Extracapsular extension on multiparametric magnetic resonance imaging better predicts pT3 disease at radical prostatectomy compared to perineural invasion on biopsy

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Abstract

Introduction: Risk assessment for non-organ-confined prostate cancer (PCa) is important in the surgical planning for radical prostatectomy (RP). Perineural invasion (PNI) on prostate biopsy has been associated with adverse pathological outcomes at prostatectomy. Similarly, the identification of suspected extracapsular extension (ECE) on multiparametric magnetic resonance imaging (mpMRI) has been shown to predict non-organ-confined disease. However, no prior study has compared these factors in predicting adverse pathology at prostatectomy. We evaluated mpMRI ECE and prostate biopsy PNI on multivariable analysis to determine their ability to predict pathological stage at time of RP.

Methods: We retrospectively investigated the prostatectomy database at our institution to identify men who underwent prostate biopsy with pre-biopsy mpMRI and subsequent RP from 2013–2017. Multivariable regression analysis was performed to compare the association of mpMRI ECE (mECE) and PNI on prostate biopsy on the likelihood of finding pT3 disease on pathology post-prostatectomy.

Results: Of a total 454 RP between 2013 and 2017, 191 patients met our inclusion criteria. Stage pT2 and pT3+ were found in 120 (62.8%) and 71 (37.2%) patients, respectively. Patients with mECE had 4.84 cumulative odds of worse pathological stage on RP (p=0.045) compared to PNI on biopsy, which showed cumulative odds of 2.25 (p=0.048). When controlling only for those patients without PNI, mECE was still found to be a significant predictor of pT3 disease at RP (p=0.030); however, in patients without mECE, PNI was not significant (p=0.062).

Conclusions: While mECE and biopsy PNI were both associated with worse pathological stage on RP, mECE had significantly higher cumulative odds compared to PNI. The significant predictive ability of mECE adds further clinical value to the use of mpMRI in PCa management. While validation in a larger cohort is required, these factors have important clinical implications with regards to early diagnosis of advanced disease and surgical planning.

Introduction

In patients with localized prostate cancer (PCa), nerve-sparing surgery offers improved postoperative functional outcomes and is considered the standard approach to radical prostatectomy (RP).1,2 Implementing a nerve-sparing approach, however, is not appropriate for every case and the decision to nerve spare is influenced by the estimated probability of extraprostatic extension (EPE) on prostatectomy.3,4 To optimize surgical and treatment planning, risk assessment for features of non-organ-confined disease is performed through nomogram assessment of preoperative clinical features, such as preoperative prostate-specific biopsy (PSA), biopsy Gleason score, and clinical stage.5,6 However, preoperative estimates are often inaccurate, and in patients where the tumor extends outside of the prostate, the risk of positive surgical margin and adverse pathology may be particularly high.7,8

In an effort to improve existing predictive risk models, additional preoperative parameters have been studied for their utility. Perineural invasion (PNI) is defined as the tracking of tumor cells along or around nerve fibers; in PCa, this extension is commonly reported on prostate biopsy pathology.9,10 While not traditionally used in nomogram assessment, PNI on prostate biopsy (PBx) is associated with EPE at RP, as well as margin positivity and biochemical recurrence.11-16

Multiparametric magnetic resonance imaging (mpMRI) is a PCa diagnostic tool that is increasingly popular...
and is being used to direct PBx and surgical planning. Preoperative mpMRI is useful for assessing the presence of significant cancer, predicting organ-confined disease, and assessing seminal vesicle invasion. While having a low to moderate sensitivity, mpMRI has achieved high specificity and positive predictive values of up to 90% in detecting extracapsular extension (ECE) and seminal vesicle invasion. Interestingly, while both PNI on PBx and mpMRI extracapsular extension (mECE) have been shown as independent predictors of adverse outcomes and non-organ-confined disease, no prior study has compared these factors. To query whether PNI is a significant prognostic variable in the era of mpMRI, we evaluated and compared the ability of mECE and PNI on PBx to predict advanced pathological stage at time of RP.

**Methods**

**Patient population**

We retrospectively investigated the prostatectomy database at our institution to identify men who underwent RP from July 2013 to March 2017. Patients were included if they received an mpMRI and subsequent PBx before RP at our institution. Patients were excluded if they had missing information, incomplete mpMRI, if biopsy or mpMRI were conducted at an outside institution, or if they had a history of alternative or previous treatment for PCa, such as radiation therapy, cryoablation, or any neoadjuvant therapy. A subgroup analysis was performed to ascertain if mECE or PNI would reach significance in prediction of pT3 when excluding all mECE patients and then all PNI patients from the original cohort. All staging was assigned according to the 2017 American Joint Committee on Cancer (AJCC) staging criteria.

**Protocol for mpMRI**

Patients were referred for mpMRI in the setting of elevated PSA or PSA density. Urologists at our institution routinely obtained mpMRI prior to biopsy to use a fusion approach and for potential surgical planning. All patients underwent mpMRI (3T Verio®, Siemens, Germany) of the prostate obtained with a 16-channel cardiac coil (SENSE®, Invivo, Gainesville, FL, U.S.) on the anterior pelvis and an endorectal coil (BPX-30®, Medrad, Pittsburgh, PA, U.S.) with its balloon filled with PFC-770 (3M, St. Paul, MN, U.S.). Sequences obtained included tri-planar T2-weighted, diffusion-weighted imaging (DWI) (b-values 0, 500, 1000, and 1500) and separate b-2000 and dynamic contrast enhanced (DCE) sequences. All mpMRI were read by experienced genitourinary radiologists, who assigned risk scores using the Prostate Imaging Reporting & Data System (PI-RADS) v1 and PI-RADS v2 scoring system. If multiple suspicious lesions were present, the highest PI-RADS score reported was used. ECE on MRI was stratified based on radiology report into mECE present, meECE suspicious, or meECE absent. meECE ‘present’ was assigned if ECE was clearly seen, as described by the radiologist, with gross extension of disease. meECE ‘suspicious’ was assigned to reports including descriptions of “abutment,” capsular “bulging,” or “microscopic invasion cannot be ruled out.” meECE ‘absent’ was assigned when there was clearly no evidence of ECE on MRI.

**Protocol for prostate biopsy**

All biopsies were performed via transrectal technique by experienced urologists at our institution. MRI-targeted images were processed on a DynaCAD® workstation (Invivo, Gainesville, FL, U.S.). MR/transrectal ultrasound (TRUS) image fusion was performed using UroNav® software in conjunction with an IU-22 (Philips Health Care, Best, Netherlands) end-fire ultrasound probe. During biopsies of the lesions, one core was obtained in the axial and sagittal planes for a total of two cores per lesion, followed by 12-core systematic biopsy as per our institution’s protocol for MRI-fusion biopsy. Patients without a targetable lesion identified on MRI received a standard 12-core TRUS biopsy whereby two cores — medial and lateral — are taken from each sextant region in a standard fashion. All biopsy pathology slides were reviewed by our institution’s experienced genitourinary pathologist.

**Statistical methods**

Univariable analysis was performed to determine if there were differences in age, PSA at MRI, prostate volume, PI-RADS score, and ECE between pT2 and pT3+. A Mann-Whitney U test was run for non-normally distributed continuous variables and Chi-squared was run for nominal parameters. A cumulative odds ordinal logistic regression with proportional odds was run to determine the effect of six variables on pathological cancer staging after RP. These variables included age, PNI on PBx, mECE, PI-RADS, PSA, and MRI prostate volume. A binomial logistic regression analysis was performed on all the variables to ascertain the likelihood of subjects having positive margins. PI-RADS, PSA, age, and prostate volume were analyzed as continuous variables. Proportional odds were assessed by a full likelihood ratio test comparing the fitted model to a model with varying location parameters. The model fit was tested through a deviance goodness-of-fit test. Collinearity of the independent variables was assessed and ruled out through inspection of correlation coefficients and tolerance/VIF values. SPSS 24.0.0 for Windows (SPSS Inc., Chicago, IL, U.S.)
was used for all statistical analysis, with $p<0.05$ considered statistically significant.

**Results**

During the study period, 454 patients underwent RP at our institution; 191 patients met our inclusion criteria. Baseline characteristics are shown in Table 1. Stage pT2 and pT3 were found in 120 (62.8%) and 71 (37.2%) patients after prostatectomy, respectively. Biopsy pathology classified 28 (14.7%), 81 (42.4%), and 82 (42.9%) patients into Gleason grade groups 1, 2, and 3–5, respectively. Prostate needle biopsy PNI was positive in 55 (22.8%) and absent in 136 (71.2%) of patients. On mpMRI, mECE was found in 12 (6.3%), suspicious mECE in 32 (16.8%), and mECE was absent in 147 (77%) patients (Table 1).

mpMRI findings for mECE and suspicious mECE were compared to final pathology to determine sensitivity, specificity, positive predictive value (PPV), and negative predictive value (NPV) of detecting pT3 disease (Table 2). mECE present had sensitivity, specificity, PPV, and NPV of 14.1% (95% confidence interval [CI] 7.3–24.8), 98.3% (95% CI 93.5–99.7), 87.5% (95% CI 50.9–97.1), and 65.9% (95% CI 58.4–72.7), respectively. On the other hand, mECE suspicious resulted in a sensitivity, specificity, PPV, and NPV of 23.9% (95% CI 14.9–35.8), 87.5% (95% CI 79.9–92.6), 53.1% (95% CI 35.0–70.5), and 66% (95% CI 58.1–73.2), respectively.

Univariable analysis revealed statistical significance between pT2 and pT3+ groups with regards to PSA, PNI, PI-RADS, and ECE. Age and prostate volumes did not reach significance (Table 1). On multivariable analysis, PNI on PBx predicted 2.25 times higher odds of pT3 disease ($p=0.045$, 95% CI 1.101–3.431). Similarly, presence of ECE on MRI also predicted pT3 disease with odds of 4.84 times ($p=0.048$, 95% CI 1.02–23.17). However, ECE suspicious on MRI did not significantly predict advanced stage at prostatectomy compared to absent ECE on MRI ($p=0.814$). Increased PI-RADS and PSA at MRI were both associated with an increase in the odds of predicting advanced stage PCA on RP (odds ratio [OR] 2.20, 95% CI 1.29–3.75, and 1.10, 95% CI 1.012–1.184, respectively) (Table 3). Age was not associated with increased odds of predicting pathological stage ($p=0.542$). When assessing for surgical margin status on binary logistic regression, only mECE present was statistically associated with an increased likelihood of exhibiting positive surgical margins, with an OR of 6.57 (95% CI 1.19–36.15, $p=0.031$) (Table 4). Fifty (90.9%) patients with PNI did not have mECE; of these, 28 (56.0%) were pT3.

In subgroup analysis, when controlling only for those patients without PNI ($n=136$), seven (5.1%) patients had mECE; of these, five (71.4%) were pT3. Within this cohort, mECE was still found to be a significant predictor of pT3 disease at RP ($p=0.03$). When analyzing mECE absent patients ($n=174$), 50 (90.9%) had PNI on PBx; of these, 28 (56.0%) were pT3 PNI on PBx ($n=50$). This cohort, however, did not significantly predict pT3 disease ($p=0.062$).

**Discussion**

To our knowledge, this is the first study to compare PNI and mECE together in a predictive model for at least pT3 disease at time of RP. We found mECE to have an increased likeli-

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**Table 1. Baseline characteristics by pathological stage on radical prostatectomy**

| Parameter                        | Total (n=191) | Pathological staging | p     |
|----------------------------------|---------------|----------------------|-------|
|                                 |               | T2 (n=120)           | T3a (n=56) | T3b (n=15) |
| PSA (mean ± SD)                  | 8.1±5.2       | 7.2±4.4              | 9.3±5.7  | 11.6±7.6   | 0.002 |
| Age (mean ± SD)                  | 61.8±7.1      | 61.8±7.1             | 61.1±7.5 | 63.3±5.9   | 0.630 |
| Prostate volume RP (mean ± SD)   | 51.2±20.7     | 52.4±21.7            | 49.0±19.2 | 49.6±18.3 | 0.266 |
| Prostate volume MRI (mean ± SD)  | 42.8±23.7     | 44.4±25.8            | 41.0±21.3 | 37.3±11.6 | 0.352 |
| Extracapsular extension (MRI)    |               |                      |         |           | <0.001|
| Present                          | 12            | 2 (16.7%)            | 5 (41.7%) | 5 (41.7%) |      |
| Suspicious                       | 32            | 15 (46.9%)           | 15 (46.9%) | 2 (6.3%)  |      |
| Absent                           | 147           | 103 (70.1%)          | 36 (24.5%) | 8 (5.4%)  |      |
| Perineural invasion on biopsy    |               |                      |         |           | <0.001|
| Present                          | 55            | 22 (40%)             | 25 (45.5%) | 8 (14.5%) |      |
| Absent                           | 136           | 98 (72.1%)           | 31 (22.8%) | 7 (5.1%)  |      |
| PI-RADS                           |               |                      |         |           | <0.001|
| 2                                | 14            | 11 (78.6%)           | 2 (14.3%) | 1 (7.1%)  |      |
| 3                                | 39            | 29 (74.4%)           | 8 (20.5%) | 2 (5.1%)  |      |
| 4                                | 94            | 67 (71.3%)           | 23 (24.5%) | 4 (4.3%)  |      |
| 5                                | 44            | 13 (29.5%)           | 23 (52.3%) | 8 (18.2%) |      |

*p-values reflect univariable analysis comparing parameters to pT2 and pT3 disease. ECE: extracapsular extension; MRI: magnetic resonance imaging; PI-RADS: Prostate Imaging-Reporting and Data System; PSA: prostate-specific antigen; RP: radical prostatectomy; SD: standard deviation.*
hood of predicting pT3 compared to that of PNI on PBx. Additionally, in our population, mECE also was shown to be an independent predictor of positive surgical margins. Importantly, when PNI was present but there was no ECE on MRI, there was no significant increase in pT3 disease.

Historically, preoperative risk-stratification methods have implemented certain clinical and laboratory elements to predict oncological outcomes. Common methods, such as the Partin tables or D’Amico risk stratification for prostatectomy oncological outcomes include the use of PSA, Gleason score, and clinical staging. While these stratification methods do not use PNI on PBx, PNI has been shown to be associated with adverse pathological outcomes at time of RP. A recent retrospective review by Celik et al stratified post-prostatectomy patients based on RP pathological stage, PNI presence on PBx, and D’Amico risk stratification. The authors suggested the presence of PNI is an important predictive marker due to positive correlation with local invasive disease after RP in all stratification groups, especially highlighting that those with intermediate D’Amico risk and clinical staging. While these stratification methods implemented certain clinical and laboratory elements to predict oncological outcomes. Common methods, such as the Partin tables or D’Amico risk stratification for prostatectomy oncological outcomes include the use of PSA, Gleason score, and clinical staging. While these stratification methods do not use PNI on PBx, PNI has been shown to be associated with adverse pathological outcomes at time of RP.

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An earlier review conducted by DeLancey et al also sought to determine the effect of PNI on adverse pathological features at time of RP, as well as to determine survival outcomes in these patients. On multivariable models controlling for biopsy Gleason score, PSA, and clinical stage, they found patients with PNI on PBx had 2.27, 2.64, and 1.64 increased odds of having ECE, seminal vesical involvement, and positive surgical margins, respectively. Additionally, their analysis of survival outcomes showed decreased disease-free and overall survival times.

While our results for predicting pT3 are in line with these findings, showing similar increased odds of 2.25, association of positive surgical margins did not achieve significance. This is likely related to controlling for different factors between our models (Table 3).

In another study examining PNI, Cozzi et al conducted a systematic review and meta-analysis in which they also found patients with PNI on PBx had a significant association with pT3 tumors at time of RP. Of their 7981 patients reported, they found PNI positivity on PBx to be 15.8%, and in patients with PNI, pT3 pathology was found in 53.4% of RP specimens. These findings agree with similar studies that showed just over half of patients with PNI having pT3. Our models (Table 3).

Table 2. Multiparametric MRI findings for mECE present and suspicious in detecting pT3 disease

| Parameter | Percentage (95% CI) |
|-----------|-------------------|
| mECE present | 14.1% (7.3–24.8) |
| Sensitivity | 98.3% (93.5–99.7) |
| Specificity | 83.3% (50.9–97.1) |
| PPV | 65.9% (58.4–72.7) |
| NPV | 66% (58.1–73.2) |
| mECE suspicious | 23.9% (14.9–35.8) |
| Sensitivity | 87.5% (79.9–92.6) |
| Specificity | 53.1% (35.0–70.5) |
| PPV | 60% (48.8–70.2) |
| NPV | 72.1% (67.1–76.5) |

Table 3. Multivariable cumulative odds ordinal regression predicting likelihood of pT3 on radical prostatectomy

| Parameter | OR (95% CI) | p |
|-----------|-------------|---|
| PNI present (Bx) | 2.25 (1.02–4.97) | 0.045 |
| PNI absent (Bx) | * | |
| ECE present (MRI) | 4.84 (1.01–23.17) | 0.048 |
| ECE suspicious (MRI) | 0.88 (0.29–2.66) | 0.814 |
| ECE absent (MRI) | * | |
| Age | 1.02 (0.94–1.07) | 0.542 |
| Prostate volume (MRI) | 0.99 (0.97–1.00) | 0.135 |
| PSA (MRI) | 1.10 (1.01–1.18) | 0.023 |
| PI-RADS | 2.20 (1.29–3.75) | 0.004 |

Table 4. Multivariable binomial regression predicting likelihood of positive surgical margins

| Parameter | OR (95% CI) | p |
|-----------|-------------|---|
| Age | 1.02 (0.97–1.08) | 0.436 |
| PNI (Bx) | 1.70 (0.75–3.86) | 0.209 |
| PSA (MRI) | 1.04 (0.96–1.13) | 0.328 |
| Prostate volume (MRI) | 0.99 (0.98–1.01) | 0.610 |
| ECE not present (MRI) | * | |
| ECE suspicious (MRI) | 2.18 (0.78–6.06) | 0.135 |
| ECE present | 6.57 (1.19–36.15) | 0.031 |

*Variable used as referent. CI: confidence interval; ECE: extracapsular extension; MRI: magnetic resonance imaging; OR: odds ratio; PI-RADS: Prostate Imaging-Reporting and Data System; PNI: perineural invasion; PSA: prostate-specific antigen.
tion imaging. Specifically, would mpMRI prediction of ECE exceed PNI on PBx as a predictor of non-organ-confined disease? Furthermore, does the presence of PNI even maintain predictive significance when mpMRI fails to show mECE?

Our study demonstrated mECE was, in fact, a better predictor of adverse pathology compared to PNI on PBx. The continued rise in mpMRI use for presurgical evaluation and management of PCa has allowed for further exploration into how these tools can be incorporated in predicting patient outcomes. While the impact of PI-RADS has been well-studied, mECE provides a logical tool to identify patients at risk for adverse postoperative pathology. In a study examining whether findings of mECE were associated with upstaging or upgrading at prostatectomy by Hedge et al., they followed the mECE T3 staging based on findings of asymmetric bulging, irregular margins, or direct extension of the lesion into periprostatic fat or neurovascular bundle staging, as reported previously by Roethke et al.26,27 On multivariable logistic regression, they found a 4.81 adjusted OR for increased likelihood of pT3 after RP compared to pT2. These findings are in line with our study results of 4.84 increased odds of pT3 compared to pT2 when mECE is present. Interestingly, our mECE suspicious group did not achieve significance for independently predicting pT3 at time of RP. This finding could perhaps be attributed to the inclusion of PI-RADS scoring into the regression model.

Using mECE in predictive models has also shown a potential benefit. Dominguez et al implemented mECE findings using a 1.5T mpMRI in patients with intermediate and high D'Amico risk to establish an area under the curve (AUC) of 0.7 for predicting non-organ-confined disease at time of RP.18 Taking a step further, when comparing these mECE against the Partin tables, Gupta et al found that mECE had an AUC of 0.82 compared to those of the Partin table's at 0.62 (p=0.04) in terms of predicting adverse pathological stage.26 Interestingly, sensitivity of mECE in predicting pT3 using a 3.0T MRI scanner is variable and generally low across the literature, ranging from 35–74%; our study was no different, reporting sensitivities of 14.1% and 23.9% for mECE present and mECE suspicious, respectively. However, specificity is consistently reported as robust for the prediction of pT3 with mECE, as we confirmed in our study, finding a specificity of 98.3% for mECE presence.22,29,30

The present study was not without its limitations. As a retrospective study, it is subject to inherent biases, including selection bias as a possible explanation for our higher observed pT3 rate at RP. Evaluating for both PNI and mECE decreased the included sample size, resulting in large CIs, specifically for mECE. Further validation with a larger cohort is warranted. Unlike the PI-RADS scoring system, reporting for mECE does not follow a strict, standardized procedure and is vulnerable to subjectivity and interobserver variability between radiologists. Evaluating mECE presents a challenge due to inherent subjectivity of proper characterization; although similar, a non-standardized and heterogeneous approach to MRI staging among studies exists in the literature.18,22,26,28 Unfortunately, there does not exist a standardized stratification for mECE at this time. Furthermore, there are a plethora of possible descriptions for addressing ECE that fall in between “no ECE” and “ECE present.” We attempted to account for this by stratifying the groups to separately analyze the less straightforward ‘mECE suspicious’ reported lesions. Although during this time period, many institutions did not routinely obtain mpMRI prior to PBx, our institution routinely obtained mpMRI prior to biopsy on patients with elevated PSA to allow for fusion biopsy approach. When challenged, we were often able to obtain high insurance approval rates through peer-to-peer authorization discussions.

Our findings support mECE as having the potential to significantly predict the presence of pT3 disease on RP. When excluding all patients with mECE present, PNI positivity on its own did not significantly predict pT3 disease. Conversely, when all PNI patients were excluded, mECE still retained its significance in predicting pT3. Although the development of an accepted standardized reporting system for ECE is necessary for inclusion into a nomogram for risk assessment, our study further supports the use of MRI preoperatively in perioperative surgical and treatment planning for organ-confined PCa.

## Conclusions

While mECE and PNI on PBx were both associated with worse pathological stage on RP, mECE had significantly higher cumulative odds compared to PNI. The significant predictive ability of ECE on multivariable analysis, and when controlling for only non-PNI patients, adds further clinical value to the use of mpMRI in PCa management. In patients with no suspicion for ECE on mpMRI, PNI is not associated with increased risk for T3 disease. This has potential implications when selecting patients for nerve-sparing prostatectomy. While validation in a larger cohort is required, these factors may have important clinical implications with regards to early diagnosis of advanced disease, while also highlighting the necessity for selective nerve sparing to be further explored.

### Competing interests:
The authors do not report any competing personal or financial interests related to this work.

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