ORIGINAL RESEARCH

Can Prostate-Specific Antigen Density Be an Index to Distinguish Patients Who Can Omit Repeat Prostate Biopsy in Patients with Negative Magnetic Resonance Imaging?

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Purpose: We evaluated the negative predictive value (NPV) of multiparametric magnetic resonance imaging (mpMRI) in detecting clinically significant prostate cancer (csPCa) according to biopsy setting and prostate-specific antigen density (PSAD) using transperineal template-guided saturation prostate biopsy (TPB) as the reference standard.

Methods: A total of 161 patients with biopsy histories and negative pre-biopsy mpMRI (Prostate Imaging Reporting and Data System version 2 scores of less than 3) participated in the study. TPB was performed on the following indications: "prior negative biopsy" in patients with persistent suspicion of prostate cancer (n = 91) or "confirmatory biopsy" in patients who were candidates for active surveillance (n = 70). The csPCa was defined as a Gleason score of 3 + 4 or greater. We calculated the NPV of mpMRI in detecting csPCa according to biopsy history and prostate-specific antigen density (PSAD) and conducted a logistic regression analysis to determine the clinical predicator for the absence of csPCa.

Results: The detection rate of csPCa was 5.5% in the prior negative biopsy group and 14.3% in the confirmatory biopsy group (P = 0.057). None of the variables in the logistic regression models including PSAD <0.15 ng/mL/cc and prior negative biopsy could predict the absence of csPCa. The NPV of mpMRI in detecting csPCa in patients with a prior negative biopsy worsen from 94.5% to 93.3% when combined with PSAD <0.15 ng/mL/cc.

Conclusion: Patients with negative mpMRI findings may not omit repeat biopsy even if their prior biopsy histories are negative and PSADs are <0.15 ng/mL/cc.

Keywords: early detection of cancer, prostatic neoplasm, prostate-specific antigen density, magnetic resonance imaging, negative predictive value, transperineal prostate biopsy

Introduction

Performing multiparametric magnetic resonance imaging (mpMRI) in patients with suspicion of prostate cancer (PCa) before prostate biopsy improves detection rates of clinically significant PCa (csPCa) by 2%-12% and reduces overdiagnosis of clinically non-significant PCa (cnsPCa) by 11%-14%.¹⁻³ In repeat biopsy settings, guidelines strongly suggested performing the mpMRI prior to repeat biopsy to identify a target lesion.^{4,5} It is currently recommended to perform a systematic biopsy when mpMRI is negative in a patient with prior negative biopsy. However, the possibility of distinguishing patients that can safely avoid biopsy by risk stratification combining mpMRI findings, clinical data, and other biomarkers has been proposed at the same time.⁶

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In this context, several studies have combined biopsy histories and other biomarkers to improve the negative predictive values (NPVs) of mpMRI in repeat biopsy setting (Table 1). Falagario et al analyzed different biopsy strategies based on the combined use of prostate-specific antigen density (PSAD) and mpMRI and suggested the individualized risk of missing PCa and csPCa.⁷ Recent study from Oishi et al investigated the NPVs of mpMRI in repeat biopsy setting using transrectal ultrasound (TRUS)-guided biopsy as the reference standard. They reported that the NPV of mpMRI were improved from 82% to 100%, when combined with negative biopsy history and PSAD <0.15 ng/mL/cc.⁸

Here, we attempted to validate the NPV of mpMRI in a repeat biopsy setting using a median of 36 core transperineal template-guided saturation prostate biopsy (TPB) as the reference standard. In particular, the csPCa detection rate and the NPV of mpMRI were investigated in combination with prior biopsy status and PSAD levels.

Patients and Methods

Study Population

This study was approved by the Institutional Review Board of Samsung Medical Center (approval number: 2018–05-192) and performed in accordance with the principles of the Declaration of Helsinki. We retrospectively reviewed the clinical data of patients who underwent TPB in repeat biopsy setting between May 2017 and December 2020. Patients who underwent mpMRI at our institution and had negative mpMRI reports, defined as a Prostate Imaging Reporting and Data System version 2 (PI-RADS v2) score of less than 3, were included in this study. While those who had undergone any prior surgical or medical treatment for benign prostatic hyperplasia or previously diagnosed PCa were excluded.

TPB was performed if clinically indicated by 1) a persistent suspicion of PCa with rising PSA despite a "prior negative biopsy" or 2) a "confirmatory biopsy" in patients with prior cnsPCa who are candidates for active surveillance (AS). Based on biopsy histology, csPCa was defined as a Gleason Score (GS) of 3 + 4 or greater, and cnsPCa was defined as a GS of 6.

MRI Protocol and Interpretation

T1-weighted, T2-weighted, and diffusion-weighted imaging (apparent diffusion coefficient maps and long-b scans) and dynamic gadolinium contrast-enhanced imaging were performed using a 3-tesla MRI instrument (Intera Achieva TX; Philips Healthcare, Best, the Netherlands) with a phased-array body coil.

All mpMRI findings were scored using the PI-RADS v2 by two dedicated uro-radiologists with more than 10 years of experience in prostate MRI interpretation. The prostate volume was calculated using the ellipsoid formula based on MRI measurements.

Transperineal Template-Guided Saturation Prostate Biopsy Protocol

The patients were given a cleaning enema and antibiotics before biopsy. All biopsies were performed in an extended lithotomy position under general anesthesia. After a digital rectal exam, biopsy was performed via a standard 5-mminterval brachytherapy grid fixed on a stepper from the medial to the lateral plane in the anterior, middle, and posterior portions of the prostate gland under the guidance of a biplanar TRUS 8848 probe (BK Medical, Herlev, Denmark). All patients underwent 24 or 36 systematic saturation biopsies depending on prostate size relative to 30 cc according to the Ginsburg protocol.⁹ No MRIdirected biopsies were performed in cases with lesions with PI-RADS scores less than 3.

Statistical Analysis

The Mann–Whitney *U*-test was used for continuous variables, and the chi-square or Fisher's exact test was used for categorical variables. The NPVs of mpMRI in detecting csPCa were calculated according to PSAD category and biopsy history. In addition, we assessed the NPV of MRI in combination with prior negative biopsy history and PSAD cutoff of 0.15 ng/mL/cc, which had the lowest Youden index. Univariate and multivariable logistic regression analyses were performed to assess the predictors of absent csPCa with clinically relevant parameters including previous biopsy status and PSAD level. Statistical analyses were conducted using the Statistical Package for the Social Sciences version 21.0 (IBM Corp., Armonk, NY, USA). All p-values were two-tailed, and a p-value less than 0.05 was considered statistically significant.

Results

A total of 452 consecutive men underwent TPB in repeat biopsy setting at our institution between May 2017 and December 2020, and 171 had negative results in prebiopsy mpMRIs. Of the 171 patients with negative mpMRIs, we excluded 6 patients who underwent treatment for benign

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Study (Year)	No. with Negative MRI	Population	Definition of Negative MRI	Definition of csPCa	Reference Standard	Biomarkers	Marker Cutoff Value	NPV of MRI	Negative MRI + NegBx	Negative MRI + Biomarker	Negative MRI + NegBx + Biomarker
Washino (2017) ¹²	127	BxN = 127	PI-RADS version 2 <3	GS≥3+4 or CCL ≥4mm	14 cores Transperineal Bx	PSAD	0.15	83%		%00 I	
Distler (2017) ¹³	344	BxN = NR NegBx = NR	PI-RADS version 1 <3	GS≥3+4	Median 24 cores Transperineal Bx	PSAD	0.15	79%	83%	89%	93%
Druskin (2018) ¹⁴	20	BxN = NR NegBx = NR	PI-RADS version 2 <3	GS≥3+4 or GS6 in > 1 core or CCL>50% of one core	Not described	DIHA	0.44	%06		%00 I	
Perlis (2018) ¹⁵	28	NegBx = NR AS = NR	PI-RADS version 2 <3	GS≥3+4 or CCL>4mm	12–16 cores Transrectal Bx	PCA3	35	95%		%00 I	
Oishi (2019) ⁸	135	BxN = 48 NegBx = 60 PosBx = 27	PI-RADS version 2 <3	GS≥3+4	Median 13 cores Transrectal Bx	PSAD	0.15	82%	92%	%06	%00 I
Falagario (2020) ⁷	480	BxN = 349 NegBx = 131	PI-RADS version 2 <3	GS≥3+4	Transrectal Bx	QA29	0.15	79%	95 %	96%	97%
Current study	161	NegBx = 91 PosBx = 70	PI-RADS version 2 <3	GS≥3+4	Median 36 cores Transperineal Bx	PSAD	0.15	%2.06	94.5%	89.5%	93.3%
Abbreviations : MRI, π PSAD, Prostate Specific	nagnetic resonanc Antigen density;	e imaging; BxN, bi PHID, Prostate He	opsy naïve; NegBx, salth Index density;	Abbreviations: MRI, magnetic resonance imaging; BxN, biopsy naïve: NegBx, prior negative biopsy: PosBx, prior positive biopsy: PI-RADS, Prostate Imaging Reporting and Data System; GS, Gleason Score; CCL, cancer core length; PSAD, Prostate Specific Antigen density; PHID, Prostate Health Index density; PCA3, prostate cancer antigen 3; NPV, negative predictive value.	prior positive biopsy 3; NPV, negative pr	r; PI-RADS, Prostatu edictive value.	e Imaging Repo	rting and D	ata System; GS,	Gleason Score; C	.CL, cancer core length;

Table I Previous Manuscripts Addressed the Negative Predictive Value of MRI in Combination with Negative MRI and Other Biomarkers

		Biopsy Ir	ndication	
	Total	Prior Negative Biopsy	Confirmatory Biopsy	Р
Number of patients	161	91	70	
Median age (IQR) (years)	62 (58–68)	62 (58–67)	64 (58–69)	0.237
Suspicious DRE, n (%)	2 (1.2)	2 (2.2)	0 (0)	0.505*
Family history of PCa, n (%)	9 (5.6)	6 (6.6)	3 (4.3)	0.733*
Median PSA (IQR) (ng/mL)	5.6 (4.2–8.6)	6.6 (4.8–10.0)	4.6 (3.6–6.6)	<0.001
Median prostate volume (IQR) (cc)	40.4 (31.3–54.8)	43.8 (34.8–57.0)	37.7 (29.7–50.0)	0.032
Median PSAD (IQR) (ng/mL/cc)	0.14 (0.10-0.20)	0.15 (0.11–0.24)	0.13 (0.09–0.16)	0.004
<0.15 ng/mL/cc, n (%)	95 (59.0)	45 (49.5)	50 (71.4)	0.005
Prior TRUS-guided biopsy Benign prostate tissue HGPIN ASAP	83 (51.6) 5 (3.1) 3 (1.9)	83 (91.2) 5 (5.5) 3 (3.3)	0 (0) 0 (0) 0 (0)	
PCa (GS 3+3)	70 (43.5)	0 (0)	70 (100)	

Table 2 Demographics of Patients with Negative MRIs Who Underwent Transperineal Template-Guided Saturation Biopsies

Notes: *Fisher's exact test.

Abbreviations: MRI, magnetic resonance imaging; IQR, interquartile range; DRE, digital rectal examination; PSA, prostate-specific antigen; PSAD, PSA density; TRUS, transrectal ultrasound; HGPIN, high-grade prostatic intraepithelial neoplasia; ASAP, atypical small acinar proliferation; PCa, prostate cancer; GS, Gleason score.

prostatic hyperplasia and 4 patients who received androgen deprivation therapy for PCa diagnosed in a prior biopsy. Therefore, 161 patients met the inclusion criteria for this study. Among this cohort, 91 (56.5%) patients were indicated for TPB by a persistent suspicion of PCa with rising PSA despite a negative result on prior TRUSguided biopsy, and 70 (43.5%) patients were indicated for a confirmatory biopsy for the previously detected cnsPCa by TRUS-guided biopsy.

The baseline demographics and characteristics of the patients with negative mpMRIs are summarized in Table 2. Patients in the prior negative biopsy group had significantly higher PSA levels and larger prostate volumes than those in the confirmatory group (P < 0.001 and P < 0.032, respectively). PSAD levels were higher in the prior negative biopsy group, and the proportion of patients with PSAD < 0.15 ng/mL/cc was higher in the confirmatory group (P = 0.004 and P = 0.005, respectively). Of the 91 patients with prior negative biopsies, 5 (5.5%) had previously been diagnosed with high-grade prostatic intraepithelial neoplasia, and 3 (3.3%) were diagnosed with atypical small acinar proliferation.

The TPB results are presented in Table 3. According to the TPB protocol of our institution, the median number of

biopsy cores (interquartile range) was 36 (24–36) in both groups. The detection rate of any PCa was 23.1% in the prior negative biopsy group and 58.6% in the confirmatory biopsy group, including 8 patients with GS 7 (3+4) PCa and 2 patients with GS 8 (4+4) PCa. There was a significant difference in the detection rate of any PCa between the two groups (P < 0.001). However, the detection rate of csPCa was 5.5% in the prior negative biopsy group, and there was no statistical difference between the groups (P = 0.057).

The NPV of mpMRI in detecting csPCa was 90.7%, 94.5%, and 85.7% in the overall study population, previous negative biopsy group, and confirmatory biopsy group, respectively (Table 4). In the previous negative biopsy group, the NPV of mpMRI was 100% in PSAD categories of 0.15–0.20 ng/mL/cc and PSAD <0.10 ng/mL/cc. However, the PSAD category of 0.10–0.15 ng/mL/cc had the lowest NPV at 89.7%. When combined with PSAD < 0.15 ng/mL/cc, it was found to be lower at 89.5% (Figure 1). In addition, in the prior negative biopsy group, the NPV of mpMRI was 94.5% and it was found to worsen at 93.3% when combined with PSAD < 0.15 ng/mL/cc.

		Biopsy Indication		
	Total	Prior Negative Biopsy	Confirmatory Biopsy	Р
Number of patients	161	91	70	
Median number of biopsy cores (IQR)	36 (24–36)	36 (24–36)	36 (24–36)	0.959
Median number of positive cores (IQR)	0 (0-1)	0 (0–0)	I (0-4)	<0.001
Results of TPB, n (%)				
Benign prostate tissue	93 (57.8)	65 (71.4)	28 (40.0)	
HGPIN	2 (1.2)	1 (1.1)	I (1.4)	
ASAP	4 (2.5)	4 (4.4)	0 (0)	
PCa (GS 3+3)	47 (29.2)	16 (17.6)	31 (44.3)	
PCa (GS 3+4)	11 (6.8)	3 (3.3)	8 (11.4)	
PCa (GS 4+3)	2 (1.2)	2 (2.2)	0 (0)	
PCa (GS 4+4)	2 (1.2)	0 (0)	2 (2.9)	
Median maximal percent of cancer in core (IQR)	10 (5–30)	10 (5–30)	15 (10-40)	0.148
Detection of any PCa, n (%)	62 (38.5)	21 (23.1)	41 (58.6)	<0.001
Detection of csPCa, n (%)	15 (9.3)	5 (5.5)	10 (14.3)	0.057

Table 3 Results of 7	Fransperineal	Template-Guided	Saturation Bio	opsy in	Patients with	Negative MRIs
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Abbreviations: MRI, magnetic resonance imaging; IQR, interquartile range; TPB, transperineal template-guided saturation biopsy; HGPIN, high-grade prostatic intraepithelial neoplasia; ASAP, atypical small acinar proliferation; GS, Gleason score; PCa, prostate cancer; csPCa, clinically significant prostate cancer.

Univariate and multivariate logistic regression models were established for the analysis of association with the absence of any PCa or csPCa based on age, family history of PCa, PSA level, prostate volume, PSAD < 0.15 ng/mL/cc, and the indication of TPB (Table 5). In the univariate model, higher PSA level (OR 1.089, 95% CI 1.009–1.177, P = 0.029) and prior negative biopsy group (OR 4.197, 95% CI 2.073–8.496, P < 0.0001) were predictive of the absence of any PCa. The multivariate model revealed that only the prior negative biopsy group was predictive of the absence of any PCa (OR 4.197, 95% CI 2.073–8.496; P < 0.0001). In contrast, none of the variables were predictive of the absence of csPCa on TPB in either univariate or multivariate analysis.

Discussion

In this retrospective study, we examined the any PCa and csPCa detection rates in patients who had no suspected lesions on mpMRI (PI-RADS <3) and underwent repeat biopsies using TPB. We also calculated the NPV of mpMRI by combining the prior biopsy histories and PSAD levels. To the best of our knowledge, this is the first study to analyze the NPV of mpMRI scored by PI-RADS v2 by performing TPB with a median of 36 cores.

Our institution has performed TPB if clinically indicated by a persistent suspicion of PCa with rising PSA despite a previous negative result on TRUS-guided biopsy and as confirmatory biopsy in patient candidates for AS

Table 4 Negative Predictive Value of	MRI According to Prostate-S	Specific Antigen Density	Category and Biopsy History
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PSAD	Total	Biopsy Ir	ndication
		Prior Negative Biopsy	Confirmatory Biopsy
Total	90.7% (146/161)	94.5% (86/91)	85.7% (60/70)
PSAD ≥0.20 ng/mL/cc	92.7% (38/41)	93.5% (29/31)	90.0% (9/10)
0.15 ng/mL/cc≤ PSAD <0.20 ng/mL/cc	92.0% (23/25)	100.0% (15/15)	80.0% (8/10)
0.10 ng/mL/cc≤ PSAD <0.15 ng/mL/cc	87.9% (51/58)	89.7% (26/29)	86.2% (25/29)
PSAD <0.10 ng/mL/cc	91.9% (34/37)	100% (16/16)	85.7% (18/21)

Abbreviations: MRI, magnetic resonance imaging; PSAD, prostate-specific antigen density.

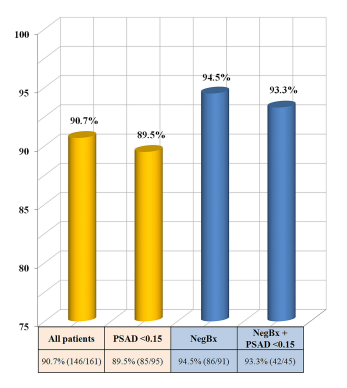


Figure I Negative predictive value of multiparametric magnetic resonance imaging in detecting clinically significant prostate cancer in combination with prior negative biopsy history and PSAD cutoff of 0.15 ng/mL/cc.

after a prior detected cnsPCa on TRUS-guided biopsy. In 91 (56.5%) patients with prior negative biopsies and 70 (43.5%) patients with prior positive biopsies, the histology of a median of 36 cores of TPB using 5-mm sampling was used as the reference standard. The incidence of biopsy-detected csPCa was 5.5% in men with negative mpMRI and prior negative biopsy history, and 14.3% in men with negative mpMRI and prior positive biopsy history. The NPV of mpMRI in the detection of csPCa in patients with prior negative biopsies worsen from 94.5% to 93.3% when combined with PSAD < 0.15 ng/mL/cc.

The multi-institutional Prostate MR Imaging Study (PROMIS) trial evaluated the detection rate of PCa in 576 patients who had undergone prebiopsy mpMRI with saturation transperineal prostate biopsy as the reference standard.¹⁰ Of the 158 patients with negative mpMRI in the study, 141 patients reported having no cancer or non-significant cancer (GS 6) on transperineal prostate biopsy, and the NPV of mpMRI predicting the absence of GS 7 (3 + 4) or greater cancer was 89% (95% confidence interval 83%–94%). PROMIS trial presents the possibility of reducing the proportion of men with unnecessary biopsies. However, limitations of the prebiopsy mpMRI were also

reported. Vargas et al confirmed that PI-RADS v2 was limited for the assessment of GS \geq 4+3 tumors with a volume of less than 0.5 mL.¹¹ In addition, in situations where approximately 30% to 40% of the patients have been reported to have negative results on prebiopsy MRIs, it is not yet clear which patients can safely omit the biopsy.

There have been several reports on improving the NPV of mpMRI in combination with clinical information with negative mpMRI findings (Table 1).^{8,12–15} In particular, two studies of Perlis et al and Oishi et al evaluated the NPV of mpMRI by adding prior biopsy histories and different biomarkers using TRUS-guided biopsy as the reference standard. Perlis et al investigated patients with a previous negative biopsy and patients on AS and reported that with a negative multiparametric MRI and normal prostate cancer antigen 3 (PCA3) score, no csPCa was detected on repeat TRUS-guided biopsy.¹⁵ Therefore, they suggested that repeat biopsies may not be needed in patients with dual negative results (negative results from both mpMRI and PCA3 scores). A recent study by Oishi et al reported outcomes of repeat biopsy in 135 patients with negative prebiopsy mpMRIs.⁸ The detection rate of csPCa (GS 3 + 4 or greater) was 18% in men with negative mpMRIs. The NPV of mpMRI was 100% for patients with prior negative biopsies who had PSAD < 0.15 ng/mL/cc, and the authors proposed that this subset of patients could avoid repeat biopsy.

However, because the reference standard of the aforementioned two studies was TRUS-guided biopsy, information regarding omitting biopsy might be limited. The TRUS-guided biopsy can be associated with both the underdetection of csPCa and the increased diagnosis of many cnsPCas.^{10,16,17} In addition, in TRUS-guided biopsies, either the size or the grade of PCa may be underestimated due to random sampling error.¹⁸ Furthermore, the pathological status derived from TRUS-guided biopsies in the repeat biopsy setting can be unreliable in discriminating csPCa from cnsPCa, such that 34% of the men have been found to have an upgrading of PCa.¹⁹ Thus, TRUS-guided biopsy may not be a sufficient modality to determine the NPV of prebiopsy mpMRI. Despite meeting the criteria for combining negative MRI results and other clinical information or biomarkers, patients could still harbor risk of csPCa, and whether biopsies can be omitted in these patients remains unclear. In patients with prior negative biopsy of this present study, PSAD category of 0.10-0.15 ng/mL/cc had the lowest NPV. Practically, in

		Absence of any PCa	í any PCa			Absence of csPCa	ıf csPCa	
	Univariate Analysis	ysis	Multivariate Analysis	ılysis	Univariate Analysis	rsis	Multivariate Analysis	ysis
	OR (95% CI)	٩	OR (95% CI)	٩	OR (95% CI)	٩	OR (95% CI)	٩
Age	0.970 (0.928–1.013)	0.172			0.965 (0.896–1.039)	0.347		
FHx (No vs Yes)	1.297 (0.33 4–5 .026)	0.707			3.055 (0.574–16.247)	061.0		
PSA	(771.1–600.1) 680.1	0.029	1.044 (0.967–1.128)	0.269	1.079 (0.935–1.246)	0.297		
Prostate volume	1.014 (0.998–1.031)	0.094			1.023 (0.987–1.059)	0.208		
PSAD <0.15 ng/mL/cc	0.549 (0.283–1.065)	0.076			0.697 (0.227–2.141)	0.528	I.142 (0.358–3.645)	0.822
Biopsy indication Confirmatory biopsy Prior negative biopsy	Ref 4.713 (2.385–9.313)	100:0>	Ref 4.197 (2.073–8.496)	100.00	Ref 2.867 (0.933–8.812)	0.066	Ref 2.786 (0.883–8.791)	0.081
Abbreviations: MRI, magne Gleason score; PCa, prostate	Abbreviations: MRI, magnetic resonance imaging; IQR, interquartile range; TPI Gleason score; PCa, prostate cancer; csPCa, clinically significant prostate cancer.	erquartile range; TP ant prostate cancer.	B, transperineal template-guide	d saturation biopsy:	Abbreviations: MRI, magnetic resonance imaging; IQR, interquartile range; TPB, transperineal template-guided saturation biopsy; HGPIN, high-grade prostatic intraepithelial neoplasia; ASAP, atypical small acinar proliferation; GS, Gleason score; PCa, prostate cancer; csPCa, clinically significant prostate cancer.	traepithelial neopla	sia; ASAP, atypical small acinar _I	oroliferation; GS,

our study, among the five patients diagnosed with csPCa in the prior negative biopsy group, three patients had PSAD < 0.15 ng/mL/cc (0.10, 0.11, and 0.14 ng/mL/cc). Furthermore, the logistic regression analysis showed none of the variables were predictive of the absence of csPCa. These results suggest that there would be limitations in distinguishing between patients who can safely avoid biopsy with PSAD, even in patients with prior negative biopsy and negative MRI.

In contrast to TRUS-guided biopsy, TPB has very high sensitivity and specificity for csPCa detection, and the results of TPB more accurately reflect the subsequent pathological results of radical prostatectomy than those of TRUS-guided biopsy.²⁰⁻²² The TPB has the advantage of approaching the anterior prostate and the transitional zone and produces a histological map of the entire prostate gland.²³⁻²⁵ This biopsy modality not only produces a three-dimensional map of the entire prostate gland but also evaluates small lesions (<0.5 mL) that mpMRI may not assess.^{13,26,27}

There was a previous study of the NPV of mpMRI using transperineal approach. Distler et al combined mpMRI with PSAD to predict the detection rate of csPCa (GS 3 + 4 or greater) by performing transperineal systematic biopsy in biopsy-naive or previously negative patients.13 They described the NPV of mpMRI for detecting csPCa in patients with negative mpMRI (PI-RADS version 1 score less than 3). Unlike our study, the NPV of mpMRI in patients with prior negative biopsies increased from 83% to 93% in combination with PSAD < 0.15 ng/mL/cc. This might be explained that the NPV of mpMRI was too low due to the low sensitivity of PI-RADS version 1; thus, this could be interpreted that the NPV improvement was observed when PSAD was combined. In our study, the NPV of mpMRI in patients with a prior negative biopsy was 94.5%, which was much higher than the result of the study (83%). Statistically, as the sensitivity of a test increases, the NPV also increases. Regarding the sensitivities of PI-RADS version 1 and PI-RADS v2, Tewes et al evaluated the diagnostic accuracy according to PI-RADS version 1 and PI-RADS v2 of two independent readers, and reported sensitivities of reader 1 and reader 2 of 84% and 74% for PI-RADS version 1, respectively, and 100% and 81% for PI-RADS v2, respectively.²⁸ A head-to-head comparison of PI-RADS version 1 and PI-RADS v2 stratified by cancer location reported higher sensitivity of PI-RADS v2 than version 1 (87.5-100% vs 75%) in transitional zone cancer, and the

similar sensitivity of the two versions in peripheral zone cancer (both 100%).²⁹ Furthermore, the relatively low prevalence of PCa in our Asian cohort compared to that in the Western population could affect higher NPV of mpMRI.^{30,31}

Our study also has several limitations. First, due to its nonrandomized retrospective cohort study design, the patient information in our data might be incomplete. Because our institution also performed repeat prostate biopsy with the transrectal approach in some patients who did not want general anesthesia, selection bias that can impact the results could not be eradicated. Second, the cohort size was relatively small. However, TPB under general anesthesia was used to obtain more detailed information on the prostate of the participants. Furthermore, to maintain the uniformity of our study cohort, we excluded patients who had undergone medical or surgical treatment for benign prostatic hyperplasia or PCa. Third, the present study focused primarily on the pathological findings from biopsies, but not radical prostatectomy specimens. Nevertheless, we performed TPB in all patients with the most accurate biopsy modality to reflect the pathologic results of the prostatectomy specimens.²¹ Finally, MRI scans were not specifically reviewed in this study. Nonetheless, highquality mpMRI was performed with an identical instrument for all patients, and two experienced uro-radiologists interpreted and reported the images according to the PI-RADS criteria.5

Conclusion

The NPV of mpMRI in patients with a prior negative biopsy did not improve when PSAD was combined. Patients with negative MRIs, prior negative biopsies and PSAD < 0.15 ng/mL/cc may not omit repeat biopsies, as some patients could be diagnosed with csPCa, especially when performing TPB.

Ethical Approval

This study was approved by the institutional review board at Samsung Medical Center and abided with the Helsinki Declaration. For this type of study, formal consent was not required and data were anonymous.

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Author Contributions

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agree to be accountable for all aspects of the work.

Disclosure

All authors have no direct or indirect conflict of interest with any institution or product.

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