigg H, Wengert G, Spick C, et al. Diffusion-weighted imaging of breast lesions: region-of-interest placement and different ADC parameters influence apparent diffusion coefficient values. Eur Radiol. 2017;27(5):1883–92.

53. Baltzer PA, Dietzel M, Vag T, Burmeister H, Gajda M, Camara O, et al. Clinical MR mammography: impact of hormonal status on background enhancement and diagnostic accuracy. RoFo. 2011;183(5):441–7.

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