Automated volumetric radiomic analysis of breast cancer vascularization improves survival prediction in primary breast cancer

Matthias Dietzel¹, Rüdiger Schulz-Wendtland¹, Stephan Ellmann¹, Ramy Zoubi³, Evelyn Wenkel¹, Matthias Hammon¹, Paola Clauser², Michael Uder¹, Ingo B. Runnebaum⁴ & Pascal A. T. Baltzer²*

To investigate whether automated volumetric radiomic analysis of breast cancer vascularization (VAV) can improve survival prediction in primary breast cancer. 314 consecutive patients with primary invasive breast cancer received standard clinical MRI before the initiation of treatment according to international recommendations. Diagnostic work-up, treatment, and follow-up was done at one tertiary care, academic breast-center (outcome: disease specific survival/DSS vs. disease specific death/DSD). The Nottingham Prognostic Index (NPI) was used as the reference method with which to predict survival of breast cancer. Based on the MRI scans, VAV was accomplished by commercially available, FDA-cleared software. DSD served as endpoint. Integration of VAV into the NPI gave NPI\textsubscript{VAV}. Prediction of DSD by NPI\textsubscript{VAV} compared to standard NPI alone was investigated (cox regression, likelihood-test, predictive accuracy: Harrell’s 𝐶, Kaplan Meier statistics and corresponding hazard ratios/HR, confidence intervals/CI). DSD occurred in 35 and DSS in 279 patients. Prognostication of the survival outcome by NPI (Harrell’s 𝐶 = 75.3%) was enhanced by VAV (NPI\textsubscript{VAV}; Harrell’s 𝐶 = 81.0%). Most of all, the NPI\textsubscript{VAV} identified patients with unfavourable outcome more reliably than NPI alone (hazard ratio/HR = 4.5; confidence interval/CI = 2.14-9.58; 𝑃 = 0.0001). Automated volumetric radiomic analysis of breast cancer vascularization improved survival prediction in primary breast cancer. Most of all, it optimized the identification of patients at higher risk of an unfavorable outcome. Future studies should integrate MRI as a “gate keeper” in the management of breast cancer patients. Such a “gate keeper” could assist in selecting patients benefitting from more advanced diagnostic procedures (genetic profiling etc.) in order to decide whether a more aggressive therapy (chemotherapy) is warranted.

Breast cancer is the most common malignant neoplasm of women in the western world. Despite substantial advances in diagnosis and treatment, morbidity and mortality of breast cancer remain high. By adjusting the treatment to the individual cancer biology, precision medicine aims to both improve disease survival and to reduce the rate of unnecessary cytotoxic treatment.

To achieve this goal, accurate biomarkers are essential tools for precision medicine. They help us to understand tumor biology and allow estimation of patient outcome. The Nottingham Prognostic Index (NPI) is a well-established and widely used method for the prediction of survival of primary breast cancer. It is based on three classic biomarkers of breast cancer: tumor size, tumor grade, and the number of metastatic loco-regional lymph nodes.

Breast MRI (MRI) is a functional imaging method that has been validated in clinical practice for over 30 years. Objective semi-quantitative analysis of MRI by automated, FDA-cleared software has been intensely...
evaluated over the last decade and has been clinically established for several years. As automated analysis of MRI provides objective measures of pathological tissue vascularization and quantifies tumor response to cytotoxic treatment, it can be considered a typical imaging biomarker. In the past several years, the feasibility of MRI as a prognostic biomarker has been demonstrated. Hereby, breast MRI dynamically investigates the cancer in real-time within the whole tumor volume in vivo. This concept offers some advantages over traditional biomarkers which are typically evaluated on a small tumor sample ex vivo. To achieve a high prognostic accuracy, the complex imaging data of breast MRI has to be investigated by a dedicated method. This method of correlating the extensive MRI imaging data with outcome parameters is referred to as radiomic analysis.

This study aimed to investigate whether automated volumetric radiomic analysis of breast cancer vascularization (VAV) can improve survival prediction in primary breast cancer.

**Methods**

**Patients.** This investigation was designed as a retrospective observational single-center study. Data were collected at an academic, tertiary care institution run by interdisciplinary breast cancer specialists from the departments of gynaecology, oncology, pathology, radiation oncology, and radiology. This interdisciplinary breast center is certified according to national quality management criteria. The local ethical review board approved this study and waived the necessity for informed consent.

The following inclusion criteria were applied:

- Patients were recruited over a consecutive time period
- Indication for MRI: Pre-treatment staging of lesions rated as category IV and V according to the Breast Imaging Reporting and Data System (BI-RADS) upon mammography and/or ultrasound
- Histologically verified diagnosis of invasive breast cancer
- Treatment and follow-up at our center.
- No neoadjuvant chemotherapy performed.

In case of recurrent or in situ breast cancer and known malignant neoplasms other than breast cancer, patients were excluded due to potential bias on MRI enhancement results or on outcome data. In addition to image-based analyses, data on demographic characteristics (sex, age, date of diagnosis), clinicopathological features, and individual patient outcome were collected.

**Patient outcome.** The end point was defined according to the “Standardized Definitions for Efficacy End Points in Adjuvant Breast Cancer Trials” document. Disease specific death (DSD) was used as the primary end point. The latter is an established variant to “overall survival” only. For DSD only, “death from breast cancer” is considered as “event”. So different from OS, cases with either “death from non-breast cancer cause” or “death from unknown cause” were not considered, but were censored. The remaining patients were defined as showing “disease specific survival” (DSS).

The analysis of patient outcome was based on routine follow-up at our center after completion of treatment. Survival time was documented as the interval [months] between the initial diagnosis and the last follow-up. The minimum follow-up time after treatment was 27 months for inclusion into the DSS group. Classification of patient outcome was established by the medical record of the last follow-up. In order to achieve valid results upon survival analysis, a mean survival time of at least 80 months was specified.

**Clinicopathological features.** Board-certified pathologists investigated the clinicopathological features, including standard measures of tumor size and disease extension (T1-T3, T4a–T4d). The number of resected lymph nodes was documented and further classified as N0–N3. Histological subtypes were specified as proposed by the World Health Organization Classification of Tumours. Tumor aggressiveness was classified as Grade 1 to 3 according to. Further details on clinicopathological features, including those not relevant for the NPI, are provided within the supplementary material.

Percentage of tumor cells expressing steroid receptors (estrogen/progesterone receptors: ER/PR) was evaluated (cut-off 1% of positive cells). The expression of the HER2 (Human epidermal growth factor receptor) was analyzed as specified in. Ki-67 was not generally recommended at date of study conception, and so it was not included in the present analysis. Further details on the clinicopathological features are provided within the supplementary material.

**Magnetic resonance imaging.** For data acquisition, we used standard MRI protocols, with an overall examination time of 14 minutes. To achieve homogeneous enhancement data, every patient was imaged with the same imaging protocol using two 1.5 Tesla whole-body magnetic resonance imaging scanners (Magnetom Symphony; Siemens Healthineers, Erlangen, Germany) and a dedicated, vendor-supplied, bilateral, four-channel breast coil.

This imaging protocol was in accordance with international recommendations. For the present analysis, dynamic T1-weighted gradient-echo images only were investigated, with 60 seconds temporal resolution and an examination time of eight minutes (fast low angle shot/FLASH; flip angle 80°, repetition time 113 ms, echo time 5 ms, voxel size: 1.1 × 0.9, slice thickness 3 mm). This dynamic sequence was acquired before and after intravenous bolus injection (automatic injector: Spectris, Medrad, Pittsburgh, USA) of gadopentetate dimeglumine (Bayer HealthCare, Leverkusen, Germany) at a dosage of 0.1 mmol/kg body weight followed by 20 ml saline solution. A delay of 30 seconds after contrast medium application was set before post-contrast images were acquired.
under identical tuning conditions for a total of 7 measurements. Further technical details of the MRI protocol are provided within the supplementary material.

**Volumetric analysis of breast cancer vascularization (VAV).** Raw data of MRI were analyzed by automated, FDA-approved, commercially available software (BreVis, Siemens Healthineers, Erlangen, Germany). The software was operated by a reader blinded to clinicopathological features and patient outcome. The reader received special training in handling the software and was supervised by two radiologists with at least 7 years of experience in breast MRI. This kind of software has been scientifically evaluated since 2005 in multiple institutions, and has been in clinical use for several years. A detailed description can be found in the literature10–12.

The only user-dependent step during analysis of MRI was defining the volume of interest. This was achieved by encircling a rectangular cuboid around the tumor (see Fig. 1)29. Based on our experience with the given protocol over many years, the minimum initial enhancement threshold was set to 30% (change of signal intensity first minute after contrast relative to baseline in [%]). As a consequence, perifocal tissue or necrotic tumor components were automatically excluded from further analysis. In case of multiple tumors per breast, the largest one was defined as index lesion.

Based on established dynamic enhancement patterns28, the **vascularization pattern of every voxel** within the tumor was characterized. In total, the software automatically extracted 14 quantitative and semi-quantitative radiomic features for the VAV. These parameters assessed three aspects of tumor vascularization:

I. **Total enhancing tumor volume (TTV)** characterized the amount of vital cancer tissue and was measured as [cm³]. It was defined as the sum of tumor voxels surpassing the initial enhancement threshold (see above). Accordingly, the TTV addressed only the vital cancer tissue and did not consider areas of necrosis. Therefore, the TTV has also been called “functional tumor volume” previously (Table 1, parameter #1)18.

II. **Heterogeneity of vascularization (within the TTV).** Breast cancer is a heterogeneous disease and various vascularization patterns can be present in the same cancer in parallel. Accordingly, a wide range of vascularisation patterns present within the same tumor. A relevant proportion of the TTV (10.3%) exhibited a pattern typical of hypervascularization and vascular shunting (category I: strong wash-in and wash-out). Nevertheless, the majority of the tumor demonstrated patterns indicative of a less pronounced neovascularization (weak or intermediate wash-in and persistent: 55.9% of the TTV). Both the NPI and the VAV were suggestive of a poor outcome. This patient died from breast cancer 18 months after initial diagnosis.

---

**Figure 1.** Clinical case illustrating the assessment of VAV. MRI of a 51 year old female patient (invasive ductal cancer, G2, diameter 5.8 cm, T4c, HER2 negative, hormonal receptor negative, 5 positive lymph nodes, NPI = 6.2). (A) Heat map of the color-coded CAD overlay of the T1 pre-contrast scan reflecting heterogeneity of vascularization. The colors are coded from red to blue and reflect the wash-out ratio (see main text for definition). A large heterogeneously enhancing mass with infiltration of skin and thoracic wall and central necrosis is visualized, consistent with T4c. Note the discrepancy between the vital tumor assessed by MRI (color coded parts. TTV) and the larger morphologic extensions including the central necrosis. (B) Vascularization of the most suspect tumor compartment. A strong wash-in (170%) followed by a significant wash-out (50%) is demonstrated. The time to peak enhancement (TTP) was one minute. (C) Heterogeneity of vascularization summarized in a 3 × 3 matrix. There was a wide range of vascularisation patterns present within the same tumor. A relevant proportion of the TTV (10.3%) exhibited a pattern typical of hypervascularization and vascular shunting (category I: strong wash-in and wash-out). Nevertheless, the majority of the tumor demonstrated patterns indicative of a less pronounced neovascularization (weak or intermediate wash-in and persistent: 55.9% of the TTV). Both the NPI and the VAV were suggestive of a poor outcome. This patient died from breast cancer 18 months after initial diagnosis.
Initially, the enhancement patterns within each voxel were identified following international breast MRI standards. The initial enhancement phase investigates the first minute after contrast application. It was categorized as weak (30% to 50%), intermediate (50% to 100%), or strong "wash-in" (>100%), with percentages expressing the change of signal intensity during the first minute after contrast media application, relative to baseline.

The delayed enhancement phase investigates the change of contrast enhancement between the first and the last minute after contrast media application. It was categorized as persistent (>10%), plateau (±10%), or wash-out (<10%). Percentages express the change of signal intensity during the last minute after contrast media application, relative to the first postcontrast scan.

Combining the initial phase (n = 3) and delayed phase categories (n = 3) yielded nine enhancement patterns to classify the vascularization of each voxel within the TTV (for instance as "weak wash-in" and "persistent", etc.).

Finally, the tumor volume was subdivided according to these nine enhancement patterns. Hereby, the proportion relative to the TTV [%] was identified for every enhancement pattern (Table 1, parameters #2–10).

III. Vascularization of the most suspect tumor compartment. In this analysis only one particular compartment of the entire tumor volume was investigated. For this reason, the TTV was divided into clusters (size: 3 × 3 voxels). Within every cluster, the ratio of the corresponding initial versus delayed enhancement was computed (wash-out ratio). According to the literature, the cluster achieving the highest wash-out ratio was defined as the "most suspicious tumor compartment".

The "most suspicious tumor compartment" was characterized by the following four radiomic features (Table 1, parameters #11–14): "wash-out ratio" (see above), "wash-in" (signal intensity first minute after contrast relative to baseline [%]), "peak enhancement" (maximum contrast uptake by the tumor during the whole dynamic scan), and time to "peak enhancement" (TTP [minutes]).

A clinical case illustrating the workflow of the volumetric analysis of breast cancer vascularization is given in Fig. 1. Table 1 summarizes all 14 parameters provided by the VAV.
**Statistical analysis.** Disease specific death (DSD) was defined as the primary study endpoint. Association of clinicopathological features, patient age, NPI, and NPI\textsubscript{VAV} with survival outcome was explored by cox regression and likelihood ratio tests. All tests were two-sided. In this exploratory study, an alpha error below 10% was defined as appropriate to reject the null hypothesis. Considering the total number of variables (n = 14), we applied a conventional Bonferroni correction for alpha-error accumulation. Accordingly, the significance level was set to $P = 0.007$.

**NPI.** According to\textsuperscript{7}, the NPI was calculated as follows:

$$NPI = [0.2 \times S] + N + G$$

Hereby, $S$ is the size of the index lesion in centimeters, $N$ is the nodal status (0 nodes = 1; 1–4 nodes = 2; >4 nodes = 3), and $G$ is the tumor grade (Grade I = 1; Grade II = 2; Grade III = 3).

The association of NPI with survival was investigated by cox regression, and predictive values were saved. The corresponding performance of NPI to predict DSD was validated by the Harrell’s C index. The latter is an extension of the area under the ROC curve for censored survival data\textsuperscript{31}.

**NPI\textsubscript{VAV}.** NPI\textsubscript{VAV} was designed as a compound measure of NPI and VAV. VAV revealing significant association with DSD upon univariate analysis were eligible for this step. Multivariate cox regression with backward feature selection was applied ($P$ for enter/removal: 0.001/0.05). Feature selection removed redundant or irrelevant VAV. Furthermore, feature selection enabled simplification and – by reducing overfitting – generalization of the model. Corresponding predictive values of the multivariate model were saved and investigated by the Harrell’s C index as described for NPI.

**Comparison of outcome prediction.** Finally, predictions of DSD by NPI and VAV were compared. For this purpose, the corresponding predictive values by the cox regressions were compared\textsuperscript{32}.

Kaplan Meier survival analysis was used to graphically illustrate differences between NPI\textsubscript{VAV} and NPI. Hereby, the cut-off criterion for the prediction of DSD was optimized to the value that maximized Youden’s J statistic on the time-dependent data. For the latter, we used the predictive values of the corresponding cox regression analysis of NPI and NPI\textsubscript{VAV}\textsuperscript{31,34}.

The differences between the corresponding Kaplan Meier survival curves were explored by hazard ratios (HR), corresponding confidence intervals (CI), and the logrank test\textsuperscript{23}.

**Statement of human rights.** 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. For this type of study formal consent is not required.

**Results**

A detailed listing of background clinical data and clinicopathological factors is given in the supplementary material. This also contains a detailed uni- and multivariate analysis of study data including subgroup analysis.

**Characteristics of patients and subgroups.** There were 314 patients with primary breast cancer who were included (DSS = 279; DSD = 35). Mean survival time was 84.6 months (CI: 81.9–87.2, standard error: 1.3). Range of follow-up interval was 27–93 months for DSS. On average, DSD occurred 27.9 months after initial diagnosis (range: 3–70 months).

Mean patient age was 57.6 years (SD: 11.7). Mean age within the DSS (mean: 57.4 years; SD: 11.7) and the DSD group was similar (mean: 59.6 years; SD: 11.5). Patient age could not be identified as a predictive parameter upon univariate analysis ($P = 0.23$). Inclusion of patient age into the NPI did not significantly increase prediction of DSD ($P = 0.13$). The same results were observed when including patient age into the NPI\textsubscript{VAV} ($P = 0.23$).

**Clinicopathological features.** Clinicopathological features showed expected distributions between the DSS and DSD group: There was a significant association with T- and N-Stage versus survival outcome ($P < 0.001$). Notably, the DSD exhibited larger tumors (e.g., T3: HR = 9.4; $P < 0.0001$) with a higher proportion of nodal-positive cases (e.g., N2: HR = 3.8; $P = 0.0002$). There was no significant association with Grading ($P = 0.96$) and histological subtypes ($P > 0.77$) on survival outcome. Key results of the clinicopathological features, stratified by the endpoints, are summarized in Table 2.

**NPI.** Mean NPI was 4.4 (95% CI = 4.3–4.5). In patients with DSD, the mean NPI (5.2; CI = 4.8–5.6) was significantly higher compared to the DSS group (4.3; CI = 4.1–4.4; $P < 0.001$). Harrell’s C was 75.3% for NPI. Descriptive statistics of the NPI are summarized in Table 3.

**VAV.** VAV revealed significant association with the endpoint: Time to peak enhancement (TTP: $P = 0.003$), total tumor volume [cm$^3$] ($P < 0.0001$) and one parameter focusing on the heterogeneity of vascularization...
(category A: \( P = 0.006 \)) were significant predictors of DSD. Predictive performance of VAV reached Harrell's \( C = 74.0\% \) and was equal to that of NPI \( (P = 0.92) \). Key VAV stratified by the endpoints are summarized in Table 3.

In addition, peak enhancement was slightly higher in the DSD group (DSD/DSS = 123.5/110%; \( P = 0.07 \)), as was wash-in (DSD/DSS: 119.6/107.1%; \( P = 0.08 \)). Both parameters, however, missed the significance level.

**NPI versus NPI\textsubscript{VAV}.** Feature selection retained the NPI and all three parameters of VAV in the final NPI\textsubscript{VAV} model (Table 4). Corresponding model fit was superior compared to NPI alone (Chi-squared: 51.5 vs. 27.9). Harrell's C was 81.0% for NPI\textsubscript{VAV} and surpassed the value for NPI by 5.6%.

### Table 2. Summary table of tumor characteristics. Note: tumor characteristics stratified by the clinical outcome given as DSD and DSS (disease specific survival and death). *A more detailed overview of clinicopathological characteristics is given in the supplementary material.*

| Parameter                        | T1a | T1b | T1c | T2 | T3 | T4 |
|----------------------------------|-----|-----|-----|----|----|----|
| **Outcome**                      |     |     |     |    |    |    |
| DSS                              | 26  | 43  | 117 | 80 | 7  | 6  |
| DSD                              | 1   | 3   | 6   | 17 | 4  | 4  |
| **Total**                        | 27  | 46  | 123 | 97 | 11 | 10 |

| Parameter                        | Outcome | T-stage | Histological subtype | Histological Grading |
|----------------------------------|---------|---------|-----------------------|----------------------|
|                                  |         | T1a     | T1b                   | T1c                  |
| **T-stage**                      |         | 26      | 43                    | 117                  |
| **Histological subtype**         |         |         |                       |                      |
| Invasive ductal (not otherwise specified) | 212 | 23 | 38 | 3  |
| Invasive lobular                  | 76.00%  | 8.20%   | 13.60%                | 1.10%                |
| Mixed (Invasive lobular and ductal) | 77.10% | 11.40%  | 11.40%                | 0.00%                |
| Invasive medullary                | 76.10%  | 8.60%   | 13.40%                | 1.00%                |
| Invasive mucinous                 | 76.00%  | 8.20%   | 13.60%                | 1.10%                |
| **Histological Grading**         |         |         |                       |                      |
| G1                                | 12      | 0       | 3                     | 12                   |
| G2                                | 108     | 13      | 3                     | 121                  |
| G3                                | 159     | 22      | 3                     | 181                  |

| Survival                         | Total tumor volume (TTV [cm³]) | Heterogeneity of vascularization* | Time-to-peak enhancement (TTP [min]) | NPI | Age [years] |
|----------------------------------|---------------------------------|----------------------------------|-------------------------------------|-----|-------------|
| Minimum                          | 0.1                             | 2.4                              | 1                                   | 1.18| 43          |
| Maximum                          | 313.1                           | 249.7                            | 5.4                                 | 7.5 | 87          |
| Mean                              | 7.1                             | 29.9                             | 23                                  | 1.18| 43          |
| Median                            | 2.9                             | 7.6                              | 23                                  | 1   | 43          |
| SD                                | 21.1                            | 60.4                             | 8.6                                 | 1.12| 12          |

**Table 3.** Descriptive statistics of parameters retained in the NPI\textsubscript{VAV} model and patient age. Note: Given are all four parameters retained in the NPI\textsubscript{VAV} model and patient age (see Table 4). Corresponding results of descriptive statistics are listed. Except patient age \((P = 0.23)\), all parameters were predictive of the endpoint (for \( P \)-values, see Table 4). *Weak wash-in (initial phase) and persistent (delayed phase) [%] (details see Table 1 and main text).**
The improved survival prediction by VAV was also demonstrated by the Kaplan Meier analysis (Fig. 2):

**Prediction of DSD.** The survival curves of patients in whom NPIVAV predicted DSD significantly differed from those in whom NPI alone predicted DSD (logrank test: \( P = 0.0001 \)). DSD was more likely to occur, if this event was predicted by NPIVAV, compared to NPI alone (HR = 4.5; CI: 2.14–9.58).

**Probability of DSS.** The survival curves of patients classified as DSS either by NPIVAV or by NPI were not significantly different (logrank test: \( P = 0.65 \)). The risk of DSS was similar, if DSS was predicted by NPIVAV, compared to the patients in whom NPI alone predicted DSD (HR = 1.05; CI: 0.86–1.27, \( P_{\text{logrank}} = 0.65 \)).

Table 4. NPI\textsubscript{VAV} model. Note: Given is the NPI\textsubscript{VAV} model as provided by the Cox regression. After applying feature selection, four covariates were retained. Besides NPI this included three parameters for MRI analysis (for details, see Table 1). CI: 95% confidence interval of the hazard ratio. SE: standard error of the coefficient. *Weak wash-in (initial phase) and persistent (delayed phase) [%] (details see Table 1 and main text).

| Covariate                              | HR   | CI              | P     | Coefficient | SE  |
|---------------------------------------|------|-----------------|-------|-------------|-----|
| Total tumor volume (TTV)              | 1.01 | 1.00 to 1.01    | 0.04  | 0.01        | 0.00|
| Heterogeneity of vascularization*     | 0.95 | 0.92 to 0.98    | 0.002 | –0.05       | 0.02|
| Time-to-peak enhancement (TTP)        | 1.84 | 1.60 to 2.42    | <0.0001 | 0.61      | 0.14|
| Nottingham Prognostic Index (NPI)     | 2.01 | 1.48 to 2.73    | <0.0001 | 0.70      | 0.16|

Figure 2. Prediction of good (DSS) and poor (DSD) patient outcome: Comparison of NPI\textsubscript{VAV} vs. standard NPI. NPI\textsubscript{VAV} enabled a better identification of patients at risk for DSD compared to standard NPI (HR = 4.5, CI: 2.14–9.58). This was also evident by a faster decline of the corresponding survival curve (\( P_{\text{logrank}} = 0.0001 \)). Optimized identification of high risk patients by the NPI\textsubscript{VAV} did not come on the price of a worse identification of patients with a more favorable outcome. Indeed, the likelihood of DSS was alike if NPI\textsubscript{VAV} was used to predict DSS, compared to standard NPI alone (HR = 1.05, CI: 0.86–1.27, \( P_{\text{logrank}} = 0.65 \)).

The improved survival prediction by VAV was also demonstrated by the Kaplan Meier analysis (Fig. 2):

**Prediction of DSD.** The survival curves of patients in whom NPI\textsubscript{VAV} predicted DSD significantly differed from those in whom NPI alone predicted DSD (logrank test: \( P = 0.0001 \)). DSD was more likely to occur, if this event was predicted by NPI\textsubscript{VAV} compared to NPI alone (HR = 4.5; CI: 2.14–9.58).

**Probability of DSS.** The survival curves of patients classified as DSS either by NPI\textsubscript{VAV} or by NPI were not significantly different (logrank test: \( P = 0.65 \)). The risk of DSS was similar, if DSS was predicted by NPI\textsubscript{VAV} compared to the patients in whom NPI alone predicted DSD (HR = 1.05; CI: 0.86–1.27).

**Discussion**

Breast MRI contributed to the prediction of survival in patients with primary invasive breast cancer. Automated volumetric radiomic analysis of breast cancer vascularization improved survival prediction compared to the Nottingham Prognostic Index alone (NPI). The NPI is a well-established and widely used method of predicting survival of primary breast cancer. Most of all, the use of MRI optimized the stratification of patients at risk of an unfavourable survival outcome.

**MRI showed a prognostic accuracy equal to that of the NPI.** To our knowledge, we are the first group to report a head-to-head comparison of MRI and NPI in the risk stratification of patients. One advantage of prognostic MRI is that it is performed as a one-stop-shop procedure within one session. Our approach was based on a standard MRI protocol, as recommended by international guidelines, and required only 14 minutes of magnet time\textsuperscript{28}. Based on this fast examination, both the staging — including the assessment of tumor extension plus multifocal and bilateral disease — and the risk estimation can be accomplished. Whereas classical risk factors such as nodal metastasis and histological grading are nominal or ordinal by nature, MRI provides quantitative and semi-quantitative parameters. This information is semi-automatically assessed by the software, which is an advantage over standard pathological assessment. For instance, the evaluation of grading is known to be limited by significant inter-pathologist variability\textsuperscript{35}. Prognostic MRI information moreover is available in real-time. This is a potential advantage for clinical workflow and patient management. Of note, however, some NPI cannot be assessed on core biopsy samples, but require surgical resection for definitive classification. This is frequently the
case in nodal staging, resulting in the delayed availability of these biomarkers for prognostic purposes. MRI both volumetrically and dynamically investigates the whole tumor volume in vivo. This is an advantage over most prognostic biomarkers that evaluate small tumor samples ex vivo. As MRI is not a snapshot diagnosis, it is well suited to investigate tumor heterogeneity. In fact, volumetric MRI features were among the best prognostic parameters in our analysis.

Our findings support the evaluation of MRI as a prognostic biomarker in precision medicine. The concept of precision medicine aims to adapt cancer treatments to an individual’s tumor characteristics. Thus, the goal of precision medicine is to both maximize the effectiveness of treatment and to minimize the side effects of cytotoxic drugs. As every known biomarker thus far has limitations, there is intensive ongoing research into the development of new tissue biomarkers. One of the most promising fields is gene expression profiling of breast cancer.\textsuperscript{18,36,37}. Gene expression profiling is already used in systemic breast cancer treatment planning,\textsuperscript{9} and empirical evidence shows that the decision towards more aggressive treatments such as cytotoxic drugs can be based on distinct gene profiles.\textsuperscript{37} MRI could actually tread the same path: In our analysis, the strength of MRI was a more accurate identification of patients with a less favorable outcome. Thus, our data could be used to stratify patient risk. Certainly, it would be unwise to apply a more aggressive treatment based only on the MRI information at the time being. Nevertheless, MRI might be used as a “gate keeper”. If MRI indicates an unfavorable outcome, further biomarkers such as genetic profiling might be warranted, in order to decide whether a more aggressive therapy is indicated.

Possible biological correlates for our MRI findings will be discussed. Tumor size is one of the best-established prognostic parameters of breast cancer.\textsuperscript{18} Typically, the measurement of the breast cancer size is performed as linear measurements in two spatial dimensions. Both the NPI and the T-staging are based on this method.\textsuperscript{34,35} The breast carcinoma may grow in a diffuse pattern, as is often observed in invasive lobular subtypes.\textsuperscript{39} Consequentially, the tumor size of breast cancer cannot always be accurately measured by two-dimensional linear measurement. However, TTV is a volumetric method and thus can evaluate the tumor size in any spatial extent.

However, TTV is not only a metric parameter of tumor size. According to the research by Hylton \textit{et al.} it can also be interpreted as a functional biomarker of vascularization.\textsuperscript{18} Neoangiogenesis is regarded as a pivotal step in the development of cancer.\textsuperscript{40} At the same time, neoangiogenesis significantly determines the potential of a breast cancer to metastasize and hereby limits patient outcome.\textsuperscript{41,42} One method to quantify tumor vascularization is the determination of its microvascular density (MVD). It is therefore conclusive that microvascular density can also be considered as a prognostic biomarker of breast cancer.\textsuperscript{42} Interestingly, contrast enhancement assessed by MRI was closely linked to MVD in previous radiopathological correlations studies.\textsuperscript{43} Thus, the prognostic significance of TTV may also be explained in this functional context. However, tumor size alone is not a sufficient parameter to optimally predict the prognosis of a patient. This explains why additional parameters were identified as independent predictors of survival status in the NPI/VAV model.\textsuperscript{44}

Different from the classical pathological analysis as well as from modern methods (such as genetic profiling), MRI can examine the entire tumor volume.\textsuperscript{45-47} For this reason, the method can also be used to assess the intratumoral heterogeneity. The latter is a major aspect in breast cancer biology.\textsuperscript{46} We identified “heterogeneity of vascularization” within the tumor as a significant parameter of patient outcome. Hereby, a higher volume of compartments exhibiting slow wash-in and persistent enhancement corresponded to a more favorable prognosis. Of note, this MRI pattern has been described by Leong \textit{et al.} as typical for breast cancers.\textsuperscript{48} Considering the relationship of MRI enhancement and the MVD (see above), this finding seems conclusive: Buadu reported that breast cancer with slower initial and delayed contrast enhancement in MRI exhibited lower MVD in radiopathological correlation.\textsuperscript{45} Since a lower MVD is regarded as a predictor of a relatively better outcome, this finding seems conclusive.\textsuperscript{42}

The analysis of tumor heterogeneity also enabled the identification of the most suspicious compartment. It was identified based on the parameters wash-in and wash-out. In the context of the aforementioned MVD, both parameters could be regarded as parameters of tissue vascularisation.\textsuperscript{45-47} Within this compartment, a slightly faster wash-in, a slightly higher maximum enhancement, and, above all, a significantly longer TTP were observed in the DSD group. Wash-in can be interpreted as a parameter of tissue perfusion.\textsuperscript{47} It reflects both the blood flow and the intravascular blood volume (for instance figure 5 in\textsuperscript{45}), and correlates with the MVD.\textsuperscript{42,43,47} After the perfusion phase, the contrast medium is transmitted through the blood vessels into the interstitium. This event is referred to as leakage and here the maximum enhancement is typically observed.\textsuperscript{47} However, it needs to be emphasized that an exact separation of the different pharmacokinetic compartments is not feasible with standard breast MRI protocols.\textsuperscript{47} Leakage is not only determined by the quantity (vessel surface), but also by the quality (permeability) of the vessels.\textsuperscript{47} The quantity (vessel surface) correlates with the MVD.\textsuperscript{47,47} The permeability is triggered by numerous factors, among others by VEGF.\textsuperscript{48} VEGF is a cytokine that plays a central role in the process of angiogenesis.\textsuperscript{49} These considerations might explain why the maximum enhancement in the DSD group was slightly higher and slightly delayed compared to the DSS cases. Interestingly, the parameter TTP is rarely used in breast MRI diagnostics. In different fields of radiology, such as in neuro imaging, the TTP is a standard parameter.\textsuperscript{47} This is particularly relevant since early breast MRI research has already shown that TTV is superior to more commonly used qualitative kinetic criteria.\textsuperscript{45} In this context, an in-depth re-evaluation of this parameter may be of promise.

Some limitations of our work will be discussed. The literature reports a large number of breast cancer biomarkers. Hereby, the NPI was introduced in 1982 and remains a widely used method to predict survival in primary breast cancer.\textsuperscript{34,35} A completely different approach is taken by multigene assays. These rather new biomarkers focus on genetic analysis and are increasingly used in clinical routine.\textsuperscript{50} Interestingly, it has already been shown in the literature that there is a close correlation between MRI parameters with multigene assays such
as MammaPrint, Oncotype DX, and PAM50\(^3\). So future studies might investigate to what extent MRI analysis could substitute genetic tests, or whether MRI might actually improve the prognostic information provided by multigene assays.

We investigated radiomic parameters of tumor vascularization. We excluded morphologic features from our analysis\(^2\). Previous studies have shown that also morphologic MRI data provide prognostic information\(^17,52\). The presence of avascular, seemingly necrotic compartments have for instance been associated with surrogates of poor outcome\(^52\). So future studies should investigate textural radiomic features as well in order to determine whether predictive accuracy of VAV can be further increased.

Breast cancer can metastasize after many years and lead to DSD\(^53\). It cannot be excluded that some of our patients died after the end of data collection. This is a potential bias of the study.

We conceptualized this work as a cross-sectional study. In this respect, our data represent a typical patient collective at our practice. Therefore - as well as in the context of the patient number - no formal subgroup analysis according to histopathological parameters (clinicopathological factors, etc.) as well as the treatment regime was performed. Therefore, we cannot exclude that these two parameters had an impact on our results. In the same way, we decided to exclude patients with NAC from the analysis. These patients were predominantly included in clinical trials and were therefore treated with a therapy not yet established. Nevertheless, this implies a potential limitation of our study: Results do not apply on NAC patients.

In distinction to the classical visual MRI analysis, the influence of the reader on the VAV is significantly lower\(^54\). The software works largely automated and the only investigator-dependent step is the definition of the volume of interest. Nevertheless, some observer-dependent bias within our data cannot be excluded. Furthermore, the reproducibility of our results is also influenced by the software itself. Although the software is established on the market, our results might not be completely reproducible on other systems. Finally, the MRI protocol itself can also have influence on our results\(^55\). We aimed to control for this bias and adopted our imaging protocol to international standards and used well established criteria to assess the tumor vascularization\(^16,18,28,29,45,56\). Future studies should investigate the repeatability of our results with different software, MRI protocols and identify the potential impact of the reader on our results.

**Conclusion**

In conclusion, automated volumetric radiomic analysis of breast cancer vasculature improved the prediction of survival of patients with primary invasive breast cancer. Most of all it improved the identification of patients at higher risk of an unfavorable outcome.

These results are based on a standard, clinical MRI followed by real-time analysis by a commercially available, FDA-cleared computer algorithm. Accordingly, the method can be incorporated into the clinical routine.

One potential role of MRI would be a “gate keeper”, in order to identify patients requiring a more advanced diagnosis (genetic profiling etc.) and/or more aggressive therapies (chemotherapy). This approach should be evaluated in future trials.

**Data availability**

All relevant data are within the paper and its supplementary files.

Received: 7 January 2019; Accepted: 4 February 2020;
Published online: 28 February 2020

**References**

1. SEER Stat Fact Sheets: Female Breast Cancer. [http://seer.cancer.gov/statfacts/html/breast.html](http://seer.cancer.gov/statfacts/html/breast.html) (2019).
2. Kurian, A. W. & Friese, C. R. Precision Medicine in Breast Cancer Care: An Early Glimpse of Impact. JAMA Oncology 1, 1109 (2015).
3. Friese, C. R. et al. Chemotherapy decisions and patient experience with the recurrence score assay for early-stage breast cancer. Breast Cancer Recurrence Scores. Cancer 123, 43–51 (2017).
4. Biomarkers Definitions Working Group. Biomarkers and surrogate endpoints: Preferred definitions and conceptual framework. Clinical Pharmacology & Therapeutics 69, 89–95 (2001).
5. Fong, Y. et al. The Nottingham Prognostic Index: five- and ten-year data for all-cause Survival within a Screened Population. The Annals of The Royal College of Surgeons of England 97, 137–139 (2015).
6. Haybittle, J. L. et al. A prognostic index in primary breast cancer. Br J Cancer 45, 361–366 (1982).
7. Todd, J. H. et al. Confirmation of a prognostic index in primary breast cancer. Br J Cancer 56, 489–492 (1987).
8. Blamey, R. W. et al. Survival of invasive breast cancer according to the Nottingham Prognostic Index in cases diagnosed in 1990–1999. European Journal of Cancer 43, 1548–1555 (2007).
9. Kaiser, W. MRI of the female breast. First clinical results. Arch. Int. Physiol. Biochim. 93, 63–76 (1985).
10. Pediconi, F. et al. Color-coded automated signal intensity curves for detection and characterization of breast lesions: preliminary evaluation of a new software package for integrated magnetic resonance-based breast imaging. Invest Radiol 40, 448–457 (2005).
11. Baltzer, P. A. et al. Computer Assisted Analysis of MR-Mammography Reveals Association Between Contrast Enhancement and Occurrence of Distant Metastasis. Technology in cancer research & treatment (2012).
12. Kim, J. J. et al. Computer-aided Diagnosis-generated Kinetic Features of Breast Cancer at Preoperative MR Imaging: Association with Desease-free Survival of Patients with Primary Operable Invasive Breast Cancer. Radiology 162079, [https://doi.org/10.1148/radiol.2017162079](https://doi.org/10.1148/radiol.2017162079) (2017).
13. Johansen, R. et al. Predicting survival and early clinical response to primary chemotherapy for patients with locally advanced breast cancer using DCE-MRI. Journal of Magnetic Resonance Imaging 29, 1300–1307 (2009).
14. Li, S. P. et al. Use of dynamic contrast-enhanced MR imaging to predict survival in patients with primary breast cancer undergoing neoadjuvant chemotherapy. Radiology 260, 68–78 (2011).
15. Pickles, M. D., Lowry, M. & Gibbs, P. Pretreatment Prognostic Value of Dynamic Contrast-Enhanced Magnetic Resonance Imaging Vascular, Texture, Shape, and Size Parameters Compared With Traditional Survival Indicators Obtained From Locally Advanced Breast Cancer Patients. Investigative Radiology 51, 177–185 (2016).
16. Dietzel, M. et al. Association between survival in patients with primary invasive breast cancer and computer aided MRI. J Magn Reson Imaging 37, 146–155 (2013).

---

**Scientific Reports** (2020) 10:3664 | https://doi.org/10.1038/s41598-020-60393-9
Saha, A., Yu, X., Sahoo, D. & Mazurowski, M. A. Effects of MRI scanner parameters on breast cancer radiomics. *Scientific Reports* 10, 3664 (2020).

Li, H.

Dietzel, M.

Gilles, R. J., Kinahan, D. E. & Hricak, H. Radiomics. Images Are More than Pictures, They Are Data. *Radiology* 278, 563–577 (2016).

Hudis, C. A. et al. Proposal for standardized definitions for efficacy end point in adjuvant breast cancer trials: the STEEP system. *J. Clin. Oncol.* 25, 2127–2132 (2007).

Kweldam, C. F., Wildhagen, M. F., Bangma, C. H. & van. Leenders, G. J. L. H. Disease-specific death and metastasis do not occur in patients with Gleason score ≤ 6 at radical prostatectomy. *BJU International* 116, 230–235 (2015).

Survival analysis. MedCalc. https://www.medcalc.org/manual/kaplan-meier-meter.php (2019).

Fattaneh, A. & Tavassoli, F. Tumours of the breast. In World Health Organization Classification of Tumours. Pathology and Genetics. Tumours of the Breast and Female Genital Organs 9–112 (IARC Press, 2003).

Edge, S., Byrd, D., Carducci, M. & Wittke-thind, C. TNM Classification of Malignant Tumours. (Springer, 2009).

Hammond, M. E. H. et al. American Society of Clinical Oncology/College of American Pathologists Guideline Recommendations for Immunohistochemical Testing of Estrogen and Progesterone Receptors in Breast Cancer. *Arch. Pathol. Lab. Med.* 134(6), 907–922 (2010).

Wolff, A. C. et al. American Society of Clinical Oncology/College of American Pathologists guideline recommendations for human epidermal growth factor receptor 2 testing in breast cancer. *Arch. Pathol. Lab. Med.* 131, 18–43 (2007).

Barn, R. M., Kühli, C. K., Kinkel, K. & Boetes, C. Breast MRI: guidelines from the European Society of Breast Imaging. *Eur Radiol* 18, 1307–1318 (2008).

Morris, E. A. et al. ACR BI-RADS® Magnetic Resonance Imaging, in ACR BI-RADS® Atlas, Breast Imaging Reporting and Data System (American College of Radiology, 2013).

Dietzel, M. & Baltzer, P. A. T. How to use the Kaiser score as a clinical decision rule for diagnosis in multiparametric breast MRI: a pictorial essay. *Insights into Imaging* 9, 325 (2018).

Antolins, L., Boracchi, P. & Biganzoli, E. A time-dependent discrimination index for survival data. *Stat Med* 24, 3927–3944 (2005).

DeLong, E. R., DeLong, D. M. & Clarke-Pearson, D. L. Comparing the area under two or more correlated receiver operating characteristic curves: a nonparametric approach. *Biometrics* 44, 837–845 (1988).

Okonkou, E. K. et al. Non-invasive detection of coronary inflammation using computed tomography and prediction of residual cardiovascular risk (the CRISP CT study): a post-hoc analysis of prospective outcome data. *The Lancet* 392, 929–939 (2018).

Youden, W. J. Index for rating diagnostic tests. *Cancer* 3, 32–33 (1950).

Boiesen, P. et al. Histologic grading in breast cancer—reproducibility between seven pathologic departments. South Sweden Breast Cancer Group. *Acta Oncol* 39, 41–45 (2000).

Potosky, A. L. et al. Population-based study of the effect of gene expression profiling on adjuvant chemotherapy use in breast cancer patients under the age of 65 years: Breast Cancer Genetics and Chemotherapy. *Cancer* 121, 4062–4070 (2015).

Barclay, C. H. et al. Outcomes in patients with early-stage breast cancer who underwent a 21-gene expression assay: Outcomes With 21-Gene Expression. *Cancer* 123, 2422–2431 (2017).

Vescera, E. C. et al. Modeling the Effect of Tumor Size in Early Breast Cancer. *Ann Surg* 241, 309–318 (2005).

Reed, A. E. M., Kutasovic, J. R., Lakhani, S. R. & Simpson, P. T. Invasive lobular carcinoma of the breast: morphology, biomarkers and andromics. *Breast Cancer Research* 17, 12 (2015).

Folkman, J. What Is the Evidence That Tumors Are Angiogenesis Dependent? *J Natl Cancer Inst* 82, 4–7 (1990).

Folkman, J. Role of angiogenesis in tumor growth and metastasis. *Semin. Oncol.* 29, 15–18 (2002).

Uzzan, B., Nicolas, P., Cacheret, M. & Perret, G. Y. MicrovesSEL density as a prognostic factor in women with breast cancer: a systematic review of the literature and meta-analysis. *Cancer Res.* 64, 2941–2955 (2004).

Buadu, L. D. et al. Breast lesions: correlation of contrast medium enhancement patterns on MR images with histopathologic findings and tumor angiogenesis. *Radiology* 200, 639–649 (1996).

Sopik, V. & Narod, S. A. The relationship between tumour size, nodal status and distant metastases: on the origins of breast cancer. *Breast Cancer Res Treat* 170, 647–656 (2018).

Leong, I. C. H., Gombos, E. C., Jagadeesan, J. & Fook-Chong, S. M. C. MRI Genetics With Volumetric Analysis in Correlation With Hormonal Receptor Subtypes and Histologic Grade of Invasive Breast Cancers. *AJR Am J Roentgenol* 204, W348–W356 (2015).

Turashvili, G. & Brogi, E. Tumor Heterogeneity in Breast Cancer. Front Med (Lausanne) 4, (2017).

Cuengod, C. A. & Balvay, D. Perfusion and vascular permeability: basic concepts and measurement in DCE-CT and DCE-MRI. *Diagn Interv Imaging* 94, 1187–1204 (2013).

Dvorak, H. F. Vascular permeability factor/vascular endothelial growth factor: a critical cytokine in tumor angiogenesis and a potential target for diagnosis and therapy. *J. Clin. Oncol.* 20, 4368–4380 (2002).

Szabó, B. K., Aspelin, P., Wiberg, M. K. & Boné, B. Dynamic MR imaging of the breast. Analysis of kinetic and morphologic diagnostic criteria. *Acta Radiol* 44, 379–386 (2003).

Vieira, A. F. & Schmitt, F. An Update on Breast Cancer Multigene Prognostic Tests—Emergent Clinical Biomarkers. Front Med (Lausanne) 5, (2018).

Li, H. et al. MR Imaging Radiomics Signatures for Predicting the Risk of Breast Cancer Recurrence as Given by Research Versions of MammaPrint, Oncotype DX, and PAM50 Gene Assays. *Radiology* 281, 382–391 (2016).

Dietzel, M. et al. The Necrosis Sign in Magnetic Resonance–Mammography: Diagnostic Accuracy in 1,084 Histologically Verified Breast Lesions. *The Breast Journal* 16, 603–608 (2010).

Pan, H. et al. 20-Year Risks of Breast-Cancer Recurrence after Stopping Endocrine Therapy at 5 Years. *N Engl J Med* 377, 1836–1846 (2017).

Grimm, L. J. et al. Interobserver Variability Between Breast Imagers Using the Fifth Edition of the BI-RADS MRI Lexicon. *American Journal of Roentgenology* 204, 1120–1124 (2015).

Saha, A., Yu, X., Sahoo, D. & Mazurowski, M. A. Effects of MRI scanner parameters on breast cancer radiomics. *Expert Syst Appl* 87, 384–391 (2017).

Cheng, Z. et al. Discrimination between benign and malignant breast lesions using volumetric quantitative dynamic contrast-enhanced MR imaging. *Eur Radiol* 28, 982–991 (2018).

Acknowledgements

The authors dedicate this work to the memory of Werner Alois Kaiser. His death marked the loss of an unprejudiced, creative, and critical scientist, a remarkable clinical radiologist, a generous, untiring mentor and a dear and dependable friend. Among many other research projects, he inspired us to study prognostic breast MRI long before this became a hot research topic and the buzz word “radiomics” was invented. The database created during this project is also the basis of the present study. Accordingly, some subjects of our study have been used for previous projects in a different clinical and/or scientific context (e.g. 12, 16).
Author contributions
Conceptualization: M.D., P.A.T.B. (lead) with contributions by all co-authors. Data curation and software: M.D., P.A.T.B., R.Z., I.B.R. Formal analysis M.D., P.A.T.B., S.E., R.Z. Investigation: M.D., P.A.T.B., R.Z., I.B.R. Methodology: M.D. & P.A.T.B. (lead) with contributions by all co-authors. Project administration: M.D., P.A.T.B., R.S.W., M.U. Resources: M.D., P.A.T.B., I.B.R., R.S.W., M.U. Software: M.D., P.A.T.B., R.Z. Supervision: M.D., P.A.T.B., R.Z. Validation: M.D., P.A.T.B. Visualization: M.D., P.A.T.B., Writing (original draft; review and editing): All authors.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-60393-9.

Correspondence and requests for materials should be addressed to P.A.T.B.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020