Title: Towards a MRI-based nomogram for the prediction of transperineal prostate biopsy outcome: a physician and patient decision tool

Article Type: Original Article

Keywords: Magnetic Resonance Imaging; Nomogram; Prostate Biopsy; Prostate Cancer; Risk Assessment

Abstract: Abstract:

Purpose: To develop and internally validate a nomogram utilising biparametric magnetic resonance imaging (B-MRI)-derived variables for the prediction prostate cancer at transperineal sector-guided prostate biopsy (TPSB).

Subjects/Patients and Methods:
Consecutive patients referred to our institution with raised PSA, abnormal prostate examination or persistent suspicion of prostate cancer after previous transrectal biopsy between July 2012 and November 2015 were reviewed from a prospective database.

All patients underwent pre-biopsy B-MRI with T2-weighted and diffusion-weighted imaging sequences, followed by 24-40 core TPSB with additional targeted cores using cognitive registration.

Univariable and multivariable logistic regression analysis was used to determine predictors of prostate cancer outcomes. Multivariable coefficients were used to construct two MRI-based nomograms to predict any and significant (Gleason 4 or Maximum Cancer Core Length ≥6mm) prostate cancer at TPSB. Bootstrap resamples were used for internal validation. Accuracy was assessed by calculating the concordance index (c-index).

Results:
In total, 615 men were included in the study. Prostate cancer was diagnosed in 317 (51.5%) men with significant cancer diagnosed in 237 (38.5%) men.

Age, PI-RADS score, PSA, PSA Density (PSAD) and Primary Biopsy were predictors of prostate cancer at TPSB on univariable analysis (p<0.0001). PSA showed strong correlation with PSAD and was excluded. The remaining variables were all independent predictors of prostate cancer on multivariable analysis (p<0.0001) and used to generate the nomograms.
Both nomograms showed good discrimination for prostate cancer, with a c-index of 87% for any cancer and 92% for significant disease. Using a nomogram-derived probability threshold of <15%, 111 (18.0%) of biopsies can be saved, at the expense of 3 missed significant prostate cancers.

Conclusions:
These internally-validated MR-based nomograms were able to accurately predict TPSB outcomes for prostate cancer, especially significant disease. Our findings support the combination of pre-biopsy MRI results and clinical factors as part of the biopsy decision-making process.
Conflict of Interest Statement

Article Title: Towards a MRI-based nomogram for the prediction of transperineal prostate biopsy outcome: a physician and patient decision tool

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Highlights

- Two MRI-based nomograms for prediction of prostate biopsy outcomes are proposed
- Variables used include MRI PI-RADS scores, PSA density, age and primary biopsy
- The nomograms show good discrimination for any and significant prostate cancer
- MRI and clinical findings can be used within patient decision-making process
To the editor:

We are very grateful for the reviewers’ comments. Please find the responses below in red. Where possible, these have been incorporated into the text.

**Reviewer #1:** Authors present 2 nomograms to predict clinically significant prostate cancer using B-MRI prior to perineal saturation biopsy. Several issues with this study which hamper my enthusiasm:

1. B-MRI is not sufficient to characterize prostate cancer lesions.
2. Authors used PIRAD v1 not V2 (likely for this reason)

Both the MRI protocols and PI-RADS are evolving. The nature of any study like this is that when results are available, newer technologies will have emerged, that can be fed back in to improve the process. When the study began, dynamic contrast enhancement (DCE) was not available in our institution. Nevertheless, we were able to demonstrate high sensitivity of B-MRI (Grey et al.) without DCE. B-MRI has compared favourably against mpMRI protocols that include DCE, including the recently published PROMIS study. A further study has directly compared B-MRI to mpMRI and found equivalent performance characteristics. We have included these references, etc. within the Discussion section.

At the present time, the use of DCE is less resource friendly than simple B-MRI and there are debates with regards to its safely. Therefore the present study is applicable to institutions using MRI with and without DCE. It is envisaged that further studies will include DCE and nomograms will incorporate those results as an additional feature.

3. There was no registered fusion biopsy, just cognitive

We used systematic sector guided biopsies as well as cognitive-guided biopsies (2-4 cores) as the reference standard, which is widely available to clinicians and avoids the variability associated with different fusion biopsy systems. This has been highlighted in the text.

4. PSA density was based on formulaic calculation of prostate size (which can be very inaccurate) compared to image segmentation

The authors agree that segmental calculation of prostate volume is more accurate than formulaic calculation; however, for the purposes of this nomogram, the aim was to make calculations as straightforward and user-friendly as possible. Formulaic calculation is relatively quick, requires very little training and does not require further software, thus making the nomogram input data relatively accessible. We have found that PSA density using this method provides an improvement in ROC area under the curve over total PSA and is a useful adjunct in order to predict biopsy outcome. This has been incorporated into the manuscript.

5. Why saturation perineal biopsy?? The only benefit is decreased infection rate

We aimed to provide as comprehensive a reference standard as possible. Short of radical prostatectomy, our view was that saturation perineal biopsy provided this. A comment to reflect this has been added to the discussion section.

**Reviewer #2:** In this study the authors developed a novel nomogram for predicting the presence of
cancer on prostate biopsy. Unique to their nomogram is the incorporation of biparametric MRI findings (PIRADS values) and PSA density (as determined using MRI). Two novel aspects of this study are (1) the fact that patients underwent a perineal biopsy with >20 cores taken, therefore the study includes an excellent truth standard for presence of cancer and (2) the authors use of biparametric rather than multiparametric MRI. The paper is well written and it is presented in an unbiased and logical fashion. The authors should consider the following edits to the paper:

1. In the last sentence of the Introduction the authors state: "In the present study, we aimed to develop a new nomogram, utilising the information provided by pre-biopsy biparametric MRI (B-MRI) and PSAD." This gives the impression that the nomogram was built without considering other variables. Reading the methods this is not so. Would reword to say "we aimed to develop a novel nomogram utilizing both clinical and biparametric MRI data for predicting the presence of cancer on prostate biopsy."

Thank you, this has been revised.

2. The authors utilized biparametric MRI. References should be provided for this non-standard form of prostate MRI. In addition, the authors should provide reference to or comment on the validity of the PIRADs system with only use of T2 and DWI imaging. This would be most appropriate for the Discussion section.

We agree that the majority of studies use DCE and our B-MRI protocol differs from this. As per reviewer #1, points #1/2, we have added additional discussion with regards to this.

3. In the methods section the authors need to clearly define their definition of "clinically significant prostate cancer."

Thank you, this has been further emphasised in the Methods section.

4. c-index values are not typically presented as percentages but rather as values <1 with two to three decimals. For example c = 0.528. The authors should consider presenting in this manner rather than as percentages.

Thank you, this has been revised as suggested.

5. Did the authors try a model of PSA + Prostate vol + PIRADs? I wonder if this would perform better than density which is a calculation of PSA and prostate vol. This should be commented on.

We ran multivariable analyses for each outcome using two models: the first using PSA and volume separately and the second using PSAD instead. We compared the fit between both models using analysis of covariance and found the model using PSAD had significantly greater reduction in the residual sum of squares (p<0.05), indicating a better fit, for all outcomes. Therefore, the final model was performed using PSAD. This has been clarified in the results section.

Reviewer #4: In this study Su-Min Lee et al conduct an analysis of their prospectively collected patient who were referred to their institution for possible prostate biopsy. They conducted an analysis to identify significant factors to predict any and significant cancer on TPSB. The manuscript is well written and the statistical analysis conducted is appropriate for such study. Below are my comments.
1. The authors discuss use of DRE in assessing patients for possible prostate biopsy and this is mentioned in the introduction. While many have their bias about how useful the exam is, the authors do not appear to have incorporated this as a patient variable in their analysis. In a prospectively collected database this was likely collected or at least the clinical stage was noted I imagine. A reason for not doing so should be stated and ideally supported.

DRE in the clinic setting was carried out by a large number of clinicians in primary and secondary care settings, and thus, subject to high levels of variability. Rightly or wrongly, many patients who were listed for general anaesthetic biopsy did not have an expert DRE prior to biopsy when an examination under anaesthetic was undertaken. We felt it would be wrong to use this as we aimed to provide a predictive nomogram prior to biopsy.

2. The authors make mention several times in the introduction and in the discussion about the benefits of TPSB over TRUS and mention several statistics regarding the possible complications of TRUS. In order to provide a statistical comparison the authors should comments on their patient cohort in regards to complications (infections, hematuria, anesthesia risk etc).

We acknowledge the comments regarding the complication rates following TRUS vs transperineal biopsy. We have included data regarding our patient cohort and subsequent complications in the Results text.

3. They also comment on cost effectiveness of TPSB and the decreased morbidity with its use. Data to support this statement should be provided.

The statement was intended to convey that the use of risk assessment could allow for increased cost-effectiveness of our current diagnostic pathway by reducing negative biopsies. We note that our overall phrasing regarding transperineal biopsy was confusing and this paragraph has been revised to avoid misinterpretation.

4. One major issue with this manuscript is the authors do not mention several other recent studies that he looked at nomograms or analysis that aimed at incorporating prostate MRI in decision making for prostate biopsy. They mention a two in the discussion but there are several others that should be directly commented on in the discussion about what makes their study different/novel/better. Below are just a few.

- Wang R et al- Pre-biopsy mp-MRI can help to improve the predictive performance in prostate cancer: a prospective study in 1478 consecutive patients. Clin Cancer Res. 2017 Jan 31
- van Leeuwen PJ et al- A multiparametric magnetic resonance imaging-based risk model to determine the risk of significant prostate cancer prior to biopsy. BJU Int. 2017 Feb 16

Thank you – we have discussed additional studies from the literature that develop nomograms or use MRIs in the decision making process for prostate biopsy, including the additional references as mentioned by the reviewer. What is different about ours? Our population consists of largely unscreened patients, primarily presenting with LUTS secondary to BPH with associated raised PSA. This may explain the higher discrimination seen in our study and is discussed in the Discussion section.
5. The authors appear to not use DCE in their model, which if they are using PI-RADS v1, this would be included in all cases. They offer reason in the discussion about this (PI-RADS v2) but do not comment on why it was not included in the introduction or the analysis. This needs to be mentioned and a reason why they, despite reporting PI-RADS v1 data, do not have in their model.

We have added additional discussion as to our use of B-MRI and the absence of DCE imaging as per Reviewer #1, point #1/2. We also comment on PI-RADS DWI/DCE imaging in the introduction as requested, and reiterate this in the Discussion.

6. Another major drawback from the study is they mention PSAD a way to improve upon the limits of serum PSA. Several readily available blood tests, 4K and PHI, also do this and do not require an MRI. The authors need to comment on these tests and how they may make several of their proposed nomogram factors obsolete (PSA, PSAD) or if they could be integrated in their model. They comment on PCA3 which has largely been replaced by the above mentioned test. Even if their reason is "availability" they need to discuss more since most readers will have access to them. These test are becoming more commonly used by clinicians and a reason to use MRI for the attainment of PSAD and PI-RADS as a first test should be discussed.

Blood markers have been discussed in further detail, including 4K and PHI, as well as the aforementioned PCA3. These weren’t available to us at the time of the study and currently are not in widespread use. We do feel strongly that any nomogram should evolve and as newer markers become available and tested, they should be incorporated appropriately.

7. The reason and number of patients excluded from analysis should be in the methods, not the results.

Thank you, this has been incorporated appropriately.

8. The authors said they excluded patient on active surveillance (n=68), this enriches this cohort for high grade disease and together with the fact this cohort comes from a referral center makes their analysis on the finding of "significant cancer" questionable. This should be listed as a limitation. Also the reason for excluding them is not clear. They included others with previous biopsies and many AS protocols include a confirmation biopsy. If there was no plans for biopsy then why get the MRI?

Active surveillance patients were excluded as noted. We do agree that this would enrich the cohort. However, they were excluded for various reasons:

a) The overall aim for the prostate cancer nomogram was to determine, in patients without prostate cancer, the risk of finding any of significant disease at biopsy. As patients undergoing active surveillance previously has positive biopsies, we felt that this cohort was not appropriate for inclusion.

b) The inclusion of active surveillance would have added reader bias from the radiologist. Our protocol for the reporting of active surveillance patients precludes the blinding of clinical details. When reporting such patients, our radiologist was aware of both the clinical details and previous histology, and able to compare the new MRI with these results.

c) Many of the patients on active surveillance did not go on to have biopsy after MRI which is often used as a surveillance tool in its own right. For example, patients with stable PSA at one year would have an MRI and if there were no new changes, would defer repeat biopsy
at that time, thus skewing the availability of a reference standard for these patients. This has been clarified in the text.

9. The use of PI-RADS v1 will likely have readers feeling that the study is antiquated although I agree with the authors discussion about the clinical significance between v1 and v2. If they could update the reads that would be great.

We acknowledge the reviewers’ comments regarding PI-RADS v1. Our comments about PI-RADS v1 vs v2 are included in the Discussion section. We have added additional references demonstrating no difference in diagnostic accuracy between v1 and v2.

As previously stated, this is an evolving process – the nature of any study like this is that when the results are available, newer technologies will have emerged that can be fed back in to improve the process.

10. When discussing their TPSB procedure they need to say "In glands more than 30ml". Also they should explain why additional basal biopsies where taken from these patients. Usually for large prostates and those with BPH the changes are in the transition zone, not the peripheral zone.

We have edited our description of the TPSB procedure, adding additional detail as per the original description as per Vyas et al (referenced). The basal sectors were taken to ensure adequate sampling of the whole peripheral zone and central zone at the base. These were taken via the middle and posterior sectors, to avoid the transitional zone.

11. The authors state they excluded PSA in their multivariable analysis in favor of PSAD. They need to state the reason they picked PSAD over PSA.

PSA was excluded from multivariable analysis as it was highly correlated with PSAD, indicating potential collinearity. As per Reviewer #2, point #5, we did compare our model to one utilising PSA and volume. An analysis of covariance found the model using PSAD had significantly greater reduction in the residual sum of squares (p<0.05), indicating a better fit, for all outcomes.

12. The authors report on a sensitivity analysis for the first 400 patients. They need to state why they did this study for the readers.

As noted we performed a sensitivity analysis for the first 400 patients. After these initial patients, the decision to allow patients to make informed decisions based upon their MRI results was made. We performed the sensitivity analysis to assess for any differences between the first 400 patients vs the overall cohort and identify any bias caused by the decision making process. This has been further clarified in the discussion.

13. The authors report a very high c-index for significant prostate cancer, likely related to selection bias mentioned above. What is not seen at all in the manuscript and needs to be reported is the sensitivity and specifically of each model along with negative and positive predictive values.

Sensitivities, specificities, positive predictive values and negative predictive values have been included as requested.

14. They comment on not using DCE in their MRI index as contrast being a limiting step in either patient administration or image acquisition (not sure which they mean). Reason for this need to be provided as this is not a limiting factor in common practice. Again if they are using PI-RADS v1 this sequence is needed for scoring.
We have revised our paragraph regarding DCE. In the UK, mpMRI use is increasing following the PROMIS study, placing increased pressure on radiology departments. We highlight that B-MRI does offer an alternative with lower acquisition time. We note the accuracy of B-MRI in the response to Reviewer #1, point #1/2.

15. The authors use a 10% cut off in their decision analysis. They should state how or why they came to this number. A decision analysis curve may help provide context.

We have revised the nomogram to range between 5% and >90%. Given that nomograms provide risk estimates for prostate cancer, we selected a lower limit of 5%. This 5% value is cautious and highlights that these predictions are merely estimates as opposed to definitive values. At this level, the nomogram should be adequate for readers and patients to decide whether to proceed to biopsy.
Towards a MRI-based nomogram for the prediction of transperineal prostate biopsy outcome: a physician and patient decision tool

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Keywords: Magnetic Resonance Imaging; Nomogram; Prostate Biopsy; Prostate Cancer; Risk Assessment

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Conflicts of Interest: None disclosed.
Abstract:

Purpose: To develop and internally validate a nomogram utilising biparametric magnetic resonance imaging (B-MRI)-derived variables for the prediction prostate cancer at transperineal sector-guided prostate biopsy (TPSB).

Subjects/Patients and Methods: Consecutive patients referred to our institution with raised PSA, abnormal prostate examination or persistent suspicion of prostate cancer after previous transrectal biopsy between July 2012 and November 2015 were reviewed from a prospective database.

All patients underwent pre-biopsy B-MRI with T2-weighted and diffusion-weighted imaging sequences, followed by 24-40 core TPSB with additional targeted cores using cognitive registration.

Univariable and multivariable logistic regression analysis was used to determine predictors of prostate cancer outcomes. Multivariable coefficients were used to construct two MRI-based nomograms to predict any and significant (Gleason 4 or Maximum Cancer Core Length ≥6mm) prostate cancer at TPSB. Bootstrap resamples were used for internal validation. Accuracy was assessed by calculating the concordance index (c-index).

Results: In total, 615 men were included in the study. Prostate cancer was diagnosed in 317 (51.5%) men with significant cancer diagnosed in 237 (38.5%) men.

Age, PI-RADS score, PSA, PSA Density (PSAD) and Primary Biopsy were predictors of prostate cancer at TPSB on univariable analysis (p<0.0001). PSA showed strong correlation with PSAD and was excluded. The remaining variables were all independent predictors of prostate cancer on multivariable analysis (p<0.0001) and used to generate the nomograms.
Both nomograms showed good discrimination for prostate cancer, with a c-index of 87% for any cancer and 92% for significant disease. Using a nomogram-derived probability threshold of <15%, 111 (18.0%) of biopsies can be saved, at the expense of 3 missed significant prostate cancers.

**Conclusions:** These internally-validated MR-based nomograms were able to accurately predict TPSB outcomes for prostate cancer, especially significant disease. Our findings support the combination of pre-biopsy MRI results and clinical factors as part of the biopsy decision-making process.
Introduction

In current practice, men suspected of harbouring prostate cancer undergo initial transrectal prostate biopsy based upon abnormal digital rectal examination (DRE) or raised prostate specific antigen (PSA). Of these biopsies, only 40% will be positive, and detection rates of subsequent biopsies fall below 20% [1-4].

Multiple factors underlie this low detection rate. Serum PSA is used for screening, but, by itself cannot accurately distinguish between benign and malignant conditions, contributing to poor detection rates [5]. Furthermore, a proportion of cancers detected through PSA screening will be small volume, low-risk disease that may not require treatment [5].

Transrectal biopsies also suffer from anatomical limitations. Random needle biopsy often leads to sampling error due to heterogeneity within prostate tumours, resulting in misdiagnosis [6]. As the prostate is approached posteriorly via the rectum, specific anatomical areas are undersampled. The anterior aspect of the prostate is challenging to biopsy, while the detection of apical tumours is limited by the needle angle attainable through the rectum [7, 8].

In addition to poor detection rates, men undergoing transrectal biopsy are placed at risk of biopsy-associated morbidity. The Prostate Biopsy Effects (ProBE) study, nested within the Prostate Cancer for Testing and Treatment (ProtecT) study, gives insight into this; pain is reported by 44%, fever by 18% and haematuria by 66%, with 1.3% requiring hospital admission [9]. One-fifth of men would consider repeat biopsies a moderate to major problem [9]. Improvements to current prostate cancer risk assessments would reduce unnecessary biopsies and morbidity.
The prostate biopsy decision-making process is complex. Tools that provide accurate risk analysis can aid the clinician and patient when considering whether to undertake biopsy. Previously reported biopsy nomograms, based on clinical variables have demonstrated predictive accuracies of only 65-77% [10, 11]. Novel diagnostic parameters can improve on the low PSA sensitivity and optimise the predictive accuracy of these nomograms. As magnetic resonance imaging (MRI) technology has improved through the use of additional sequences, including diffusion-weighted imaging (DWI) and dynamic contrast enhancement (DCE), it is increasingly being used within the diagnostic pathway for prostate cancer [12-14]. The recent Prostate MRI Imaging Study (PROMIS) confirmed multiparametric MRI (T2-weighted, DWI, DCE) to be a useful triage test, with greater sensitivity (93%) as compared to transrectal biopsy (48%) [15]. Since the results of this trial were released, the role of pre-biopsy MRI has been increasing within the United Kingdom, placing increased pressure on radiology departments [16]. Biparametric MRI (B-MRI) represents a valid alternative to mpMRI, with its shorter acquisition times, and we have validated its accuracy previously [13].

Furthermore, PSA density (PSAD) has had potential to improve serum PSA specificity whilst preserving sensitivity. Despite this, use in clinical practice is limited and poor uptake may be related to the use of transrectal ultrasound for prostate volume estimation and transrectal biopsy as the reference standard, two techniques subject to human error and inaccuracy [7, 8, 17]. Separate volume assessments are expensive, inconvenient and uncomfortable for patients. With pre-biopsy MRI, prostate volume can be accurately and conveniently assessed prior to biopsy [18].
In the present study, we aimed to develop a new nomogram, utilising both clinical and biparametric MRI (B-MRI) data for the predicting the presence of cancer on prostate biopsy.
**Patients and Methods**

The study was approved by the local governance committee as a prospective audit and conforms to the Standards for the Reporting of Diagnostic Accuracy (STARD).

**Patient Population**

Consecutive patients referred to our institution for biparametric MRI (B-MRI) and transperineal sector-guided prostate biopsy (TPSB) between July 2012 and November 2015 were reviewed from a prospectively-collected database. Referral reasons included raised PSA, abnormal DRE, or persistent suspicion of prostate cancer after previous transrectal biopsy. Patients excluded included: those unable to undergo B-MRI (n=29), B-MRI older than 6 months before biopsy (n=38) or undertaken at other institutions (n=67) and those undergoing active surveillance (n=68).

**Biparametric MRI (B-MRI)**

All patients underwent B-MRI prior to biopsy and at least 6 months after previous biopsy or prostate instrumentation to avoid haemorrhage artefact. B-MRI was undertaken using a 1.5 Tesla machine (Signa Excite, GE Healthcare, Little Chalfont, UK) and 8-channel phased array body coils. Protocol included axial oblique, sagittal and coronal T2-weighted imaging and axial DWI (Supplementary Table 1). DWI sequences were obtained with b-values of 0, 700 and 1000 in 18 patients, 0, 1000 and 1400 in 26 patients and 0 and 1400 in 571 patients. High b-value images and apparent diffusion coefficient maps were used for analysis.

Reporting was performed by a single uroradiologist (S.H.L.) with 5 years prostate MRI experience and blinded to clinical details. Images were reported using Prostate Imaging
Reporting and Data System (PI-RADS) v1, indicating likelihood of malignancy from 1 to 5 [19]. PI-RADS reporting at our institution has been previously described [13]. Analysis was based upon the highest overall score reported for either B-MRI sequence (T2 or DWI). Final report did not influence the decision to proceed to biopsy for the initial 400 cases.

Subsequent patients with negative MRIs, i.e. PI-RADS 1 or 2, were given the option to proceed to biopsy.

Prostate volume was determined with T2-weighted images, using the ellipsoid method, defined as $\pi/6 \times \text{length} \times \text{width} \times \text{height}$. PSAD was defined as total serum PSA divided by prostate volume.

_Psage_ Prostate Biopsy

TPSB was performed as previously described [20, 21], preferentially targeting the peripheral zone. The PZ was divided into anterior, mid and posterior sectors for each lobe, and the mid and posterior sectors occupying the PZ posterior to the prostatic urethra, verumontanum and bladder neck. In glands <30ml, four cores were taken from each sector, yielding 24 cores. In glands >30ml, basal biopsies were taken via the mid and posterior sectors to sample the central zone at the base (on either side of the midline), whilst avoiding the TZ. In prostates >50ml, 38 cores were taken: 5 per sector, with 8 basal biopsies. In patients with suspicious MRI lesions, a further 2-4 cores were taken using cognitive registration. The use of TPSB provided a comprehensive reference standard, allowing for accurate prostate cancer diagnosis.

Two outcomes were assessed, any cancer and significant prostate cancer, defined as:

*presence of Gleason pattern 4 or maximum cancer core length (MCCL) of $\geq 6mm$. This core*
length was chosen as it corresponded to a lesion volume of 0.5ml, below the calculated threshold of 1.3ml for significant tumour volume [14].

Data Analysis

Data analysis was performed using R version 3.1.2 (R Foundation for Statistical Computing, Vienna, Austria). Patient characteristics were compared using chi-squared tests and independent sample t-test for categorical and continuous variables, respectively. Odds ratios (OR) and 95% confidence intervals (95% CI) for any and significant prostate cancer were estimated by univariable logistic regression analysis assessing age, PSA, PSAD, PI-RADS score and primary biopsy as potential predictors. Multivariable logistic regression analysis including all potential predictors was used to determine independent predictors of prostate cancer outcomes. A further sensitivity analysis was performed including only the first 400 patients, as subsequent MRI results were used in the decision-making process in subsequent patients.

A nomogram was developed using the rms package based on the final logistic regression model to estimate the risk of being diagnosed with each prostate cancer outcome. Performance was evaluated through internal validation, performed by generating 200 bootstrap samples. Nomogram discrimination was assessed by calculating the concordance index (c-index) from the bootstrap samples. A value of 0.5 indicated that 50% of the patients were correctly classified. Calibration curves were generated for the actual and bootstrap samples, demonstrating prediction model accuracy. Various nomogram probability cut-offs were tested to assess the ability to identify patients with and without prostate cancer.
Results

From July 2012 to November 2015, 817 men undergoing TPSB were considered for the study. After exclusions, 615 patients were included, including: patients undergoing primary biopsy for raised PSA or abnormal DRE (n=484) and patients with previous negative transrectal prostate biopsy and ongoing suspicion of prostate cancer (n=131).

Patient characteristics are shown in Table 1 after stratification according to presence and absence of prostate cancer. Overall, 51.5% of the men (n=317) were diagnosed with prostate cancer. Of these, 38.5% (n=237) had significant prostate cancer. 28% (n=175) of patients had negative MRIs (PI-RADS 1 or 2), whereas 51% (n=312) were deemed positive (PI-RADS 4 or 5). In total, 34 (5.5%) patients suffered complications from TPSB, consisting of: 23 (3.7%) episodes of urinary retention, 8 (1.3%) patients with significant haematuria with clots, 1 (0.2%) case of wound bleeding, 1 (0.2%) prostatitis, and 1 (0.2%) patient with SVT following general anaesthetic.

On univariable analysis (Table 2), age, PSA, PSAD, PI-RADS score and primary biopsy were all significant predictors of any and significant prostate cancer at TPSB. A strong correlation between PSA and PSAD density was noted (r=0.81), indicating potential collinearity; therefore, PSA was excluded from multivariable analysis. Multivariable logistic regression analysis (Table 2) showed the remaining variables remained independent predictors of both outcomes.

After the first 400 patients, patients were given the informed choice to proceed to biopsy based upon the B-MRI results. Therefore, a further sensitivity analysis was performed,
including only the first 400 patients. Similar albeit slightly weaker findings were observed with all associations remaining statistically significant apart from age (data not displayed).

Using age, PSAD, PI-RADS score and primary biopsy, a nomogram was developed to predict any prostate cancer, as shown in Figure 1. An additional nomogram for the prediction of significant prostate cancer is demonstrated in Figure 2. We ran an additional multivariable analysis for each outcome, using PSA and volume separately. We compared the fit between this and the PSAD model, using analysis of covariance and found that the model using PSAD had significantly greater reduction in the residual sum of squares (p<0.05), indicating a better fit, for all outcomes. Therefore the final model was performed using PSAD.

The nomograms were internally validated using 200 bootstrap samples; calibration curves are shown in Figure 1(b) for any prostate cancer and Figure 2(b) for significant prostate cancer. The c-index for the any prostate cancer nomogram was 0.87 (95% CI 0.84-0.90). For significant prostate cancer, the c-index was 0.92 (95% CI 0.89-0.94).

Finally, Table 3 shows the numbers of biopsies performed and the number of missed prostate cancers at various ‘any cancer’ probability scores as calculated by our MR-based any prostate cancer nomogram. Sensitivity, specificity, positive predictive values and negative predictive values for significant cancer are also demonstrated at each cut-off value. By utilising this calculator, at a calculated probability cut-off of 5%, 23 (3.7%) biopsies would be avoided, at the expense of 1 missed significant prostate cancers. At a cut-off as high as 30%, 213 (34.6%) biopsies would be avoided, with 6 missed significant prostate cancers.
Discussion

Nomograms for prostate cancer risk assessment based on clinical, laboratory and ultrasound parameters have attempted to improve detection rates and reduce unnecessary biopsies. However, accuracy has been limited and additional prognostic variables can improve their performance [10, 11]. We present nomograms, incorporating MRI-derived and clinical information for the prediction of prostate cancer on TPSB. Internal validation showed strong discrimination with a c-index of 0.87 for any cancer and 0.92 for significant prostate cancer.

The use of MRI has progressed from staging to detection and localisation of tumours [12]. The addition of functional sequences has led to accurate prostate cancer localisation, which can be used in risk stratification before biopsy and studies have shown the sensitivity of mpMRI to be over 90% [12, 15]. The role of prostate MRI has been increasing within the United Kingdom, placing demand on radiology departments, due to the long acquisition times, particularly for DCE sequences [16]. Our MRI index test uses two straightforward, resource-friendly sequences: T2-weighted and DWI sequences. These sequences are widely available, and the lack of DCE reduces sequence times, representing a practical solution to many hospitals.

We have previously validated our B-MRI protocol with the use of PI-RADS scoring [13]; when PI-RADS 1 and 2 lesions were deemed negative, B-MRI had a sensitivity of 97% and specificity of 60% for significant prostate cancer (Gleason pattern 4 or MCCL ≥6mm). These compare favourably to the values reported by the PROMIS study. Ahmed et al. reported mpMRI images using a validated Likert system; again, when Likert 1 and 2 lesions were negative, the authors demonstrated a sensitivity of 87% and specificity of 47% for significant prostate cancer (UCL definition 1: Gleason ≥3+4 or MCCL ≥4mm) [15]. Furthermore, a recent
study by Thstrup et al. [22] showed similar prostate cancer detection rates between B-MRI (sensitivity 0.94-0.96) vs mpMRI (sensitivity 0.93-1.0) across 2 independent readers.

Pre-biopsy imaging also lends itself well to MRI-derived PSAD, which has been demonstrated to more accurately predict TPSB outcomes compared to PSA alone [18, 23]. Given that certain prostate cancers are MRI invisible, inclusion of PSAD within our nomogram aids in the identification of patients harbouring MRI-invisible cancers.

For our reference standard, we used TPSB with its high overall detection rate [13, 24]. This arises from the approach used: by entering the prostate through the skin, systematic investigation of the prostate may be achieved, including the anterior and apical aspects, two areas which are difficult to sample through the transrectal approach [7, 8]. We aimed to provide comprehensive a reference standard as possible, and short of radical prostatectomy, our view is that TPSB provides this. Cognitive-guided cores were included, which are widely available to clinicians and avoid the variability associated with different fusion biopsy systems [25].

These nomograms for prostate cancer may prove to be a valuable patient and clinician decision making tool, potentially avoiding unnecessary biopsies and reducing adverse events, through improved prostate cancer prediction. At a probability cut-off of 5% for any prostate cancer, 27 out of 615 biopsies could be avoided, at the expense of 4 missed prostate cancers, of which 1 is significant. At 10%, this increases to 65 saved biopsies and 11 missed prostate cancers, 2 of which are significant. The acceptability of these missed cancers is open to debate, given the morbidity associated with prostate biopsy.
Table 4 gives an example case of a 60-year old man undergoing primary biopsy. Risk of significant and any (in parentheses) prostate cancer is shown and stratified into categories: low (<5%), medium (5-15%) and high risk (>15%). This example highlights the utility of both MRI and PSAD in establishing prostate cancer risk. Patients falling into lower risk categories may choose to avoid prostate biopsy. These men would continue to undergo surveillance through repeat PSA tests, only proceeding to biopsy when a chosen PSA threshold was reached.

Previous studies have looked at the use of both mpMRI and PSAD for the prediction of prostate cancer. Van Leeuwen et al. [26] developed a predictive model from 393 patients for the prediction of prostate cancer utilising similar variables including PI-RADS v1, with a similar transperineal biopsy reference standard (median 30 cores). The study benefits from a multicentre cohort and retrospective external validation, but demonstrates slightly lower accuracy to our model (AUC 0.80 for any cancer, 0.88 for significant cancer). This may arise from the differing biopsy strategy used during validation (transrectal or transperineal, median 18-cores).

Distler et al. [27] studied the value of adding PSAD to pre-biopsy mpMRI in 1040 men undergoing transperineal biopsy, developing a nomogram utilising these two variables. Accuracy of the nomogram was lower (AUC 0.79) than that seen in this study. In a further study by Radtke et al, of the same study group, risk models for prostate cancer were developed utilising clinical (age, DRE, PSA, volume) and PI-RADS v1 scores [28]. The authors reported good accuracy with 0.83 ROC area under the curve. It is interesting to note that within the latest risk model, the PI-RADS score accounts for fewer overall points as
compared to our model. Both the Van Leeuwen and Radtke nomograms, as well as the present study require prospective external validation.

In a separate large study, Wang et al. [29] examined the value of pre-biopsy MRI in a cohort of 985 Chinese patients, developing a nomogram utilising mpMRI and clinical information for the prediction of prostate cancer over a 2-year follow-up period. This was internally validated showing greater accuracy (AUC 0.938). Despite this, a major shortcoming in the study was the reference standard, which varied considerably, including transrectal biopsy, transurethral resection of prostate (TURP) and radical prostatectomy specimens. A further 101 patients were diagnosed clinically, with no available histology. Given that patients were diagnosed over a subsequent two year period, it would be difficult to apply this nomogram to clinical practice, and is not directly comparable to the current study.

In addition to MRI, the ongoing search and development of novel prostate cancer biomarkers has been documented and could aid the development of new nomograms. These markers, including the four kallikrein panel (4K), prostate health index (PHI, a panel of total PSA, free PSA and [-2]proPSA) and urinary prostate cancer gene 3 (PCA3) have been integrated into clinical nomograms to improve their accuracy, with accuracy reported in the range 0.73-0.80 on ROC analysis [30-32]. These assays have the potential to supplant PSA and PSAD, and how they will integrate into the diagnostic pathway along with MRI remains to be determined. However, we note that these assays are not freely available in all centres, including in the United Kingdom, from which this study group arises.

The strengths of the present study include the fact it is a large, prospective study, involving 615 patients. MRI was reported prospectively by a single uroradiologist, blinded to clinical details, avoiding reporter bias. Our study uses TPSB as a reference standard, interrogating all
sectors of the prostate and avoiding the inherent sampling error associated with transrectal biopsies.

MRI reporting used PI-RADS v1. PI-RADS has changed since initial publication; in v1, T2, DWI and DCE sequences were considered equivalent, whereas in v2, DCE became a secondary sequence, used to delineate indeterminate PI-RADS 3 lesions [33]. This highlights the importance of the two primary sequences used in this study. Furthermore, studies have shown similar accuracy between both v1 and v2 [34, 35].

We note potential sources of bias. Firstly, the ellipsoid method was used for the calculation of prostate volume. For the purposes of this nomogram, our aim was to create a user friendly prediction tool with straightforward calculations. Formulaic prostate volume calculation is quick, requires little training and does not require further software, making data input very accessible. We have found that PSAD utilising this volume calculation method provides an improvement in the prediction of TPSB outcomes over total PSA alone [23].

While clinical details were blinded during MRI reporting, the study is prone to a learning curve, as MRI scans were re-reviewed with histology results. However, our published B-MRI series found little learning curve effect [13]. For this reason, active surveillance patients were excluded from our cohort. Our protocol for the reporting of active surveillance patients precludes the blinding of clinical details. When reporting such patients, our radiologist was both aware of the clinical details and previous histology, and able to compare new MRIs with previous imaging. In addition, many patients undergoing active surveillance did not go on to have biopsy after MRI; for example, patients with stable PSA
and negative MRI may have biopsies deferred, skewing the reference standard availability in this set of patients.

Furthermore, after the initial 400 cases in our cohort, patients were given the choice to proceed to TPSB, based on MRI results, potentially adding bias. We performed a sensitivity analysis to assess for any variations between the initial 400 patients versus the entire cohort. Results were similar, albeit with slightly weaker associations, when limiting analyses to these first 400 patients. The influence of informed patient decision making was small, represents current urological practice and is justifiable based on our initial results.

Finally, this nomogram requires external validation to be used in different cohorts. Our population is predominantly Caucasian, presenting with rising PSA and lower urinary tract symptoms; therefore, benign patients often have low PSAD due to increased prostate volume. It may not be representative of populations at other institutions, which may include screened patients with lower PSA values and prostate volumes.
Conclusions

We demonstrate an MRI-derived nomogram for the prediction of TPSB outcomes. Our nomogram shows good discrimination for any (87%) and significant (92%) prostate cancer on internal validation and supports the combination of pre-biopsy MRI findings and clinical risk factors. It can be used in clinic for patient counselling as part of the biopsy decision-making process.
References


Figure Legends

Figure 1: (a) MR-based Nomogram predicting any prostate cancer on transperineal prostate biopsy. Instructions: To obtain nomogram-predicted probability of prostate cancer (PCa), patient values are located for each variable. A vertical line is drawn to the ‘Points’ axis to determine the number of points attributed to each variable. The points are added up for all variables. The sum is located on the ‘Total Points’ line to show the individual’s probability of cancer on transperineal prostate biopsy on the ‘PCa probability’ line; (b) calibration curve of MR-based nomogram. Perfect prediction corresponds to the dashed 45° line. Points estimated above the 45° line represent nomogram underprediction, whereas points below the 45° line correspond to nomogram overprediction. The nonparametric, smoothed dotted curve represents the relationship between predicted probability and observed frequency of PCa at transperineal prostate biopsy across the entire cohort. The solid curve was bias-corrected by bootstrapping using 200 resamples.

Figure 2: (a) MR-based Nomogram for predicting significant prostate cancer (Gleason 4 or 3+3 (MCCL ≥6mm)) on transperineal prostate biopsy; (b) calibration curve of MR-based nomogram for significant prostate cancer. Instructions as per Figure 1.
Figure 1a
Click here to download high resolution image
Figure 1b

B = 200 repetitions, boot

Mean absolute error = 0.026 n = 615
Figure 2b

B = 200 repetitions, boot
Mean absolute error = 0.009 n = 614
Table 1: Patient Characteristics

<table>
<thead>
<tr>
<th></th>
<th>All patients (n=615)</th>
<th>Prostate Cancer (n=317)</th>
<th>No Cancer (n=298)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age, Mean ± SD</strong></td>
<td>65.4 ± 6.9</td>
<td>66.7 ± 6.9</td>
<td>64.1 ± 6.5</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><strong>PSA, Mean ± SD</strong></td>
<td>15.4 ± 39.2</td>
<td>20.6 ± 53.4</td>
<td>9.9 ± 9.2</td>
<td>0.0005</td>
</tr>
<tr>
<td><strong>PSA Density, Mean ± SD</strong></td>
<td>0.37 ± 2.03</td>
<td>0.58 ± 2.81</td>
<td>0.15 ± 0.14</td>
<td>0.0088</td>
</tr>
<tr>
<td><strong>Primary Biopsy</strong></td>
<td>484 (79%)</td>
<td>277 (87%)</td>
<td>207 (69%)</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>
### Table 2: Univariable and Multivariable Associations of Predictor Variables for Any and Significant Prostate Cancer

#### Univariable Analysis

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Any Prostate Cancer</th>
<th>Significant Prostate Cancer</th>
<th>P</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>1.06 (1.03-1.09)</td>
<td>1.07 (1.04-1.10)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td><strong>PSA</strong></td>
<td>1.04 (1.02-1.06)</td>
<td>1.06 (1.04-1.09)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td><strong>PSAD</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;0.10</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.10-0.16</td>
<td>2.11 (1.30-3.46)</td>
<td>3.75 (1.99-7.44)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>0.16-0.28</td>
<td>5.19 (3.20-8.57)</td>
<td>9.01 (4.91-17.59)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>≥0.28</td>
<td>10.37 (6.21-17.79)</td>
<td>24.22 (13.01-48.07)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td><strong>PI-RADS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>-</td>
<td></td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>2</td>
<td>1.69 (0.64-5.29)</td>
<td>1.00 (0.33-3.85)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>3</td>
<td>3.09 (1.22-9.51)</td>
<td>6.00 (1.17-109.96)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>4</td>
<td>12.16 (4.83-37.36)</td>
<td>38.56 (7.97-694.81)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>5</td>
<td>79.58 (29.97-256.39)</td>
<td>179.23 (37.17-3230.92)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
</tbody>
</table>

#### Primary Biopsy

<table>
<thead>
<tr>
<th></th>
<th>Any Prostate Cancer</th>
<th>Significant Prostate Cancer</th>
<th>P</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>3.04 (2.03-4.64)</td>
<td>3.75 (2.35-6.22)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
</tbody>
</table>

#### Multivariable Analysis

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Any Prostate Cancer</th>
<th>Significant Prostate Cancer</th>
<th>P</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>1.51 (1.13-2.01)</td>
<td>1.05 (1.02-1.09)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td><strong>PSAD</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;0.10</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.10-0.16</td>
<td>1.87 (1.05-3.36)</td>
<td>3.57 (1.66-8.02)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>0.16-0.28</td>
<td>3.17 (1.74-5.77)</td>
<td>5.81 (2.76-12.42)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>≥0.28</td>
<td>4.26 (2.22-8.16)</td>
<td>12.26 (5.71-27.80)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td><strong>PI-RADS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>-</td>
<td></td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>2</td>
<td>1.59 (0.54-4.61)</td>
<td>1.90 (0.29-37.38)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>3</td>
<td>2.84 (0.99-8.13)</td>
<td>5.61 (1.01-105.78)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>4</td>
<td>8.38 (2.94-23.90)</td>
<td>25.47 (4.87-471.35)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>5</td>
<td>43.12 (14.34-129.72)</td>
<td>91.52 (17.51-1694.02)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
</tbody>
</table>

#### Primary Biopsy

<table>
<thead>
<tr>
<th></th>
<th>Any Prostate Cancer</th>
<th>Significant Prostate Cancer</th>
<th>P</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>2.16 (1.28-3.68)</td>
<td>2.68 (1.42-5.20)</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
</tr>
</tbody>
</table>
Table 3: Numbers of biopsies performed and missed prostate cancer (any or significant) according to different MR-based nomogram-derived probability cut-offs

<table>
<thead>
<tr>
<th>Probability (any cancer) cut-off (%)</th>
<th>Biopsies performed, n (%)</th>
<th>Biopsies not performed, n (%)</th>
<th>Any cancer missed, n</th>
<th>Significant cancer missed, n</th>
<th>For significant prostate cancer:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Sensitivity (%)</td>
</tr>
<tr>
<td>5</td>
<td>588/615 (95.6)</td>
<td>27 (3.7)</td>
<td>4</td>
<td>1</td>
<td>99.6</td>
</tr>
<tr>
<td>10</td>
<td>550/615 (89.4)</td>
<td>65 (10.6)</td>
<td>11</td>
<td>2</td>
<td>99.2</td>
</tr>
<tr>
<td>15</td>
<td>504/615 (82.0)</td>
<td>111 (18.0)</td>
<td>20</td>
<td>3</td>
<td>98.7</td>
</tr>
<tr>
<td>20</td>
<td>456/615 (74.1)</td>
<td>159 (25.9)</td>
<td>27</td>
<td>4</td>
<td>98.3</td>
</tr>
<tr>
<td>25</td>
<td>435/615 (70.7)</td>
<td>180 (29.3)</td>
<td>30</td>
<td>5</td>
<td>97.9</td>
</tr>
<tr>
<td>30</td>
<td>402/615 (65.4)</td>
<td>213 (34.6)</td>
<td>37</td>
<td>6</td>
<td>97.5</td>
</tr>
</tbody>
</table>
Table 4: Percentage risk of significant and any (parentheses) prostate cancer (Gleason 4 or ≥6mm MCCL) in a 60 year old man with no previous prostate biopsy

<table>
<thead>
<tr>
<th>PI-RADS</th>
<th>60 years old</th>
<th>PSA Density</th>
<th>&lt;0.10</th>
<th>0.10-0.16</th>
<th>0.16-0.28</th>
<th>&gt;0.28</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>&lt;5% (&lt;5%)</td>
<td>&lt;5% (6%)</td>
<td>&lt;5% (11%)</td>
<td>&lt;5% (14%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>&lt;5% (9%)</td>
<td>&lt;5% (15%)</td>
<td>7% (25%)</td>
<td>13% (31%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>&lt;5% (21%)</td>
<td>14% (33%)</td>
<td>20% (47%)</td>
<td>35% (55%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>14% (42%)</td>
<td>36% (57%)</td>
<td>48% (71%)</td>
<td>66% (77%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>37% (67%)</td>
<td>67% (78%)</td>
<td>77% (87%)</td>
<td>88% (&gt;90%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
To the editor,

We wish to re-submit our manuscript entitled “Towards a MRI-based nomogram for the prediction of transperineal prostate biopsy outcome: a physician and patient decision tool” for consideration in Urologic Oncology: Seminars and Original Investigations.

We would like to you thank you and the reviewers for the valuable comments and feedback which have allowed us to improve our manuscript. I have attached a detailed list of the changes that we have made and responses to each of the reviewers’ comments.

We confirm that this work is original and has not been published elsewhere nor is it currently under consideration for publication elsewhere. We declare no conflicts of interest.

All authors have read and approved the manuscript.

Thank you for your consideration of this manuscript,

Sincerely,

Su-Min Lee
(Corresponding Author)
Email: smlee84@gmail.com
Telephone: (+44) 7533 488525