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INNOVATIONS IN PROSTATE CANCER SPECIAL FEATURE: COMMENTARY

The challenge of prostate biopsy guidance in the era of mpMRI detected lesion: ultrasound-guided versus in-bore biopsy

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ABSTRACT

The current recommendation in patients with a clinical suspicion for prostate cancer is to perform systematic biopsies extended with targeted biopsies, depending on mpMRI results. Following a positive mpMRI [i.e. Prostate Imaging Reporting and Data System (PI-RADS) ≥3], three targeted biopsy approaches can be performed: visual registration of the MRI images with real-time ultrasound imaging; software-assisted fusion of the MRI images and real-time ultrasound images, and in-bore biopsy within the MR scanner. This collaborative review discusses the advantages and disadvantages of each targeting approach and elaborates on future developments. Cancer detection rates seem to mostly depend on practitioner experience and selection criteria (biopsy naïve, previous negative biopsy, prostate-specific antigen (PSA) selection criteria, presence of a lesion on MRI), and to a lesser extent dependent on biopsy technique. There is no clear consensus on the optimal targeting approach. The choice of technique depends on local experience and availability of equipment, individual patient characteristics, and onsite cost-benefit analysis. Innovations in imaging techniques and software-based algorithms may lead to further improvements in this field.

INTRODUCTION

Currently, there is level 1A evidence that the addition of mpMRI and subsequent MRI-targeted biopsy (TBx) to the diagnostic pathway of prostate cancer (PCa) increases the detection of clinically significant prostate cancer (csPCa) compared to systemic biopsy (SBx) alone.1,2 In case of a positive mpMRI (PI-RADS ≥3), the combination of both TBx and SBx finds significantly more csPCa, than either approach on its own, especially in biopsy naïve patients.1,2 Consequently, current guidelines strongly recommend a combined biopsy approach in this patient population. In the previous negative biopsy patient population, the additional value of SBx is less obvious, and a TBx only approach is recommended.3

The additional value of SBx should be carefully considered, since omitting SBx has the benefit of finding less clinically insignificant prostate cancer (ISUP Grade 1). In the Cochrane meta-analysis, systematic biopsy detected an additional 9.8% (8.0–11.8%) ISUP Grade 1 PCa compared to the MRI-pathway.4

In case of a negative mpMRI, the practitioner has a choice to omit biopsy or to perform only SBx. MRI is reported to have a negative predictive value of >90%; however, it still misses approximately 9% of ISUP ≥2 PCa.1,5 Consequently, omitting biopsy carries the risk of missing csPCa, whereas performing SBx carries the risk of finding clinically insignificant PCa. The choice to omit or perform SBx in this patient population should be based on the level of clinical suspicion and shared decision making.4 The clinical suspicion can be judged using tools, such as the “prostate cancer-risk calculator”, a tool that weights MRI results, volume, previous negative biopsies, digital rectal exam, PSA level and age.6
Two of the authors (BRJ, BM) have previously published a review article on imaging guidance technology for prostate cancer (prostate cancer: a review article on imaging guidance technology for prostate cancer, Br J Radiol, 2015). Subsequently, articles on the subject have appeared in the literature. MRI-targeted biopsy in the form of magnetic resonance imaging (MRI) is a technique that has been used to improve diagnostic accuracy and to target biopsy samples more efficiently.

mpMRI followed by systematic and/or targeted biopsy is the currently recommended diagnostic pathway for prostate cancer detection. A current topic of discussion is which technique to use when performing MRI-targeted biopsies. Following a positive mpMRI, a practitioner can choose between three targeted biopsy techniques: in-bore MRI-guided biopsy (in-bore biopsy), ultrasound-guided fusion-targeted biopsy (fusion-TBx) or ultrasound-guided cognitive-targeted biopsy (cognitive-TBx). These three techniques show different features according to their technical characteristics (Table 1).

### Table 1. Comparison of techniques for MRI targeted prostate biopsies

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Cognitive-TBx</th>
<th>Fusion-TBx</th>
<th>MRGB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ease of use</td>
<td>1</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Time-efficient</td>
<td>1</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Real-time feedback</td>
<td>1</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Targeting inaccuracies due to prostate deformation</td>
<td>2</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Lesion directly visible on guidance imaging</td>
<td>3</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Accurate lesion &lt;10 mm targeting</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Steep learning curve</td>
<td>2</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Cost-effectiveness</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>MRI imaging registration</td>
<td>Visual</td>
<td>Software assisted</td>
<td>Visual</td>
</tr>
<tr>
<td>Possibility for systematic biopsies</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Use by urologist</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Robotic approach</td>
<td>Not possible</td>
<td>Possible</td>
<td>Possible</td>
</tr>
</tbody>
</table>

Parameters are scored on a 5-point Likert scale: 1. Extremely applicable, 2. Very applicable 3. Moderately applicable, 4. Not very applicable 5. Not at all applicable.

Scoring results are based on a combination of referred to literature and the authors’ opinion and experience.

![Figure 1](a) Robotic MRI in-bore-targeted biopsy (SoteriaTM). 62-year-old male with a PSA value of 16.6 ng/ml\(^{-1}\). Axial T2 WI shows a 6 mm focal, PIRADS four lesion on the right peripheral zone (arrow). (b) The desired and planned target for the nodule is shown with restricted diffusion on DWI image (arrow). (c) Oblique axial Fast Imaging Employing Steady-State Acquisition (FIESTA) sequence shows the current position of the needle guide, represented by the orange line overlay from the needle guide with the optimal path after the remote movement of the guide. The needle track and sample core are represented by red line. (d) After the desired target for biopsy has been selected, the robotic device (arrow) is moved remotely to the target as shown (c); the table is moved out of the bore and a biopsy sample is taken with a standard compatible biopsy gun. Biopsy histopathology results showed a Gleason Score 4 + 3 = 7 adenocarcinoma.
prostate biopsy can also be performed using a transperineal or transgluteal approach. The transrectal procedure requires antibiotic prophylaxis without the need for local anaesthesia, for the transperineal or transgluteal approach local anaesthesia (or sedation) is necessary, but antibiotic prophylaxis can be omitted. The transgluteal approach is typically reserved for patients without rectal access and is rarely used in clinical practice.15

Ultrasound-guided biopsies

Cognitive- vs fusion-targeted biopsy

Ultrasound-guided targeted biopsies can be performed using cognitive guidance (cognitive-TBx) or with MRI-TRUS fusion software and navigation systems (fusion-TBx).

In cognitive-TBx, the practitioner visually registers the MRI lesion and uses the prostate anatomy to target this lesion on real-time US imaging (Figure 2). In fusion-TBx, the practitioner uses dedicated software to superimpose suspicious lesions detected on MRI over real-time US images and uses one of the available MRI-US fusion systems to navigate to the lesion (Figure 3). These software-based fusion platforms differ in the image registration and tracking methods. Image registration can be performed using either elastic or rigid fusion methods. It has been suggested that elastic fusion is more accurate because it accounts for prostate deformation, where rigid fusion does not. However, a meta-analysis evaluating these methods showed no difference in csPCa detection rates.16 Some platforms are available for either the transrectal and transperineal approach, others for both.

Transrectal vs transperineal approach

The prostate can be approached through either the rectal wall or the perineum. Transrectal ultrasound (US)-guided biopsy (TRUS-GB) is the standard of care for obtaining histopathological confirmation of prostate cancer. The transperineal approach has been gaining attention as a result of the associated low infection risk compared to the transrectal approach. A recent meta-analysis confirmed this association, showing a significant decrease in infectious
Complications, with a relative risk of 0.55 (95% CI 0.33–0.92).\textsuperscript{17} Consequently, the 2021 EAU guideline now strongly recommends using the transperineal approach.\textsuperscript{4} Additionally, transperineal biopsies are reported to lead to superior sampling of the apex as well as the anterior prostate. If this results in higher cancer detection rates (CDR), however, remains to be proven.\textsuperscript{18}

**DISCUSSION**

Comparing targeted biopsy techniques

**Diagnostic accuracy**

First, it is important to look at the diagnostic accuracy. A meta-analysis comparing the three techniques, which included 43 studies with a total of 2497 patients found no significant advantage of any particular technique for csPCa detection.\textsuperscript{19} Another meta-analysis, comparing only fusion- and cognitive-TBx using 1714 men from nine studies, concluded there was a trend toward improved csPCa detection rates for fusion-TBx; however, this was not statistically significant.\textsuperscript{20} Because the studies included in both meta-analyses had small sample sizes and were subject to significant heterogeneity, the results need to be interpreted with caution. In a more recent study, Costa et al evaluated biopsy results from a mixed (prior negative and biopsy naive) cohort, with MRBG (n = 103) performing significantly better than TBx combined with SBx (N = 300), reporting csPCa detection rates of 61 and 47%, respectively.\textsuperscript{21} The same group, using a partly overlapping cohort, found significantly more Grade Group upgrading when matching radical prostatectomy histopathology to biopsy results in the TBx/SBx group (27.7%) compared to MRBG (13%).\textsuperscript{22} The results of these studies show a clear advantage in diagnostic accuracy for MRGB. However, it is difficult to say if these results can be extrapolated to general practice, considering both trials originated from a single centre and have a non-randomised and retrospective design.

In the prior negative biopsy patient population, two randomised-controlled trials evaluated detection rates of the different biopsy techniques. In the FUTURE trial, 234 men with a prior negative biopsy result and a positive mpMRI (PIRADS ≥3) were randomised to cognitive-TBx, fusion TBx or MRGB. This study found no significant differences in diagnostic accuracy, with csPCa detection rates of 33.3%, 34.2 and 33.3%, respectively.\textsuperscript{23} Arsov et al\textsuperscript{18} randomised 210 patients and also did not find a significant difference, reporting csPCa detection rates of 29% for MRRBG and 26% for fusion-TBx.

**Operator experience**

A factor that does seem to be of importance is the experience of the operator performing a certain biopsy technique, being an independent predictor for csPCa detection in SBx as well as TBx with odds ratios up to 2.40.\textsuperscript{24}

**Possibility for systematic biopsies**

Another factor to take into account is whether or not SBx is indicated, considering that MRGB does not support a systematic biopsy approach. This may be an issue in biopsy naive patients where SBx leads to significantly more csPCa detection (4.3–5.0%).\textsuperscript{1}

Cost-effectiveness

When looking at the financial aspects, multiple studies have shown cost-effectiveness of the MRI pathway (MRI followed by targeted biopsy) compared to SBx.\textsuperscript{10–12} When comparing the three targeting techniques, fusion TBx and MRGB are both more expensive than cognitive-TBx, since they require additional planning, guidance material and time.\textsuperscript{13} Nevertheless, all three techniques provide a cost-effective strategy, because they improve disease stratification and facilitate more appropriate patient selection for prostate biopsies.\textsuperscript{15} Better patient selection avoids unnecessary biopsies and can possibly reduce the number of biopsy cores (in TBx only or saturation targeted biopsy (sTB) strategies), leading to a reduction in pathology costs.\textsuperscript{15,26} Additionally, it leads to a decrease in insignificant PCa detection and, therefore, alleviates the patient burden and healthcare costs associated with active surveillance.\textsuperscript{1–3,10,11}

Targeting small lesions (<10 mm)

One of the key factors for accurate targeted biopsy is reaching small lesions in specific and difficult to reach (e.g., lateral or anterior) locations. In-bore MRI-guided biopsy can be performed using a software-guided robotic device, allowing for precise target (<10 mm) localisation within any area of the prostate. A similar robotic device is available for US, allowing for pre-programmed biopsy schemes.\textsuperscript{27} Robotic assistance is currently not widely used; future development might lead to increased accuracy and could decrease the current role of operator experience.\textsuperscript{24}

Targeting inaccuracies

Finally, it is important to consider that, when applying the US-guided biopsy approach, targeting inaccuracies can occur due to prostate deformation, patient positioning, mismatching of imaging planes and, in case of cognitive-TBx, incorrect visual registration. Several MRI-US fusion platforms have been developed to minimise the impact of these issues. Whether one of these platforms has an advantage over the others, or over cognitive-TBx, remains to be proven. A systematic review evaluating a variety of different fusion platforms, could not find significant difference in cancer detection rates (CDR) between them.\textsuperscript{28} Evidence on CDR of fusion-TBx versus cognitive-TBx is conflicting, with some studies finding no difference and others finding improved CDR for fusion-TBx.\textsuperscript{29–31} The benefit of fusion TBx, as well as in-bore biopsy, compared to cognitive-TBx might be more apparent in lesions < 10 mm.\textsuperscript{31} In addition to accurate targeting, raising the number of targeted biopsy cores improves CDR in smaller lesions. The use of SBx templates, in which 10–20 cores are obtained from the lesion and adjacent sectors, shows promising results. Two recent single centre trials showed that sTB had detection rates of 91 and 99% for csPCa compared to TBx combined with SBx.\textsuperscript{11,25} Targeting inaccuracies as a result of image fusion could theoretically be solved by using US as the diagnostic imaging modality, instead of mpMRI. However, conventional greyscale US lacks the diagnostic accuracy to compete with mpMRI. Having said that, novel US modalities, such as micro-ultrasound and computer-aided quantification dispersion imaging (CUDI), have shown great improvements in PCa detection. These novel ultrasound modalities will be further discussed below.
Future perspectives: developments in ultrasound
Considering most biopsy guidance is performed using ultrasound imaging, it is interesting to elaborate on the idea of using advances ultrasound imaging as the diagnostic imaging modality for PCa. If diagnostic accuracy is sufficient, the use of one of these US modalities as biopsy guidance would resolve targeting inaccuracies caused by image fusion errors.

Micro-ultrasound, operating at high frequency (29 MHz), leads to improved resolution and more detailed prostate imaging. The micro-US system (ExactVu) supports MRI-micro-US fusion and initial evaluations show promising results. In a recent, multicentre trial, 1040 patients with a clinical suspicion for PCa underwent mpMRI and high-resolution micro-ultrasound before prostate biopsies. Micro-ultrasound reached a sensitivity of 94% for GG ≥ 2 PCa compared to 90% in mpMRI. In this study, however, most clinical operators were not blinded for mpMRI results when performing the US targeting, thereby introducing a major source of bias.

Another US modality showing promising results is shear wave elastography (SWE). SWE visualises and quantifies solid lesions in soft tissue. Studies evaluating SWE, using radical prostatectomy specimens as the reference standard, showed a direct relation between tissue stiffness and presence of PCa, with stiffness (measured as Young's Module) increasing with higher Gleason Grades. A systematic review on 2277 patients originating from 16 studies, showed specificities and sensitivities for PCa detection ranging from 0.69 to 0.85 and 0.71–0.87, respectively.

Finally, Contrast-Enhanced Ultrasound (CEUS) and its derivative imaging technique Contrast Ultrasound Dispersion Imaging (CUDI) have also shown great potential in PCa detection. The applicability of CEUS and CUDI in prostate cancer detection is based on visualisation and quantification of abnormalities in the microvascular architecture. Angiogenesis, necessary for PCa growth and progression, causes alterations in the microvascular structures. These alternations lead to abnormalities in dispersion and perfusion of the ultrasound contrast agent (UCA) that can be utilised with CEUS. The head-to-head trial comparing CUDI-TBx, mpMRI-TBx and SBx found that CUDI-TBx and mpMRI had comparable detection rates, finding 28 and 29% csPCa, respectively.

A major limitation of ultrasound, including SWE, CEUS and CUDI, have been its 2D image acquisition. Advances in US technology have now made it possible to make rapid 3D acquisition, thereby facilitating visualisation of the whole prostate using one bolus of intravascular contrast. By combining multiple US modalities some studies have shown that this multiparametric approach leads to an increase in diagnostic accuracy. This year, a multicenter trial (NCT04605276) will commence to further improve the CUDI-algorithm for 3D multiparametric ultrasound. Future clinical validation trials will have to provide further evidence for both novel US modalities to become standard diagnostic and biopsy guidance imaging modalities.

CONCLUSION
Based on the available data, a definitive answer to which is the optimal biopsy technique cannot be given. Ultimately, the choice should rely on local experience and availability of equipment. The selection of patients based on their individual characteristics and onsite cost-benefit analysis are of utmost importance as well. The development of novel imaging techniques may lead to further shifts in this important diagnostic pathway.

REFERENCES


