MRI utility in predicting extraprostatic extension of prostate cancer and biochemical recurrence after radical prostatectomy

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Introduction. The study aimed to evaluate the performance of multiparametric magnetic resonance imaging (mpMRI) and the Martini model to predict extraprostatic extension (EPE) and biochemical recurrence (BCR) of prostate cancer (PCa).

Materials and methods. 61 patients underwent a radical laparoscopic prostatectomy. The preoperative risk of EPE was determined using mpMRI and the Martini model.

Results. MpMRI predicts the presence of EPE of PCa with a sensitivity and specificity of 47.4% and 85.7%, respectively (AUC 0.66, 95% CI: 0.51–0.82, p = 0.046). The Martini model’s sensitivity was higher, but the specificity was lower than that of mpMRI and was 84.2% and 66.7%, respectively (AUC 0.78, 95% CI: 0.66–0.89, p < 0.001). Univariate and multivariate Cox analysis indicated that EPE in mpMRI (HR 6.6, 95% CI: 1.8–24.1), and the presence of positive surgical margins (PSM) (HR 7.1, 95% CI: 1.9–26.7) are independent factors increasing the probability of BCR.

Conclusions. MpMRI and Martini model are valuable tools in local staging of PCa, managing and predicting the oncological treatment outcomes of patients with PCa.

Key words: prostate cancer, multiparametric magnetic resonance imaging, radical prostatectomy, biochemical recurrence, extraprostatic extension

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According to the current standards, the local staging is based on clinical examination (digital rectal exam – DRE) or a transrectal ultrasound (TRUS) [2]. Apart from the clinical examination and TRUS, local staging can also be performed based on mpMRI. MpMRI provides valuable information about the clinical significance and localization of the lesion [6]. This information is used to qualify the patient for a prostate biopsy. According to the current guidelines, it is recommended to perform mpMRI in suspected PCa before the first prostate biopsy [2]. Moreover, mpMRI, in the case of PCa, allows for a characterization of such clinical parameters as lesion size, possible EPE or seminal vesicle invasion (SVI) [6]. The above information allows for an estimation of the appropriate risk group of disease progression, choose the treatment method (RP vs. RT), or plan the extension of the RP. Nevertheless, mpMRI is still not validated as a diagnostic tool for local staging and treatment planning [2, 7, 8].

The study aimed to evaluate the utility of mpMRI in the prediction of EPE and the impact of EPE on the occurrence of BCR in patients qualified for RP.

Material and methods

The study group consisted of 61 patients diagnosed with PCa who underwent laparoscopic RP. RP was performed either by the trans- or retropertioneal approach. An extended lymphadenectomy was performed in the case of high-risk and intermediate-risk cancer with a predicted probability of lymph node involvement above 7%, according to the Briganti 2017 model [9]. The operation was performed with bilateral or unilateral NVB preservation or without NVB preservation. The NVB preserving technique involved inter- or intra-fascial dissection of the bundles, according to Walz [10]. The procedures without NVB preservation involved a wide extra-fascial prostatectomy [10]. The decision to preserve NVB was made depending on the EAU risk group. The preoperative risk of EPE was determined according to the Martini et al. model [11]. The decision to preserve NVB was also influenced by comorbidities, erectile dysfunction present before the planned procedure, age, and the patient’s preferences.

The biopsy material and specimen acquired during RP underwent histopathological assessment conducted by three pathologists in accordance with the guidelines of the International Society of Urological Pathology (ISUP) 2014 in the field of pathomorphological diagnosis of PCa [12].

The study used mpMRI obtained when qualifying the patients for the first biopsy due to suspected PCa. 1.5T (GE Healthcare Medical System Optima MR360, Chicago, IL, USA) and 3T equipment (Siemens HealthCare Magnetom Skyra, Erlangen, Germany) were utilized using 12- or 18-channel Body Matrix coils. The mpMRI scheme followed the PI-RADS v. 2.0 guidelines of the American College of Radiology (ACR) [6]. It included a multiplanar assessment of the prostate in T1- and T2-weighted images, diffusion weighted imaging (DWI) and a dynamic contrast-enhanced (DCE) MRI. Apparent diffusion coefficient (ADC) maps were developed automatically. The mpMRI was evaluated by four radiologists experienced in prostate imaging who knew the PSA levels and rectal examination (DRE).

Based on mpMRI, a targeted cognitive biopsy of the prostate combined with a systematic biopsy was performed, guided with transrectal ultrasound using a biplane transducer with simultaneous imaging of both planes (BK Medical Flex 400, Herlev, Denmark).

The biopsy was performed according to the scheme recommended by the European Society of Urology (EAU) [2]. 6–8 specimens were collected from each lobe, plus additionally 2–4 specimens from the suspicious lesion depending on its size [2, 13, 14]. The biopsy was performed by four urologists. BCR was diagnosed when two PSA levels above 0.2 ng/ml were obtained. PSA was monitored every three months during the first year and every six months in consecutive years.

Statistical analysis

The analyzed parameters were described using an arithmetic mean, standard deviation, and median. The normality of distribution was verified using the Shapiro-Wilk test in each of the analyzed groups. In a normal distribution, the t-Student test for independent variables was used to compare the two groups. In the case of non-normal distribution, the non-parametric Mann-Whitney test was applied. Categorical variables in individual groups were described using percentage values; they were compared using the Chi2 test with the Yates correction and with Fisher’s exact test. Receiver operating characteristic (ROC) curves were determined for the analyzed parameters (Martini model and mpMRI). AUC (area under curve) was subsequently calculated, and their significance was analyzed. Cut-off points for which sensitivity and specificity reach optimal values were determined for parameters with significant AUC (Youden point).

Kaplan-Meier analysis and the log-rank test were used to compare BCR-free survival for patients with EPE in mpMRI. For BCR risk factors, hazard ratios (HRs) with 95% confidence intervals (CIs) were determined using the Cox model. The univariate and multivariate model was established. The assumed p-value was <0.05. The IBM SPSS Statistics statistical package was used for the calculations.

Results

Table I presents the clinical characteristics of 61 patients undergoing RP (n = 61). 19 patients (n = 19) were diagnosed with EPE(+) and 42 (n = 42) without EPE (–) in the final histopathological report post RP. Patients with EPE(+) differed statistically significantly from patients without EPE(–), in terms of the following clinical parameters: ISUP grade in the preoperative biopsy, maximum index lesion (IL) dimension in mpMRI, EPE diagnosis in mpMRI, ISUP Grade in the postoperative report, the incidence of PSM, and the incidence of BCR in follow-up.
Analysis using the ROC curve showed that preoperative mpMRI might predict the presence of EPE of PCa with a sensitivity and specificity of 47.4% and 85.7%, respectively (AUC 0.66, 95% CI: 0.51–0.82, p = 0.046). Taking into account the results of the first statistical analysis, which indicate that EPE may also be dependent on other clinical parameters, we examined the sensitivity and specificity of the Martini model, which uses the following data: PSA level, EPE status in mpMRI, ISUP Gleason grade, and the percentage of the biopsy specimen involvement. The Martini model’s sensitivity was higher, but the specificity was lower than that of mpMRI and was 84.2% and 66.7%, respectively (AUC 0.78, 95% CI: 0.66–0.89, p < 0.001) (fig. 1).

All patients in the study were subject to follow-up. The mean follow-up was 38 months (95% CI: 34.0–42.5 months). Patients with pEPE(+) more frequently experienced BCR 42% (n = 8) vs. pEPE(−) 7% (n = 3) (HR 6.4, 95% CI: 1.7–24). Using univariate and multivariate Cox analysis, it was examined whether other clinical factors may also influence the occurrence of BCR. Previous prognostic factors influencing the patient’s prognosis (PSA, ISUP grade, DRE, prostate volume, SM) were analyzed, taking into account mpMRI (PIRADS of IL, max diameter of IL in mpMRI (mm), mpMRI zone location: PZ, non-PZ). The final Cox model showed that EPE in mpMRI was an independent factor that increased the likelihood of BCR – HR 6.6 (95% CI: 1.8–24.1).

### Table 1. Clinical characteristics of patients. Comparison of clinical data between pEPE(−) and pEPE(+) patients

| Total n (%) or median | Overall (n = 61) | pEPE(−) (n = 42) | pEPE(+) (n = 19) | pEPE(+) vs. pEPE(−) p value |
|-----------------------|------------------|-----------------|-----------------|---------------------------|
| age (median)          | 65               | 65.7            | 63.4            | NS                        |
| PSA (ng/ml)           | 8.46             | 8.01            | 9.47            | NS                        |
| PSA (ng/ml/ml)        | 0.24             | 0.22            | 0.29            | NS                        |
| DRE:                  |                  |                 |                 | NS                        |
| • normal              | 22 (36.1)        | 18 (42.9)       | 4 (21)          | NS                        |
| • abnormal            | 39 (63.9)        | 24 (57.1)       | 15 (79)         |                           |
| biopsy ISUP grade:    |                  |                 |                 | *                         |
| • 1                   | 35 (57.4)        | 29 (69)         | 6 (31.6)        |                           |
| • >1                  | 26 (42.6)        | 13 (31)         | 13 (68.4)       |                           |
| prostate volume (ml) (median) | 38.9 | 39.32 | 37.86 | NS |
| PIRADS:               |                  |                 |                 | NS                        |
| • 1–3                 | 21 (34.4)        | 15 (35.7)       | 6 (31.6)        |                           |
| • 4–5                 | 40 (65.6)        | 27 (64.3)       | 13 (68.4)       |                           |
| max diameter of IL in mpMRI (mm) | 13.8 | 12.26 | 17.32 | *                       |
| mpMRI EPE             |                  |                 |                 | *                         |
| • EPE(−)              | 46 (75.4)        | 36 (85.7)       | 10 (52.6)       |                           |
| • EPE(+)              | 15 (24.6)        | 6 (14.3)        | 9 (47.4)        |                           |
| mpMRI zone location:  |                  |                 |                 | NS                        |
| • PZ                  | 45 (73.8)        | 29 (69)         | 16 (84.2)       |                           |
| • non-PZ              | 16 (26.2)        | 13 (31)         | 3 (15.8)        |                           |
| BCR(+)                | 11 (18)          | 3 (7.1)         | 8 (42.1)        | *                         |
| pathologic ISUP grade:|                  |                 |                 | *                         |
| • 1                   | 25 (41)          | 22 (52.4)       | 3 (15.8)        |                           |
| • >1                  | 36 (59)          | 20 (47.6)       | 16 (84.2)       |                           |
| PSM                   | 18 (29.5)        | 8 (19)          | 10 (52.6)       | *                         |

PSA – prostate specific antigen; PSAD – PSA density; DRE – digital rectal exam; ISUP grade – 2014 International Society of Urological Pathology Grade; PIRADS – Prostate Imaging Reporting and Data System; mpMRI – multiparametric magnetic resonance imaging; EPE – extraprostatic extension (“−” absent, “+” present); pEPE – pathological extraprostatic extension (“−” absent, “+” present); PZ – peripheral zone; non-PZ – zone other than peripheral; BCR – biochemical recurrence; PSM – positive surgical margin; IL – index lesion; NS – not significant; * – p < 0.05

Figure 1. ROC curve – sensitivity and specificity of the Martini nomogram for EPE prediction before RP
Another important factor, also significantly increasing the risk of BCR, was the presence of PSM in the postoperative report – HR 7.1 (95% CI: 1.9–26.7). Based on the above analysis, the Kaplan-Meier curve was determined for EPE assessment in mpMRI (fig. 2).

Discussion

The results obtained indicate that preoperative mpMRI may be a useful tool in local staging of PCA and for the prediction of recurrence; they are to a large extent consistent with the results of other similar studies [7, 8, 15, 16]. The mpMRI parameters analyzed by other researchers included: PIRADS score, volume and location of index lesions, presence of EPE, and/or SVI. Moreover, taking into account mpMRI parameters in conjunction with other clinical data, such as PSA levels and prostate biopsy reports, may contribute to a significant improvement in local staging. An example of this strategy is the model according to Martini et al., which takes into account mpMRI, PSA levels, and a report of the targeted biopsy based on MRI [11].

This retrospective study is one of the few studies available in the literature, which indicates that mpMRI and the Martini model may be tools that are helpful in determining the presence of EPE before surgical treatment [8, 11]. The sensitivity and specificity of mpMRI in assessing EPE in our study were 47.4% and 85.7%, respectively, while for the Martini model – 84.2% and 66.7%, respectively. The meta-analysis conducted by de Rooij, assessing the diagnostic utility of mpMRI in the prediction of stage T3a (EPE), indicated a sensitivity of 0.57 (95% CI: 0.49–0.64) and a specificity of 0.91 (95% CI: 0.88–0.93) [8]. When creating this predictive model for the presence of EPE, Martini estimated the sensitivity of mpMRI alone at 40.7%. [11].

Another important factor, also significantly increasing the risk of BCR, was the presence of PSM in the postoperative report – HR 7.1 (95% CI: 1.9–26.7). Based on the above analysis, the Kaplan-Meier curve was determined for EPE assessment in mpMRI (fig. 2).

Table II. Univariate and multivariate Cox regression analysis of clinical factors for BCR in patients with localized PCA after RP

| Univariate analysis | Multivariate analysis |
|---------------------|----------------------|
|                      | HR (95% CI)          | p value | HR (95% CI) | p value |
| DRE, ref. normal     | 0.96 (0.28–3.3)      | 0.94    |              |         |
| PSA                 | 0.95 (0.81-1.1)      | 0.4     |              |         |
| biopsy ISUP grade, ref. grade 1 | 1.76 (0.56–5.8) | 0.35    |              |         |
| prostate volume     | 0.97 (0.92–1.0)      | 0.33    |              |         |
| max diameter of IL in mpMRI (mm) | 1.04 (0.97–1.1) | 0.26    |              |         |
| PIRADS, ref. ≤3    | 1.025 (0.3–3.6)      | 0.97    |              |         |
| mpMRI – EPE, ref. EPE(–) | 3.9 (1.2–12.9) | 0.02    | 6.6 (1.8–24.1) | 0.005 |
| SM, ref. negative   | 4.7 (1.4–16.1)       | 0.01    | 7.1 (1.9–26.7) | 0.004 |

DRE – digital rectal exam; PSA – prostate specific antigen; ISUP grade – 2014 International Society of Urological Pathology Grade; PIRADS – Prostate Imaging Reporting and Data System; mpMRI – multiparametric magnetic resonance imaging; EPE – extraprostatic extension; BCR – biochemical recurrence; SM – surgical margins; IL – index lesion; NA – not applicable in final model; CI – confidence interval; ref – reference.

Figure 2. Kaplan-Meier survival curves for biochemical-free survival in patients with positive (+) and negative (–) mpMRI EPE.

Our study also confirmed the relationship between the presence of EPE in the postoperative histopathological report and an increased risk of BCR. Moreover, we demonstrated that EPE in preoperative mpMRI might also be an essential pre-surgical factor increasing the risk of BCR after RP. Therefore, the results obtained in our study indicate that mpMRI may improve the prediction of possible disease recurrence by improving local disease staging. In a study with a similar methodology, conducted on a large group of respondents (n = 804), Gandalgia et al. proved that preoperative information obtained from mpMRI and the report of systematic biopsy combined with a targeted biopsy based on mpMRI allows stratification of PCA recurrence after RP [15]. When assessing the influence of preoperative factors on BCR, Manceau et al. also proved that patients with EPE(+) diagnosed in mpMRI at the treatment planning stage more frequently experience BCR after RP [16]. Moreover, they showed a correlation between the occurrence of BCR and higher PIRADS scores, the greater
maximum dimension of MRI lesions, and a higher ISUP Gleason grade in fusion biopsy [16].

From a clinical perspective, the presence of EPE in mpMRI may be a predictive factor for the risk of adjuvant RT implementation after RP. This knowledge at the treatment planning stage can help in the choice of a treatment method, i.e., RP vs. RT. If surgical treatment is chosen, the information about the presence of EPE in mpMRI and its location can be used to plan the procedure’s technique and make decisions, for example, regarding the preservation of nerve bundles. It has been proven that planning RP based on mpMRI changes the operator’s decision to preserve NVB in 35% (95% CI: 29–41%) of cases; this strategy is correct in 77% (95% CI: 72–81%) of cases and does not worsen oncological outcomes [17].

Our study has several limitations. The first significant limitation is that it was conducted on a small group studied in one center. The second limitation is the lack of assessment of the extent of EPE and its detailed location, which, according to available knowledge, may also be a significant factor influencing the risk of BCR.

Conclusions

MpmRI and the Martini model are helpful tools in the local staging of patients with PCa. Preoperative use of mpMRI can predict oncological treatment outcomes in patients with PCa after RP.

Conflict of interest: none declared

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