Supervised machine learning enables non-invasive lesion characterization in primary prostate cancer with \(^{68}\text{Ga}\) Ga-PSMA-11 PET/MRI

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Abstract

Purpose Risk classification of primary prostate cancer in clinical routine is mainly based on prostate-specific antigen (PSA) levels, Gleason scores from biopsy samples, and tumor-nodes-metastasis (TNM) staging. This study aimed to investigate the diagnostic performance of positron emission tomography/magnetic resonance imaging (PET/MRI) in vivo models for predicting low-vs-high lesion risk (LH) as well as biochemical recurrence (BCR) and overall patient risk (OPR) with machine learning.

Methods Fifty-two patients who underwent multi-parametric dual-tracer \(^{18}\text{F}\)FMC and \(^{68}\text{Ga}\)Ga-PSMA-11 PET/MRI as well as radical prostatectomy between 2014 and 2015 were included as part of a single-center pilot to a randomized prospective trial (NCT02659527). Radiomics in combination with ensemble machine learning was applied including the \(^{68}\text{Ga}\)Ga-PSMA-11 PET, the apparent diffusion coefficient, and the transverse relaxation time-weighted MRI scans of each patient to establish a low-vs-high risk lesion prediction model (MLH). Furthermore, M\(_{\text{BCR}}\) and M\(_{\text{OPR}}\) predictive model schemes were built by combining MLH, PSA, and clinical stage values of patients. Performance evaluation of the established models was performed with 1000-fold Monte Carlo (MC) cross-validation. Results were additionally compared to conventional \(^{68}\text{Ga}\)Ga-PSMA-11 standardized uptake value (SUV) analyses.

Results The area under the receiver operator characteristic curve (AUC) of the MLH model (0.86) was higher than the AUC of the \(^{68}\text{Ga}\)Ga-PSMA-11 SUV\(_{\text{max}}\) analysis (0.80). MC cross-validation revealed 89% and 91% accuracies with 0.90 and 0.94 AUCs for the M\(_{\text{BCR}}\) and M\(_{\text{OPR}}\) models respectively, while standard routine analysis based on PSA, biopsy Gleason score, and TNM staging resulted in 69% and 70% accuracies to predict BCR and OPR respectively.

Conclusion Our results demonstrate the potential to enhance risk classification in primary prostate cancer patients built on PET/MRI radiomics and machine learning without biopsy sampling.

Keywords Prostate cancer · PET/MRI · Radiomics · Machine learning · Lesion risk prediction · Biochemical recurrence prediction · Overall patient risk prediction

M. Hartenbach and M. Hacker shared senior authorship

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Introduction

Prostate cancer is the second most common cancer in men worldwide, with 1.3 million new cases diagnosed in 2018 [1, 2]. The worldwide incidence rates significantly increased during the last decade, most likely due to the wider application of prostate-specific antigen (PSA) screening [2]. While the 10-year survival rate of prostate cancer is approximately 90%, advanced or late-stage prostate cancer may be life-threatening, in particular, in metastasized stages of the disease [3].

The 5-year risk stratification in patients with primary prostate cancer is mainly built on clinical stage, PSA, and Gleason scores, derived from invasive biopsy samples [4]. Despite having profound effects on treatment planning and, thus, patient’s quality of life, this approach has a number of limitations [3, 5]. First, Gleason scoring relies on biopsy sampling, hence, can neither help assess the entire prostate nor fully characterize the heterogeneity of any pertinent tumor [6]. In addition, transrectal biopsy sampling has been associated with side-effects, such as haematospermia or haematuria [3]. Second, previously published risk classification systems were reported to have the tendency of incorrectly grading primary prostate cancer [3]. In patients with a high risk score and absent metastatic disease, radical prostatectomy is the treatment-of-choice [7] despite the risk of potential overtreatment [8] and at the same time, a 20–40% chance of biochemical recurrence (BCR) [9, 10].

Combined positron emission tomography/computed tomography (PET/CT) or PET/magnetic resonance imaging (PET/MRI) using radiotracers targeting prostate-specific membrane antigen (PSMA) can help to localize suspicious lesions in the prostate [11, 12]. PSMA-PET in combination with CT has been reported to improve primary tumor localization [13] and the diagnosis of recurrent prostate cancer [14, 15] in patients after radical prostatectomy even at low PSA levels [16]. In contrast, PSMA-PET/MRI was shown to support the diagnosis of intermediate and high-risk patients as well as to detect tumor recurrence [13]. Nevertheless, the diagnosis of primary prostate cancer is still based on core-needle biopsy, with non-invasive imaging playing a role in the visual identification of lesions and/or in image-guidance for biopsy sampling [17, 18].

Recently, radiomics have been argued to add value to the diagnostic pathways and patient management [19]. Various studies have been investigating the correlation of PSMA expression and clinical end-points in prostate cancer patients [14, 15]. Furthermore, radiomics combined with machine learning in MRI [20, 21] as well as in PET/CT [22–24] demonstrated the potential feasibility to establish novel in vivo prediction models for prostate cancer risk assessment.

In light of the potential of combining PET/MR imaging, radiomics and machine learning (ML), the objectives of this study were as follows: (a) to establish and cross-validate prostate lesion low-vs-high risk in vivo ML predictive models built on PET/MRI radiomics, (b) to establish and validate biochemical recurrence and overall patient risk (OPR) models that utilize in vivo ML scores instead of biopsy grades together with PSA and clinical stage, and (c) to compare the above patient risk models to the standard risk stratification.

Materials and methods

Patient data

Patients were selected from the database (n = 122) of a monocentric pilot study to a prospective randomized trial (clinicaltrials.gov NCT02659527) conducted between 2014 and 2015. Fifty-two of the 122 patients underwent surgery; in these patients, PET/MRI, PSA values, pre-operative biopsy results, and post-operative whole-mount histopathology were documented [15] (Table 1). All the 52 patients underwent a dual-tracer, fully integrated PET/MRI scan ([18F]FMC and [68Ga]Ga-PSMA-11 sequentially). This study, however, only included the [68Ga]Ga-PSMA-11 PET image as well as the transverse relaxation time-weighted (T2w) and apparent diffusion coefficient (ADC) MRI sequences in the analysis (Supplement: Table 1). All patients were treated with radical prostatectomy according to guideline recommendations [3]. All surgical specimens were processed according to the institution’s standard pathologic procedures in whole mount sections. Staging and grading were performed according to the UICC TNM classification and WHO/ISUP 2005 system, respectively [25]. The study was approved by the local institutional ethical committee and patients provided their written informed consent. See Fig. 1 for the CONSORT study diagram.

Delineation

Delineation and annotation of prostate lesions on PET/MR images were performed using the Hybrid 3D software ver. 4.0.0 (Hermes Medical Solutions, Stockholm, Sweden). Here, [68Ga]Ga-PSMA-11 PET and T2w as well as ADC MR images were viewed side-by-side with the annotated, whole-mount histopathological slices. Delineation was done over the [68Ga]Ga-PSMA-11 image using standard three-dimensional iso-count VOIs (Fig. 2). The initial lesion delineations were cross-examined and corrected manually—if required—as part of an independent review process performed by PET and MRI specialists. This step resulted in 121 lesions in total. An additional reference region was defined in the gluteus muscle to normalize the standard uptake value (SUV) of [68Ga]Ga-PSMA-11 and the T2w arbitrary voxel values to the mean of their respective reference background (26).
Feature extraction

Each image was resampled to $2.0 \times 2.0 \times 2.0$ uniform voxel resolution via ordinary Kriging interpolation [27, 28]. Radiomic features with “very strong” or “strong” consensus values as of the Imaging Biomarker Standardization Initiative (IBSI) guidelines were extracted from the 121 resampled $[^{68}\text{Ga}]{\text{Ga}}$-PSMA-11, T2w and ADC lesions by the MUW Radiomics Engine (ver. 2.0) that was validated based on IBSI standards [29] (Supplement Table 1). Conventional standardized uptake values including $\text{SUX}_{\text{max}}$, $\text{SUV}_{\text{peak}}$, $\text{SUV}_{\text{mean}}$, and $\text{SUV}_{\text{TLG}}$ were merged with the extracted 442 radiomic features to compose a 446 long feature vector for each lesion. While total lesion glycolysis (TLG) is originally proposed for $[^{18}\text{F}]\text{FDG}$, it was involved in our analysis as it characterized $[^{68}\text{Ga}]{\text{Ga}}$-PSMA-11 accumulation in prostate lesions.

Feature redundancy reduction

Feature redundancy ranking and reduction were done across the 446 features by covariance matrix analysis [19] where features were considered redundant with higher than 0.75 absolute Pearson correlation coefficient. This step resulted in keeping 80 features for further analysis.

Reference standard

The respective whole-mount histopathology patterns of each delineated lesion were dichotomized as low ($\leq$Gleason 3, prostatic intraepithelial neoplasia (PIN), prostatitis, benign prostatic hyperplasia (BPH)) and high ($\geq$Gleason 4) risk respectively. Furthermore, BCR and OPR reference values were established for each patient. BCR was defined when two consecutive PSA rose above 0.2 ng/ml. Follow-up was generally every 3 months for the first 2 years, then semiannually until the fifth year, then annually. Mean follow-up was 41 months. OPR was defined high, if BCR was positive or the node-stage (clinical or pathological) or the metastases-stage (clinical or pathological) were positive.

Statistical analysis in $[^{68}\text{Ga}]{\text{Ga}}$-PSMA-11

Area under the receiver operator characteristic curve (AUC) was calculated for conventional SUVs and the volume of each delineated lesion in the $[^{68}\text{Ga}]{\text{Ga}}$-PSMA-11 image to estimate the performance of predicting low-vs-high lesion risk. This process included $\text{SUX}_{\text{max}}$, $\text{SUV}_{\text{peak}}$, $\text{SUV}_{\text{TLG}}$, and lesion volume values.

Cross-validation scheme

Monte Carlo (MC) cross-validation scheme was utilized to randomly assign training and validation roles to the 52 patients 1000-times. In each fold, five patients were selected for the validation role, while the remaining patients got the training role. This step was necessary to avoid mixing lesions for training and validation from the same patient. No repetitions were allowed during the generation of MC folds; thus, each of the 1000-fold configurations with their training-validation selections was unique.

Machine learning scheme

Mixed ensemble learning scheme built on random forest classifiers (RF) was utilized to build models for predicting lesion LH, patient BCR as well as OPR (models denoted as $M_{\text{LH}}$, $M_{\text{BCR}}$ and $M_{\text{OPR}}$ respectively) [26, 30, 31]. Nine RFs
with various hyperparameters were configured for each of the three model schemes (Supplemental Table 2). The final prediction was provided by majority vote of the respective nine RFs. This approach was chosen to minimize hyperparameter bias and to increase predictive performance [32]. Furthermore, the average predictive score of the nine RFs represented a continuous value range between 0.0 and 1.0 reflecting on the prediction certainty of the mixed ensemble. Therefore, this value could be the subject of AUC analysis across MC folds.

Lesion low-vs-high risk prediction

Training and validation lesion sets were generated as of the pre-generated MC scheme roles to train and validate the MLH models in each MC fold. In order to keep model complexity minimal and to reduce the chance of overfitting, selection of the top five-ranking features was performed by R-squared ranking in the training dataset prior to establishing the MLH lesion model per fold [33]. The same five features were then selected from the respective validation

Fig. 1 The analysis workflow of the collected dataset. The pre-study of the prospective randomized trial NCT02659527 provided data records of 122 patients between 2014 and 2015. Patients having a dual-tracer positron emission tomography/magnetic resonance imaging (PET/MRI), prostate-specific antigen (PSA) screening, and whole-mount histopathology through undergone surgery were included in the analysis (n = 52). Only [68Ga]Ga-PSMA-11 PET, apparent diffusion coefficient (ADC), and transverse relaxation time-weighted (T2w) MRI images were selected for radiomic analysis. Overall 121 PET/MRI-positive lesions were delineated from the 52 patients followed by radiomics feature extraction. The 121 lesions underwent prostate specific membrane antigen (PSMA) standardized uptake value (SUV) and volume area under the receiver operator characteristics curve (AUC) analysis. Monte Carlo (MC) cross-validation scheme was utilized to generate patient training and validation sets 1000-times. This MC scheme was utilized to build lesion low-vs-high (LH) prediction models via machine learning (MLH). Biochemical recurrence (BCR, n = 36) and overall patient risk (OPR, n = 50) patient prediction models were built across the same MC folds (MBCR and MOPR respectively). All machine learning models underwent confusion matrix analytics, sham data analysis, and AUC analysis across MC folds. BCR and OPR were also predicted by standard D’Amico score.
dataset to evaluate. Validation model performance was estimated via confusion matrix analytics across the predictions of the validation cases of the MC folds [26]. The MLH scheme also underwent AUC analysis by evaluating the predictive performance of its averaged nine RF vote across the MC validation cases. Last, to estimate the effect of sham data in the MLH model, confusion matrix analytics were also performed over randomly permuted labels across all MC folds [24, 34].

**Feature weighting**

The importance of each feature in predicting lesion low-vs-high risk was determined by counting the occurrence of all selected features across the MC folds by the $R^2$-squared ranking approach.

**Patient biochemical recurrence and overall risk prediction**

Patient risk models for predicting BCR and OPR were established ($M_{BCR}$ and $M_{OPR}$ respectively) analyzing the PSA, the enumerated clinical stage (Supplemental Table 3), and a composite $M_{LH}$ score (CLH) per patient calculated by eq. 1.

$$CLH = \sum_{i=1}^{k} \frac{M_{LH}(i)v_i}{V}$$

where $k$ is the number of lesions in the given patient, $M_{LH}(i)$ is the predicted low-vs-high risk score of lesion $i$ provided by the $M_{LH}$ model of the given fold, $v_i$ is the volume of lesion $i$, and $V = \sum_{i=1}^{k} v_i$ is the sum of lesion volumes in the given patient.

Training and validation patient sets containing the above value triplets were generated as of the pre-generated MC scheme roles to train and validate the $M_{BCR}$ and $M_{OPR}$ models in each MC fold. In case a patient with validation role in the given fold had no BCR or OPR reference value available, it was excluded from the respective cross-validation of the given patient model.

To handle class imbalance, the training set underwent class imbalance correction by synthetic minority oversampling technique (SMOTE) [24, 35] for both the $M_{BCR}$ and $M_{OPR}$ training independently. Confusion matrix analytics were calculated across the validation set of all MC folds of the $M_{BCR}$ and $M_{OPR}$ model schemes. The same process was repeated by reference label permutations across the MC folds to estimate the effect of sham data. Both the

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**Fig. 2** (A) Positron emission tomography/magnetic resonance imaging (PET/MRI) views of a prostate cancer patient with volumes of interests (VOIs) drawn over lesions with Gleason 4 (red) and high-grade pin (blue) patterns. Standard iso-count 3D VOIs were drawn over the $[^{68}\text{Ga}]$Ga-PSMA-11 PET in the Hermes Hybrid 3D software. First row: $[^{68}\text{Ga}]$Ga-PSMA-11 PET; second row: apparent diffusion coefficient (ADC) MRI; third row: fused $[^{68}\text{Ga}]$Ga-PSMA-11 PET and transverse relaxation time-weighted (T2w) MRI images. Note that each image is represented in its own frame of reference, while the fused PET/MRI view is aligned to the frame of reference of the T2-weighted MRI. Hence, the cross-sections of the drawn VOIs look different on each view. (B) An example histopathological slice with the same color codes as in case of the PET/MRI views (red: Gleason 4, blue: high-grade pin)
M_{BCR} and M_{OPR} models underwent AUC analysis across the MC cross-validation folds.

### Results

#### Patients

Of the 52 patients, 36 had BCR during follow-up and 50 had OPR information available at the time of conducting the study. At the time of radical prostatectomy, the average PSA was 7.5. The most common pathologic stages were stage 2 ($n = 20$, 38%) followed by 3b ($n = 17$, 33%) and 3a ($n = 11$, 21%). Total Gleason score occurrences were GS $\geq 8$ ($n = 35$, 67%) followed by GS 7 ($n = 14$, 27%) and GS 6 ($n = 3$, 6%) (Table 1). The delineated 121 lesions represented a wide-range of benign and malign pathological alterations (Table 2). The most common high-risk pattern was associated to Gleason 4 ($n = 50$, 41%), followed by Gleason 3 ($n = 17$, 14%) and Gleason 5 ($n = 11$, 9%). Low-vs-high risk pattern regions were represented with balanced occurrences ($n = 61$-vs-60) (Table 2).

![ROC curves of lesion predictors to classify low-vs-high risk prostate lesions](image)

**Table 2** Characteristics of the 121 delineated lesions in the 52 patients

| Delineated lesions, $n$ (ratio)               | Value       |
|---------------------------------------------|-------------|
| Benign prostatic hyperplasia                | 20 (0.17)   |
| Low grade PIN                               | 16 (0.13)   |
| High grade PIN                              | 5 (0.04)    |
| Prostatitis                                  | 2 (0.02)    |
| Gleason 3                                    | 17 (0.14)   |
| Gleason 4                                    | 50 (0.41)   |
| Gleason 5                                    | 11 (0.09)   |
| Lesion high-low risk pattern, $n$ (ratio)   |             |
| High risk pattern                            | 61 (0.504)  |
| Low risk pattern                             | 60 (0.496)  |

ML_{BCR} and ML_{OPR} AUC performance is a conservative estimate, as it is a Monte Carlo cross-validation AUC, while the SUV and volume curves were measured directly from the whole dataset.
Statistical analysis in $[^{68}\text{Ga}]$Ga-PSMA-11

The AUC curves of SUV metrics were SUV$_{\text{max}}$ 0.80, SUV$_{\text{peak}}$ 0.74 and SUV$_{\text{TLG}}$ 0.64. Lesion volume presented AUC of 0.53. In contrast, the low-vs-high lesion prediction model (MLH) demonstrated a cross-validation AUC of 0.86 which was the highest compared to conventional $[^{68}\text{Ga}]$Ga-PSMA-11 values (Fig. 3).

Lesion low-vs-high risk prediction

The MLH model validation performance as per the MC cross-validation scheme yielded 71% sensitivity, 90% specificity, 88% positive predictive value, 75% negative predictive value, 81% accuracy, and 0.86 AUC. Sham data analysis revealed 0.52 AUC for permuted labels in the MLH model.

Feature weighting and distribution

Overall seven features were identified as selected across the 1000 MC folds via the $R$-squared ranking method. Features that were always selected were coefficient of variation and gray level co-occurrence matrix (GLCM) information correlation type 1; ADC—apparent diffusion coefficient; stat.iqr—interquartile range; cm.joint.entr—gray level co-occurrence matrix joint entropy; dzm.hgze—gray level distance zone matrix high gray zone emphasis.
respectively in the $[^{68}\text{Ga}]$Ga-PSMA-11 image. The lowest ranking feature ($n = 58$) was high gray zone emphasis in the $[^{68}\text{Ga}]$Ga-PSMA-11 image (Fig. 4).

**Patient biochemical recurrence and overall risk prediction**

The cross-validation performance revealed an average validation accuracy of 89% and 91% as well as AUC of 0.90 and 0.94 for the $M_{\text{MCR}}$ and $M_{\text{OPR}}$ patient models respectively. The $M_{\text{OPR}}$ model outperformed the $M_{\text{MCR}}$ model with 94% specificity, 93% positive predictive value, and with 87% sensitivity. The performance of $M_{\text{OPR}}$ and $M_{\text{MCR}}$ with sham data revealed 0.54 and 0.56 AUC respectively. See Fig. 5 for the detailed performance values of the $M_{\text{MCR}}$ and $M_{\text{OPR}}$ models.

**Discussion**

In this study, we investigated the feasibility of predicting prostate lesion-specific low-vs-high risk built on PET/MRI radiomics and patient-specific biochemical recurrence as well as overall patient risk. We demonstrated excellent cross-validation performances for $M_{\text{MLH}}$ (AUC 0.86) as well as for $M_{\text{MCR}}$ (AUC 0.90) and $M_{\text{OPR}}$ (AUC 0.94). Based on the above approaches and our achieved model performances, we consider that our findings have important clinical implications in the field of primary prostate cancer risk assessment as they point towards the feasibility to estimate lesion and patient risks in vivo.

Next to establishing the above models with radiomics and machine learning, conventional $[^{68}\text{Ga}]$Ga-PSMA-11 SUV and volume analysis were also conducted. This analysis revealed that SUV$_{\text{max}}$ had the highest predictive power (AUC 0.80) to classify low-vs-high prostate lesions followed by SUV$_{\text{peak}}$ and SUV$_{\text{TLDG}}$, while lesion volume had no significant predictive power (AUC 0.53). These findings are in line with previous analyses performed in PET/CT [24].

Feature ranking across our Monte Carlo folds demonstrated that $[^{68}\text{Ga}]$Ga-PSMA-11 is the most important in vivo feature source to establish lesion risk prediction models compared to ADC and T2w MRI features. The highest-ranking $[^{68}\text{Ga}]$Ga-PSMA-11 features were either simple statistical values such as the coefficient of variation and SUV$_{\text{max}}$ or simple second-order textural ones such as information correlation from the GLCM feature category. Information correlation is a first-order GLCM feature reflecting on the information content (a.k.a. entropy) of voxel neighborhood connectivity occurrences; thus, it is a basic heterogeneity descriptor. This feature was previously also identified as highly robust across various PET imaging centers [36]. The feature ranking across MC folds identified SUV$_{\text{peak}}$, SUV$_{\text{TLDG}}$, and volume as low-ranking; however, SUV$_{\text{max}}$ was among the highest ranking ones. While the potential of PSMA SUV$_{\text{max}}$ in characterizing prostate cancer had been presented [37, 38], Cysouw et al. concluded in a recent study that prostate risk in PSMA can be better characterized by textural parameters compared to SUV$_{\text{max}}$ [24]. They utilized $[^{18}\text{F}]$-DCFPyL PET/CT and reported 0.81 AUC to differentiate high (GS $\geq 8$) and low-risk prostate cases. Our findings on the other hand demonstrate that conventional SUV parameters in combination with simple textural features can yield high-performing models in $[^{68}\text{Ga}]$Ga-PSMA-11 PET/MRI to characterize prostate risk.

While no T2w feature was selected as high-ranking, ADC interquartile range (also referred to as “robust” value range) was selected as high-ranking. Prior studies focusing on ADC analysis to predict prostate lesion risk consistently identified ADC$_{\text{min}}$, ADC$_{\text{mean}}$ as well as ADC$_{\text{median}}$ [20, 39] as highly predictive (AUC range 0.72–0.90). We consider that the above findings and ours describe the same phenomenon, namely, the strong predictive ability of simple ADC values without the need of incorporating second or higher-order radiomic features in the analysis. The above findings in prior reports demonstrate the predictive performance of PSMA PET and ADC MR images individually. Hence, we hypothesize that the high performance of our $M_{\text{MLH}}$ model is due to the fact that it combines both $[^{68}\text{Ga}]$Ga-PSMA-11 PET and ADC MRI features in one model scheme.

Further to the above findings, we also established patient biochemical recurrence ($M_{\text{MCR}}$) and overall patient risk ($M_{\text{OPR}}$) models. In order to provide an in vivo score per patient in lieu of biopsy grades in these models, we created a CLH score which weighted each $M_{\text{MLH}}$ score per lesion with its respective volume in each patient. Since volume was identified as non-predictive to classify low-vs-high risk in prostate lesions (AUC 0.53), we assumed that the volume effect [40] in our high-ranking features was negligible, and thus, lesion volume was an independent value from our lesion $M_{\text{MLH}}$ scores. This assumption allowed us to utilize volume as a weight factor for each lesion $M_{\text{MLH}}$ score to compose the patientspecific CLH score. The resulted CLH score in combination with PSA and clinical stage values resulted in high-performing $M_{\text{MCR}}$ and $M_{\text{OPR}}$ models (0.90 and 0.94 cross-validation AUCs respectively). We assume that the accuracy performance increase of +20% and +21% in our $M_{\text{MCR}}$ and $M_{\text{OPR}}$ models compared to standard risk estimation are due to the following reasons: first, the clinical standard utilizes Gleason patterns from biopsy to describe lesion pattern risks. In contrast, we incorporated PSA, clinical stage, and the CLH score without re-binning them, and thus, avoiding...
potential information loss. Third, the clinical standard score acts as a maximum filter across its pre-binned risk categories to estimate overall risk to the patient. In contrast, the random forest ensemble logics in our MBCR and MOPR models could describe more complex relationships among PSA, clinical stage, and our in vivo CLH score. Our results demonstrate that such relationships may be indeed present and that building on those relationships may lead to in vivo risk predictive models in prostate cancer patients with the potential to eliminate the need of biopsy sampling in the future.

This study had a number of limitations. First, it built on a single-center cohort; however, due to utilizing a pre-generated MC fold scheme for all training and validation processes, no training and validation samples were mixed in between the lesion and patient predictors. In addition, the utilized data preparation (redundancy reduction, feature ranking, and class imbalance correction) as well as training (mixed ensemble) and validation (1000-fold CV, sham data analysis) approaches minimized the chances of false discoveries. Second, due to the dual-tracer study design from which our images were taken, the $^{68}$Ga-Ga-PSMA-11 scans were not entirely exempt of $^{18}$F]FMC uptake remnants. Nevertheless, $^{18}$F]FMC can be regarded an irreversible tracer [43] and, thus, the $^{18}$F]FMC uptake in terms of tissue to lesion ratio is expected not to change until the $^{68}$Ga-Ga-PSMA-11 examination. Last, only patients with proven prostate cancer were included after radical prostatectomy. Nevertheless, this selection criterion was necessary to acquire stable ground truth for lesion labeling.

**Conclusions**

This study demonstrates the feasibility of $^{68}$Ga-Ga-PSMA-11 PET/MRI in combination with radiomics and machine learning to non-invasively deliver both lesion characterization and risk prediction equally to preoperative invasive biopsy in patients with primary prostate cancer. Prospective multicentric studies are required to investigate the reproducibility and clinical utility of this approach.

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**Authors’ contributions** All authors contributed to writing, critically reviewing, and approving the paper. Specific author contributions are as follows:

| Author | Contribution |
|--------|--------------|
| L. Papp | Radiomics and machine learning methodological design, study concept and execution, literature research |
| C. P. Spielvogel | Random forest implementation, validation, and parameter tuning |
| B. Grubmüller | Biochemical recurrence reference standard collection, study design, review |
| M. Grahovac | High-performance computing execution, literature research |
| D. Krajnc | Radiomic feature purification, class imbalance handling, literature research |
| B. Ecsedi | Implementation and validation of the MUW Radiomics Engine as of IBSI guidelines and reference datasets |
| R. A. M. Sareshgi, M. Grahovac, D. Mohamad, M. Hamboeck | Lesion delineation |
| I. Rausch | PET/MR protocol set up, $^{18}$F]FMC cross-effect analysis and estimation in $^{68}$Ga-Ga-PSMA-11, literature research |
| M. Mitterhauser | Study design review and approval, review |
| W. Wadsak | Supervision and set-up of radiotracer synthesis and quality control |
| A. Haug | Study concept, review |
| L. Kenner | Histopathological analysis and labelling |
| P. Mazal | Histopathological analysis and labelling |
| M. Susani | Histopathological analysis and labelling |
| S. Hartenbach | Whole-mount histopathological analysis and labelling |
| P. Balzer | MRI acquisition protocol setup |
| T. Helbich | MRI acquisition protocol setup, study concept |
| G. Kramer | Generating and processing clinical data including follow-up |
| S. F. Shariat | Generating and processing clinical data including follow-up |
| T. Beyer | Study design review and approval |
| M. Hartenbach | PET/MR acquisition protocol set up, initial data collection and preparation, delineation review and validation, study concept, study PI, review |
| M. Hacker | Study concept, study PI, review |

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**Data availability** The datasets generated and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.
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