Review

Magnetic resonance imaging-guided prostate biopsy—A review of literature

Kulthe Ramesh Seetharam Bhat a,*, Srinivas Samavedi b, Marcio Covas Moschovas a, Fikret Fatih Onol a, Shannon Roof a, Travis Rogers a, Vipul R. Patel a, Ananthakrishnan Sivaraman c

a Department of Urology, AdventHealth Global Robotics Institute, Celebration, FL, United States
b The Hays Medical Centre, University of Kansas Health System, Hays, KS, USA
c Chennai Urology and Robotics Institute, Chennai, Tamilnadu, India

Received 24 December 2019; received in revised form 22 April 2020; accepted 1 June 2020
Available online 28 July 2020

KEYWORDS
MRI targeted biopsy; MRI fusion biopsy; MRI cognitive biopsy; MRI fusion technology; Prostate biopsy

Abstract  Objective: Multiparametric magnetic resonance imaging (MP-MRI) helps to identify lesion of prostate with reasonable accuracy. We aim to describe the various uses of MP-MRI for prostate biopsy comparing different techniques of MP-MRI guided biopsy.

Materials and methods: A literature search was performed for “multiparametric MRI”, “MRI fusion biopsy”, “MRI guided biopsy”, “prostate biopsy”, “MRI cognitive biopsy”, “MRI fusion biopsy systems”, “prostate biopsy” and “cost analysis”. The search operation was performed using the operator “OR” and “AND” with the above key words. All relevant systematic reviews, original articles, case series, and case reports were selected for this review.

Results: The sensitivity of MRI targeted biopsy (MRI-TB) is between 91%–93%, and the specificity is between 36%–41% in various studies. It also has a high negative predictive value (NPV) of 89%–92% and a positive predictive value (PPV) of 51%–52%. The yield of MRI fusion biopsy (MRI-FB) is similar, if not superior to MR cognitive biopsy. In-bore MRI-TB had better detection rates compared to MR cognitive biopsy, but were similar to MR fusion biopsy.

Conclusions: The use of MRI guidance in prostate biopsy is inevitable, subject to availability, cost, and experience. Any one of the three modalities (i.e. MRI cognitive, MRI fusion and MRI in-bore approach) can be used. MRI-FB has a fine balance with regards to accuracy, practicality and affordability.

© 2021 Editorial Office of Asian Journal of Urology. Production and hosting by Elsevier B.V. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

* Corresponding author.
E-mail address: Seetharam_bhat2003@yahoo.co.in (K.R. Seetharam Bhat).

Peer review under responsibility of Second Military Medical University.

https://doi.org/10.1016/j.ajur.2020.07.001
2214-3882/© 2021 Editorial Office of Asian Journal of Urology. Production and hosting by Elsevier B.V. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
1. Introduction

Transrectal ultrasound (TRUS)-guided prostate biopsy is commonly used to diagnose prostate cancer (PCa). The use of TRUS for prostate biopsy was first described by Watanabe et al. in 1968 [1]. The next major advancement in prostate biopsy is the sextant biopsy, described by Hodge et al. in 1989 [2]. The appearance of the prostate malignant lesion on a TRUS could be either hypoechoic, isoechoic, or hyperechoic, none of which are pathognomonic. With these limitations in TRUS biopsy, magnetic resonance imaging (MRI) guided biopsy was investigated due to increased quality of images obtained by MRI.

In a Nobel prize winning paper in 1946, Felix Bloch proposed that the atomic nucleus acts like a magnet with momentum due to spinning protons, and its first ever clinical application [3]. In 1960, Damadian et al. [4] differentiated between malignant and normal tissue in a rat, which formed the basis of use of MRI today. The use of MRI is popular due to its contrast resolution, especially in detecting soft tissue lesions. With further advancement in MRI and the use of multiparametric MRI (MP-MRI), radiologists were able to identify lesions with reasonable accuracy, especially in PCa. These advances triggered the use of MP-MRI for prostate biopsy, either in the form of in-bore magnetic resonance imaging targeted biopsy (MRI-TB), MRI cognitive biopsy (MRI-CB) or MRI fusion biopsy (MRI-FB).

2. Methods

A literature search for articles in English was performed with PubMed, Google Scholar, WHO Hinari, Web of Science, Cochrane database and Cochrane library using the terms “multiparametric MRI”, “MRI fusion biopsy”, “MRI guided biopsy”, “prostate biopsy”, “MRI cognitive biopsy”, “MRI fusion biopsy systems” and “cost analysis”. Articles primarily after 2014 and other important studies prior to 2014 were selected and reviewed in this article. The search operation was performed using operator “OR” and "AND" with the above key words. All relevant systematic reviews, original articles, case series and case reports were selected for this review.

3. MP-MRI

MP-MRI has been used for the diagnosis and staging of PCa [5]. The MP-MRI is a combination of high-resolution T2-weighted images (T2WI), dynamic contrast-enhanced MRI (DCE-MRI), and diffusion-weighted imaging (DWI) to assess the anatomy and detect tumours >0.5 cm³ [6–9]. The prostate imaging reporting and data system (PIRADS) is a scoring system proposed by the European Society of Urogenital Radiology (ESUR) to diagnose PCa in the year 2012 [10]. Later, magnetic resonance spectroscopic imaging (MRSI), which was initially part of PIRADS, has been discontinued in PIRADS v2.0 [11]. Specific MP-MRI performed using “detection protocol” is used in MP-MRI guided biopsy [12–14]. Prior to the PIRADS scoring, the radiologist used the Likert scale, wherein without strict criteria the radiologist used a 5 point grading system based on overall impression. More radiologists are familiar with this system, though it is subject to interpreter variability [15]. Both PIRADS and LIKERT scales had similar rates of decision to biopsy, with LIKERT performing better in identifying clinically significant prostate cancer (csPCa). Unlike PIRADS, LIKERT is flexible, intuitive, and allows the radiologist to use clinical data. However, it is useful only if the radiologist has sufficient experience and it has the drawback of being subjective [16,17].

In 2019, an international group comprised of the American College of Radiology (ACR), the ESUR and the AdMeTech Foundation published an updated PIRADS v2.1 [18]. DWI MRI is the dominant sequence in the peripheral zone, and T2 weighted (T2w) MRI is the dominant sequence in the transitional zone. The latest scoring system is more specific about the uses of b values in DWI and DCE temporal resolution, especially in scores 2 and 3, thus adding more clarity to PIRADS score. The changes in T2w are in assessment in scores 1 and 2 defining encapsulated nodule and atypical nodules [18].

Final PIRADS v2.1 assessment categories [18]:

- 1 Very low (clinically significant cancer highly unlikely)
- 2 Low (clinically significant cancer unlikely)
- 3 Intermediate (clinically significant cancer equivocal)
- 4 High (clinically significant cancer likely)
- 5 Very high (clinically significant cancer highly likely)

Recently published prospective validation studies of PIRADS v2 show the detection rates for PCa were 35%–39%, 60%–72% and 91% for PIRADS 3, 4, and 5 lesions, respectively. The rates of csPCa were 17%–23%, 34%–49% and 67%–77% for PIRADS 3, 4, and 5, respectively [19,20]. Pepe et al. [21] suggested the use of PIRADS 3 or above as the safe cutoff for MRI-TB, wherein 83.8% of the csPCa were diagnosed, with a false-negative rate of 16.2%, which negated the need for saturation biopsy. In a meta-analysis of 21 studies evaluating the diagnostic performance of PIRADS v2, the reported pooled sensitivity and specificity were 0.89 (95% confidence interval [CI] 0.86–0.92) and 0.73 (95% CI 0.60–0.83) [22]. In another meta-analysis by Zhang et al. [23], 13 studies were reviewed and it was concluded that the pooled sensitivity and specificity were 0.85 (95% CI 0.78–0.91) and 0.71 (95% CI 0.60–0.80), respectively. A study comparing the reader agreement of six highly experienced uroradiologists from six institutions showed moderate reproducibility (kappa = 0.55) [24]. A subsequent study by Muller et al. [25] showed kappa interpreter agreement for overall suspicion score, T2W in the peripheral zone (PZ), T2W in the transitional zone (TZ), DWI, and DCE-MRI of 0.46, 0.47, 0.37, 0.40 and 0.46, respectively. The limitations of MP-MRI include limited sensitivity for the detection of PCa in the TZ, especially in the setting of benign prostatic hyperplasia (BPH). Similarly, BPH nodules can mimic cancers, lesions especially in DWI. Numerous benign and premalignant lesions like granulomatous prostatitis, adenosis, and prostatic intra-epithelial neoplasia can mimic PCa [26].

4. MRI-TB

With many multicentric prospectively designed studies and systematic reviews (Table 1) confirming superiority of the
use of MRI guided TRUS biopsy, the current debate is if one should continue doing the systematic biopsy along with MRI guided biopsy. MRI-TB is highly sensitive when compared to standard TRUS biopsy and has better detection rates of csPCA. Combining both has been suggested to increase the yield of the biopsy [27]. The sensitivity of MRI-TB is between 91%—93% and the specificity is between 36%—41% in various studies. Also, it has a high negative predictive value of 89%—92% and a positive predictive value of 51%—52% [28–30].

4.1. Approaches to MRI-TB

MRI-TB has three main approaches:

4.1.1. MRI-CB

MRI-CB involves visual registration by the operator, creating a mental map of the MRI images including suspicious lesions, and targeting those spots while doing a TRUS-biopsy. The operator mentally analyses the images and measures various distances using three-dimensional (3D) spatial reasoning and recognition of set patterns in MR images, thus locating the target spot. With regards to its yield, MRI-CB is superior to the systematic prostate biopsy, and in expert hands it could probably equal the MRI-FB in diagnostic yield.

Sciarrà et al. [40] prospectively compared two groups of men, one undergoing systematic biopsy and the other undergoing MRI-CB, with significant differences in PCa detection rates in both groups (24.5% vs. 45.5%, \( p = 0.01 \)). Similarly, Lee et al. [41] demonstrated a higher yield in previous negative biopsy patients who had MRI-CB vs. systematic core biopsy (28.8% vs. 3.6%, \( p = 0.012 \)). MRI-CB was particularly useful in anterior and apical tumours that are often missed in systematic biopsy.

4.1.2. MRI-FB

This method involves a real time fusion of MP-MRI images which are superimposed on the real time TRUS images, thus targeting the lesion seen on MP-MRI during the prostate biopsy. There are several commercially available systems for co-registering MP-MRI scans with real time ultrasound (US) (Table 2). Of these, URONAV and Artemis are the two most commonly used systems throughout the United States. These machines differ in the following ways [42]:

- Registration algorithm—The prostate can become deformed for a number of reasons, such as a full bladder, endorectal coil during MP-MRI, patient position, haemorrhage, intra procedural distortion, etc. Based on the ability to compensate for the distortion, the type of fusion is classified as:
  a. Rigid—Does not consider the organ deformation and could potentially lead to suboptimal anatomic registration.
  b. Elastic—Accounts for the gland deformation and provides a better fusion leading to improvement in accuracy.
- Strategy of navigation
  a. Organ-based—This involves retrospective guidance, where the device performs an organ-based navigation in which the location of the US transducer, the proposed needle target, and the prostate are mapped retrospectively using software programs.
  b. Electromagnetic tracking—This is a real time tracking system using a Global Positioning System (GPS) like tracking tool that tracks using electromagnetic fields and angle sensors. The drawback to this type of tracking is that it is cumbersome and expensive, as it uses extra hardware, like an electromagnetic field generator near the patient and a sensor on the US probe.
    - Post biopsy needle track documentation
      a. Operator performs a US of the prostate with biopsy needle in situ to exactly track the area of interest.
      b. Approximation of the biopsy needle based on orientation relative to the target and US transducer.
    - Design of the robotic arms
      
      Some systems have articulating arms with steppers and coders to reduce operator dependency, thus improving consistency and accuracy, while other systems merely guide the operator to target the desired area for sampling.

4.1.2.1. Artemis

Using computer software, the operator identifies the lesions and marks them prior to the biopsy. During the procedure, the Artemis fusion platform initially calibrates the MP-MRI images and the real-time US image to the mechanical arm, developing a 3D fusion mode. The machine can track and save the needle tract, which is useful for patients who are on active surveillance during repeat biopsy [43]. This system is highly accurate, with an accuracy of about 1.2 mm±1.1 mm [44]. Artemis device had thrice more detection rates of cancer versus the standard biopsy [43]. However, the main drawback of this system is that the biopsy involves two devices, and change in patient position can alter the fusion requiring the procedure to be repeated. The robotic arm has a limited degree of freedom (doF), the prostate can become deformed during apposition of the TRUS probe and the fusion may not be accurate.

4.1.2.2. UroNav

UroNav uses a free hand approach to electromagnetically track the motion of a real-time transrectal US probe to MP-MRI loaded and has an accuracy of 2.3 mm±0.9 mm [45]. The advantage of this system is that it enables rigid and elastic registration, and due to its freehand nature, it allows the operator to perform the procedure in the office.

There have been reports of MRI-FB being performed via the transperineal route using the conventional robotic system, by switching from transrectal to transperineal modules. The BiopSee system is used to perform transperineal biopsy, and it includes a transrectal probe with a mechanical stepper that aids in performing the biopsy transperineally [46].

The major challenges these systems face are the associated cost and learning curve. It requires continuous learning and feedback between different specialties. The MRI fusion technology comes with significant cost and requires an initial investment with a steep learning
| Authors                  | Trial design                                                                 | Number of patients | Conclusion                                                                 |
|-------------------------|------------------------------------------------------------------------------|--------------------|---------------------------------------------------------------------------|
| Kasivisvanathan et al.  | - Multicenter, randomized, noninferiority trial at 25 centers in 11 countries (PRECISION STUDY) | 500                | - There was significantly higher rates of detection of prostate cancer using MRI-TB vs. systematic biopsy ($p=0.005$). |
| van der leest et al.    | - Multicentric prospective paired cohort—Patients underwent prebiopsy MRI   | 626                | - MRI-TB and systematic TRUS biopsy had similar rates of clinically significant prostate cancer. MRI-TB detected a lower proportion of clinically insignificant disease. |
| Rouvière et al.         | - Multicentric-prospective paired cohort—16 centre in France.               | 275                | - 20% of clinically significant prostate cancer was diagnosed by MRI-TB, 14% by TRUS biopsy. Combined biopsy had maximal detection rates (66%). |
| Ahmed et al.            | - Multicentric-prospective paired cohort (PROMIS trial)—Patients underwent all three test. | 576                | - MP-MRI guided biopsy leads to 18% more clinically significant cancer compared to standard pathway. MP-MRI when used to triage men can avoid unnecessary biopsy in 27%, but can miss 5% of clinically significant prostate cancer. |
| Thompson et al.         | - Single centre—prospective paired cohort                                  | 344                | - MP-MRI can avoid 23% of unnecessary biopsies and enhanced the detection of low risk PCa by 34%. |
| Panebianco et al.       | - Single centre—RCT                                                       | 570 in each group  | - Accuracy of MP-MRI was 97%.                                             |
| Baco et al.             | - Single centre—RCT                                                       | 175 (86 MRI vs. 89 control) | - Prostate cancer detection rates were similar in both groups.           |
| Siddiqui et al.         | - Single centre—Paired cohort                                              | 1 003              | - MRI-TB diagnosed 30% more high risk PCa and 17% lesser low-risk PCa. Sensitivity for high risk PCa was 77% vs. 53% with similar specificity. |
Mendhiratta et al. [47] showed the yield of MRI-FB increased over 33 months from 63% to 86%, thus supporting its use and demonstrating a learning curve. Inaccurate segmentation of MP-MRI images and misregistration of the MP-MRI images or transrectal US images can lead to discrepancies in targeting. In order to overcome

| Authors | Trial design | Number of patients | Conclusion |
|---------|--------------|--------------------|------------|
| Drost et al. [30] | A Cochrane systematic review and meta-analysis | 43 studies (6,871 men) | - 43 studies (6,871 men) - This review used template biopsy as standard reference and compared the diagnostic accuracy of MRI index only lesions, MRI-targeted biopsy, MRI pathway (MRI + / MRI targeted biopsy) and systematic biopsy. - The primary end point was ISUP 2 and above (csPCa), and secondary end point was ISUP 1. |
| Elwenspoek et al. [35] | Systematic review and meta-analysis - Evaluated three biopsy 1) Pathway systematic biopsy 2) MRI-TB 3) Both pathway | 7 RCTs (2,582 men) csPca definition varied in different series | - 7 RCTs (2,582 men) csPca definition varied in different series - 57% improvement in detection of csPca, 33% reduction in number of biopsy and 77% reduction in the number of cores. |
| Woo et al. [36] | Systematic reviews and meta-analysis - MRI-TB 1) Pathway systematic biopsy 2) MRI plus systematic biopsy as intervention 3) Systemic TRUS as comparator | 9 RCTs (2,908 men) csPca definition varied in different series | - MRI stratified pathway detected more clinically significant PCa than TRUS biopsy (relative detection rate 1.45 for all men, 1.42 for biopsy naïve and 1.6 for men with prior negative biopsy). |
| Schoots et al. [37] | Systematic review and meta-analysis - MRI-TB vs. TB | 16 studies (1,926 men) varied definition | - MRI-TB vs. TB | MRI-TB had higher rate of detection of significant prostate cancer (MRI-TB vs. TB sensitivity: 0.91 vs. 0.76) and lower rate of detection of insignificant prostate cancer (0.44 vs. 0.83). |
| Moore et al. [38] | Systematic review | 599 patients | - MRI-TB detects clinically significant PCa with less number of cores (MRI-TB [3.8 cores] vs. TB [12 cores]). |
| Wegelin et al. [39] | Systematic review and meta-analysis - Three techniques of MRI-TB are available: 1) In-bore MRI target biopsy (MRI-TB), 2) MRI-transrectal ultrasound fusion (MRI-FB), and 3) cognitive registration (MRI-CB). | 43 studies | - Overall detection rates are similar in different biopsy techniques. - Increased rate of csPCa and decrease in clinically insignificant PCa. |

MCCL, maximum cancer core length; MRI, magnetic resonance imaging; MRI-TB, MRI targeted biopsy; MRI-FB, MRI fusion biopsy; MRI-CB, MRI cognitive biopsy; MP-MRI, multiparametric MRI; NPV, negative predictive value; PIRADS, Prostate Imaging Reporting and Data Systems; csPca, clinically significant prostate carcinoma; PPV, positive predictive value; RCT, randomised control trial; PSA, prostate-specific antigen; ISUP, International Society of Urological Pathology; TPM, template prostate mapping.
the registration and targeting errors, it is a norm to obtain at least two spatially distributed samples from the target [48].

4.1.3. In-bore MRI target biopsy
This technique involves obtaining tissue samples with direct MRI guidance in MRI gantry, thus allowing the operator to target the lesion in real time. Though traditionally done using the open MRI system, this technique is now being performed in closed systems, as well using either 1.5 T or 3 T MRI systems. The biopsy can be performed either transrectally or transperineally. DynaTRIM is a commonly used portable device used to perform the in-bore biopsy transrectally [49]. The patient is placed in a prone position and the device is fixed underneath the patient; it has an adjustable needle guide that has three degrees of freedom (cranial/caudal, anterior/posterior and left/right). Once positioned, a rectal sleeve is placed and Sagittal T2WIs are obtained to position the arm in a neutral position. Axial images are then processed in the DynaTRIM workstation to target the lesion [49]. A key advantage of this method is that any series of MRI can be used to target the lesion effectively. After identification of the lesion, the software plans the trajectory of the 18-gauge needle to biopsy the prostate. This procedure has a learning curve of about 25–30 patients, and takes about 30 min with an additional target taking about 15 min [48].

The main advantage of this technique is that one can target the biopsy site more accurately. However, the evidence to support its usefulness in small lesions is at best anecdotal, and the data on the exact size of the lesion where this might be advantageous are lacking. The main limitations of this technique are its limited availability, long procedure time, and cost. Another drawback is the logistic issue to perform such a procedure by a urologist amidst their busy practice (Table 3).

4.1.4. Comparison between different MRI biopsy techniques
There are various studies that show the superiority of the MRI-TB over the conventional standard 12 core biopsy. However, when it comes the comparison between the individual modalities of MRI-TB, the data are scarce. Though the current available literature does not show if one is superior over the other, MRI-FB is more popular, probably because it is practical and can easily be incorporated into the workflow. The MRI-CB may have similar advantages in terms of its practicality and lower cost. There is evidence suggesting the superiority of MRI-fusion technology over the
| Study                  | Methods                                                                 | Conclusion                                                                 | Significant findings                                                                 |
|-----------------------|-------------------------------------------------------------------------|----------------------------------------------------------------------------|--------------------------------------------------------------------------------------|
| Puech et al. [50]     | - Intraindividual comparison of systematic vs. MRI-CB vs. MRI-FB       | - Cognitive and software fusion biopsy were significantly superior to systematic biopsy. | • Prostate carcinoma positivity rates for SB (59%) vs. MRI-TB (both) (69%).       |
|                       |                                                                         | - The yield was similar between the cognitive and the fusion biopsy.         | • MRI-CB vs. MRI-FB positivity rate (47% vs. 53%; no significant difference). |
|                       |                                                                         |                                                                             | • MRI-FB 20% vs. MRI-CB 15%; \( p = 0.052 \).                                    |
|                       |                                                                         |                                                                             | • MRI-FB 77% vs. MRI-CB 60; \( p = 0.0104 \).                                    |
|                       |                                                                         |                                                                             | • Multivariable analysis reveals size of the lesion as an independent factor predicting cancer in MRI-FB. |
| Wysock et al. [51]    | - Intra individual comparison between MRI-CB and MRI-FB by two different urologist | - MR-FB had higher detection rate of Gleason score 7 or above.               | • MRI-FB had 100% sampling rate but does not translate to 100% cancer detection rates as the prostate MP-MRI specificity ranges between 44% and 67%. |
|                       |                                                                         | - MRI-FB also better characterised nonbenign histology.                      | • Detection rates of csPSA did not differ between MRI-FB and MR-in bore guided biopsy (49% vs. 61% respectively). |
| Cool et al. [52]      | - Comparison between 2D vs. 3D vs. MRI-FB using simulation models form 100 patients | - MRI-CB was inferior to MRI—TRUS fusion biopsy, irrespective of operator level of experience. | • Overall detection rates for MRI-FB vs. MRI in-bore biopsy were 66.7% vs. 85% (\( p < 0.05 \)). |
|                       |                                                                         |                                                                             | • The mean distance of cognitive targets was 10.6 mm from the MRI fusion targets with 15.3% patients having less than 5 mm discrepancy. |
|                       |                                                                         |                                                                             | • The difference between spatial difference between the experience and inexperienced were 9.7±5.1 mm vs. 13.4±7.4 mm; \( p = 0.042 \). |
| Venderink et al. [53] | - Retrospective comparison between in bore MRI-guided biopsy vs. MRI-FB | - No significant differences in detecting csPSA between in bore MR biopsy and MR fusion guided biopsy. | • 267 patients (106 in MRI-guided in-bore group vs. 104 in other approaches) |
|                       |                                                                         |                                                                             | • In-bore approach vs. other approaches (mean cores: 5.6 cores vs. 17 cores, \( p < 0.001 \)). |
| Kwak et al. [54]      | - Two operators of different experience performed visual registration (MR-CB) and MRI-FB biopsy | - The amount of mistargeting in MRI-CB was high regardless of site the lesion. | • No significant difference \( p < 0.05 \). |
|                       |                                                                         | - Different levels of experience led to substantial difference in visual registration leading to missed diagnosis. | • Pathological diagnosis |
|                       |                                                                         |                                                                             | • Grade |
|                       |                                                                         |                                                                             | • Volume of tumour |
|                       |                                                                         |                                                                             | • Significant fewer cores in the in-bore approach. |
| Arsov et al. [55]     | - Randomized patient to in-bore MR imaging targeted biopsy vs. systematic biopsy/MRI-FB | - No significant difference | • Pathological difference |
|                       |                                                                         |                                                                             | • Grade |
|                       |                                                                         |                                                                             | • Volume of tumour |
|                       |                                                                         |                                                                             | • Significant fewer cores in the in-bore approach. |

(continued on next page)
4.1.5. MRI guided transperineal biopsy

Transperineal biopsy has recently gained popularity, as transrectal biopsy has some disadvantages that are overcome by a transperineal route. A transrectal biopsy necessitates the use of prophylactic antibiotics because the needle passes through the rectum. Until now, fluoroquinolone has been advocated as the antibiotic of choice, but the use of this particular group of antibiotics has been questioned for several reasons. First, its use is now restricted by the Federal Drug Agency (FDA). In 2016, the FDA advised against its use unless deemed necessary, due to the risk of permanent disabling joint issues [57]. Second, the readmission rates following the quinolone use were similar to no antibiotic use, suggesting increasing antibiotic resistance due to the widespread use of fluoroquinolone [58, 59]. Therefore, this has led to the use of either a combination of antibiotics or the use of carbapenems as prophylaxis for prostate biopsy [60]. In this context, rectal swab culture-specific antibiotic use is more effective [61]. For the above-mentioned reasons, the transperineal route has been adopted in practice by many clinicians [58]. The advantage of the transperineal biopsy compared to transrectal biopsy is the reduced chance of infection and better detection of cancer in the anterior zone of the prostate, with an almost 10% increased detection rate of these tumours [62–64]. However, transperineal biopsy may have increased rate of urine retention [65]. The targeting is done using either the freehand approach, the brachytherapy grid, or with robotic guidance. It is considered a relatively clean procedure, as the needle passes through the perineal skin. In a multicentric study by Pepdjonovic et al. [66], the readmission rate following transperineal biopsy was zero wherein the last 710 cases (59.5%) patients received a single dose of cephazolin as prophylaxis.

Historically, transperineal biopsy is usually performed under general anaesthesia. However, recently there are many studies reporting the feasibility of using local anaesthesia [67–69]. Of note, the Cambridge Prostate Biopsy (CAMPROBE), developed by Thurtle et al. [70], has shown to be effective in administering local anaesthesia with reduced pain and 87% preference over TRUS biopsy. Technically, this is similar to the transrectal biopsy; the MRI targeted transperineal biopsy can be either cognitive guided, MRI software fusion biopsy, or in-bore MRI transperineal biopsy. MRI guided cognitive transperineal biopsy is done by placing the TRUS probe transrectally and using a brachytherapy grid. Using a stepper and a stabilizer, the prostate is imaged in sagittal and transverse views, and the needle is directed towards the target lesion site. The 5 mm grid spacing can accurately help in targeting the lesion.

MRI guided fusion transperineal biopsy has the same principles as the MRI fusion transrectal biopsy. The difference is that the software compensates for the needle tracking transperineally. One can perform this either with or without the brachytherapy grid. After capturing the US images, the software performs rigid/elastic registrations and suggests appropriate grid holes that can be used for target sampling. The MRI-FB systems capable of performing transperineal biopsy are listed in Table 3. In a comparison between the systematic template transperineal biopsy (TPB) and MRI fusion biopsy, the template biopsy missed 21% of clinically significant cancer and MRI-TB missed 20%, thus concluding that their detection rates are similar and should be used in combination [71].

The first transperineal in-bore imaging biopsy was reported in 2001, and was useful in patients with limited rectal access due to previous proctocolectomy or rectal stenosis [72]. Visualase is a commercially available system used to perform transperineal biopsy, wherein the patient is supine and a needle guide template with fiducial marker is strapped
onto the perineum. The software then plans the needle adjustments and positioning to target the correct hole to sample the target on the prostate [48]. In view of its cumbersome nature and difficulty in incorporating the in-bore MRI transperineal biopsy into the routine workflow of a clinician’s practice, it is not very popular. Of note is a study comparing the use of manual versus robotic template in the in-bore MR TB, which found the robotic template had statistically higher accuracy in needle placement (p<0.032) [72].

The iSR’obot Mona Lisa is a robot exclusively developed for the transperineal biopsy. It performs the biopsy through two transperineal skin punctures that act as a pivot, thus giving a biconical configuration of core positions. The TRUS probe images the prostate from base to apex and generates a 3D model of the prostate. This image is then fused with MRI images of the prostate, thus marking out the targets for biopsy. Once the Mona Lisa device is positioned, the robot automatically creates a map and moves to the desired position for the biopsy. Under the US guidance, the clinician fires the needle, which is monitored, and the robot positions itself to the next target. A standard transperineal biopsy can be completed in half an hour [74].

Pepe et al. [64] compared MRI targeted TRB versus MRI targeted cognitive transperineal biopsy. The study involved 200 patients who underwent standard template transperineal biopsy followed by MRI fusion transrectal biopsy of the suspicious lesions. These patients were then targeted again using MRI-targeted cognitive transperineal biopsy. Sixty cases were diagnosed with csPca, which was confirmed using an MP-MRI. Also, 20 of these cases were missed by MRI fusion, and only four were missed by MRI-CB. The major drawback of this study is that the definition of csPca was restricted to two or more cores with Gleason score 6 or above, which is debatable.

4.1.6. Cost analysis of MRI-TB

With so many available options, the cost comes into consideration in choosing the optimal modality, based on the indication of the prostate biopsy. Compared to a standard TRUS biopsy under local anaesthesia, the cost of TRUS biopsy under sedation, the cost of transperineal template biopsy under general anaesthesia, MRI-FB under sedation and sedation in-bore prostate biopsy were significantly higher (1.9 vs. 2.5 vs. 2.5 vs. 2.2, p<0.001). In the same series, cancer detection rates when compared to TRUS biopsy were higher in fusion biopsy (16% vs. 36%, p<0.001) and transperineal template biopsy (16% vs. 34%, p<0.001) [75].

In another cost analysis by Venderink et al. [76], the incremental cost-effectiveness ratio following a MRI-FB versus the systematic TRUS biopsy was $1 470 per quality-adjusted life year gained. An in-bore MRI guided biopsy would be cost effective if its sensitivity for csPca is 11.8% higher than the sensitivity of MRI-FB [76]. To improve the sensitivity of either modality, newer definitions of csPca can be used. Also, to improve cost-effectiveness, lowering the upper limit of the willingness to pay threshold recommended by National Institute for Health and Care Excellence (NICE) has been suggested. The major limitation of this study is that it depends on the input parameters into the complex calculation and as it was performed in Netherlands, one cannot extrapolate the findings to other populations.

[76]. De Rooij et al. [77] concluded that MRI-guided biopsy is cost effective if the sensitivity of MRI guided biopsy is 90% for any PCa.

In order to cut down cost and avoid unnecessary biopsy in patients with lower suspicion of PCa, a biparametric MRI (bp-MRI) with axial T2WIs and diffusion weighted images (b values: 0, 100, 800 and 2000) can be performed, as they are the two dominant parameters of PIRADS scoring. This exam has a high negative predictive value of 97%, and takes approximately 15 min to perform. Also, the corresponding apparent diffusion coefficient maps can be generated. However, the potential disadvantage is that dynamic contrast-enhanced imaging, which is a part of the MP-MRI, is lacking; as a result, lesions of equivocal score 3 in the peripheral zone may not be upgraded to 4. Seventeen percent of these PIRADS 3 lesions on bp-MRI had clinically significant malignancy, thus justifying need for performing biopsy on these lesions [78]. Also, there are several reports suggesting follow-up using repeat MP-MRI in PIRADS 3 lesion, to avoid unnecessary biopsies, thus cutting overall cost [79–81].

5. Conclusion

The use of MP-MRI guidance in prostate biopsy is inevitable, and we have passed the point of doubt on its usage. However, depending on the availability, cost, and experience, any one of the three modalities (i.e. MRI cognitive, MRI fusion and MRI in-bore approach) can be used. MRI-FB has a fine balance with regards to accuracy, practicality and affordability.

Author contributions

Study concept and design: Vipul R. Patel, Kulthe Ramesh Seetharam Bhat, Ananthkrishnan Sivaraman.
Data acquisition: Kulthe Ramesh Seetharam Bhat, Marcio Covas Moschovas.
Data analysis: Travis Rogers, Fikret Fatih Onol.
Drafting of manuscript: Srinivas Samavedi, Kulthe Ramesh Seetharam Bhat.
Critical revision of the manuscript: Vipul R. Patel, Shannon Roof, Ananthkrishnan Sivaraman.

Conflicts of interest

The authors declare no conflict of interest.

References

[1] Watanabe H, Kato H, Kato T, Morita M, Tanaka M. [Diagnostic application of ultrasonotomography to the prostate]. Nippon Hinyokika Gakkai Zasshi 1968;59:273–9 [Article in Japanese].
[2] Hodge KK, McNeal JE, Terris MK, Stamey TA. Random systematic versus directed ultrasound guided transrectal core biopsies of the prostate. J Urol 1989;142:71–4.
[3] Shampo MA, Kyle RA. Felix Bloch—developer of magnetic resonance imaging. Mayo Clin Proc 1995;70:889. https://doi.org/10.4065/70.9.889.
[4] Damadian R, Minkoff L, Goldsmith M, Stanford M, Koutcher J. Field focusing nuclear magnetic resonance (FONAR):
visualization of a tumor in a live animal. Science 1976;194:1430–2.

[5] Sivaraman A, Bhat KRS. Screening and detection of prostate cancer—review of literature and current perspective. Indian J Surg Oncol 2017;8. https://doi.org/10.1007/s13193-016-0584-3.

[6] van As NJ, de Souza NM, Riches SF, Morgan VA, Sohalb SA, Dearnaley DP, et al. A study of diffusion-weighted magnetic resonance imaging in men with untreated localised prostate cancer on active surveillance. Eur Urol 2009;56:981–7.

[7] Tamada T, Sone T, Jo Y, Toshimitsu S, Yamashita T, Yamamoto A, et al. Apparent diffusion coefficient values in peripheral and transition zones of the prostate: comparison between normal and malignant prostatic tissues and correlation with histologic grade. J Magn Reson Imaging 2008;28:720–6.

[8] Villers A, Puech M, Mouton D, Leroy X, Ballereau C, Lemaitre L. Dynamic contrast enhanced, pelvic phased array magnetic resonance imaging of localized prostate cancer for predicting tumor volume: correlation with radical prostatectomy findings. J Urol 2006;176:2432–7.

[9] Girouin N, Mège-Lechevallier F, Tonina Senes A, Bissery A, Rabilloud M, Maréchal J-M, et al. Prostate dynamic contrast-enhanced MRI with simple visual diagnostic criteria: is it reasonable? Eur Radiol 2007;17:1498–509.

[10] Barentsz JO, Richenberg J, Clements R, Choyke P, Verma S, Hambrock T, Somford DM, Hoeks C, Bouwense SAW. PI-RADS 2.0: what is new? Diagn Interv Radiol 2015;21:382–4.

[11] Hambrock T, Somford DM, Hoeks C, Bouwense SAW, Huisman H, Yakar D, et al. Magnetic resonance imaging guided prostate biopsy in men with repeat negative biopsies and increased prostate specific antigen. J Urol 2010;183:520–7.

[12] Amsellem-Ouazana D, Younes P, Conquy S, Peyromaure M, Flam T, Debré B, et al. Negative prostate biopsies in patients with a high risk of prostate cancer. Is the combination of endorectal MRI and magnetic resonance spectroscopy imaging (MRSI) a useful tool? A preliminary study. Eur Urol 2005;47:582–6.

[13] Coakley FV, Kurhanewicz J, Lu Y, Jones KD, Swanson MG, Amsellem-Ouazana D, Younes P, Conquy S, Peyromaure M, Flam T, Debré B, et al. Negative prostate biopsies in patients with a high risk of prostate cancer. Is the combination of endorectal MRI and magnetic resonance spectroscopy imaging (MRSI) a useful tool? A preliminary study. Eur Urol 2005;47:582–6.

[14] Rougeiro O, Puech P, Renard-Penna R, Claudon M, Roy C, Mège-Lechevallier F, et al. Use of prostate systematic and targeted biopsy on the basis of multiparametric MRI in biopsy-naïve patients (MRI-FIRST): a prospective, multicentre, paired diagnostic study. Lancet Oncol 2019;20:100–9.

[15] Thompson JE, Van Leeuwen PJ, Moses D, Shnier R, Brenner P, Delpredo W, et al. The diagnostic performance of multi-parametric magnetic resonance imaging to detect significant prostate cancer. J Urol 2017;295:815–22.

[16] Drost FJH, Osess D, Nieboer D, Bangma CH, Stiermetz EE, Roobol MJ, et al. Prostate magnetic resonance imaging, with or without magnetic resonance imaging-targeted biopsy, and systematic biopsy for detecting prostate cancer: a Cochrane systematic review and meta-analysis. Eur Urol 2020;77:78–89.

[17] van der Leest M, Cornel E, Israël B, Hendriks R, Padhani AR, Drost FJH, Osses D, Nieboer D, Bangma CH, Steyerberg EW. Validation of prostate imaging reporting and data system at multiparametric MR imaging. Radiology 2015;277:741–50.

[18] van der Leest M, Cornel E, Israël B, Hendriks R, Padhani AR, Drost FJH, Osses D, Nieboer D, Bangma CH, Steyerberg EW. Validation of prostate imaging reporting and data system at multiparametric MR imaging. Radiology 2015;277:741–50.

[19] van der Leest M, Cornel E, Israël B, Hendriks R, Padhani AR, Drost FJH, Osses D, Nieboer D, Bangma CH, Steyerberg EW. Validation of prostate imaging reporting and data system at multiparametric MR imaging. Radiology 2015;277:741–50.

[20] van der Leest M, Cornel E, Israël B, Hendriks R, Padhani AR, Drost FJH, Osses D, Nieboer D, Bangma CH, Steyerberg EW. Validation of prostate imaging reporting and data system at multiparametric MR imaging. Radiology 2015;277:741–50.
Schoots IG, Roobol MJ, Nieboer D, Bangma CH, Steyerberg EW, Imaging and MRI guided biopsy 115
Wegelin O, van Melick HHE, Hooft L, Bosch JLHR, Reitsma HB, Lee SH, Chung MS, Kim JH, Oh YT, Rha KH, Chung BH. Magnetic
Xu S, Kruecker J, Turkbey B, Glossop N, Singh AK, Choyke P, Hadaschik BA, Kuru TH, Tulea C, Rieker P, Popeneciu IV,
Natarajan S, Marks LS, Margolis DJA, MacAiran M, Lieu P, Huang J, et al. Clinical application of a 3D ultrasound-guided prostate biopsy system. Urol Oncol Semin Orig Investig 2011; 29:334–42.
Kongnyuy M, George AK, Rastinehad AR, Pinto PA. Magnetic resonance imaging-ultrasound-guided prostate biopsy: review of technology, techniques, and outcomes. Curr Urol Rep 2016;17:1–9.
Sonn GA, Natarajan S, Margolis DJA, MacAiran M, Lieu P, Huang J, et al. Targeted biopsy in the detection of prostate cancer using an office based magnetic resonance ultrasound fusion device. J Urol 2013;189:86–92.
Natarajan S, Marks LS, Margolis DJA, Huang J, MacAiran ML, Lieu P, et al. Clinical application of a 3D ultrasound-guided prostate biopsy system. Urol Oncol Semin Orig Investig 2011; 29:334–42.
Xu S, Kruecker J, Turkbey B, Glossop N, Singh AK, Choyke P, et al. Real-time MRI-TRUS fusion for guidance of targeted prostate biopsies. Comput Aided Surg 2008;13:255–64.
Hadaschik BA, Kuru TH, Tulea C, Rieker P, Popeneivuc IU, Simpfendorfer T, et al. A novel stereotactic prostate biopsy system integrating pre-interventional magnetic resonance imaging and live ultrasound fusion. J Urol 2011;186:2214–20.
Mendhiratta N, Rosenkrantz A, Meng X, Huang R, Taneja S. MP16-03 the impact of a learning curve in the performance of MRI-US fusion-targeted prostate biopsy: Improvements in cancer detection over time. J Urol 2016;195:e161. https://doi.org/10.1016/j.juro.2016.02.2568.
Verma S, Choyke PL, Eberhardt SC, Oto A, Tempany CM, Turkbey B, et al. The current state of MR imaging-targeted biopsy techniques for detection of prostate cancer. Radiology 2017;285:343–56.
Woodrum D, Gorny K, Greenwood B, Mynderse L. MRI-guided prostate biopsy of native and recurrent prostate cancer. Semin Interv Radiol 2016;33:196–205.

Ewenspeok AMC, Sheppard AL, McInnes MDF, Merriel SWD, Rowe EWJ, Bryant RJ, et al. Comparison of multiparametric magnetic resonance imaging and targeted biopsy with systematic biopsy alone for the diagnosis of prostate cancer. JAMA Netw Open 2019;2:e198427. https://doi.org/10.1001/jamanetworkopen.2019.8427.
Woo S, Suh CH, Eastham JA, Zelefsky MJ, Morris MJ, Abida W, et al. Comparison of magnetic resonance imaging-stratified clinical pathways and systematic transrectal ultrasound-guided biopsy pathway for the detection of clinically significant prostate cancer: a systematic review and meta-analysis of randomized controlled trials. Eur Urol Oncol 2019;2: 605–16.
Schoots IG, Roobol MJ, Nieboer D, Bangma CH, Steyerberg EW, Hunink MGM. Magnetic resonance imaging-targeted biopsy: a systematic review and meta-analysis. Eur Urol 2015;68:438–50.
Moore CM, Robertson NL, Arsanious N, Middleton T, Villers A, Klotz L, et al. Image-guided prostate biopsy using magnetic resonance imaging-derived targets: a systematic review. Eur Urol 2013;63:125–40.
Wegelin O, van Melick HHE, Hooft L, Bosch JLHR, Reitsma HB, Barentsz JO, et al. Comparing three different techniques for magnetic resonance imaging-targeted prostate biopsies: a systematic review of in-bore versus magnetic resonance imaging-transrectal ultrasound fusion versus cognitive registration. Is there a preferred technique? Eur Urol 2017;71:517–31.
Sciarra A, Panebianco V, Cicciariello M, Salciccia S, Cattarino S, Lisi D, et al. Value of magnetic resonance spectroscopy imaging and dynamic contrast-enhanced imaging for detecting prostate cancer foci in men with prior negative biopsy. Clin Cancer Res 2010;16:1875–83.
Lee SH, Chung MS, Kim JH, Oh YT, Rha KH, Chung BH. Magnetic resonance imaging targeting biopsy in men with previously negative prostate biopsy results. J Endourol 2012;26:787–91.
Kongnyuy M, George AK, Rastinehad AR, Pinto PA. Magnetic resonance imaging-ultrasound-guided prostate biopsy: review of technology, techniques, and outcomes. Curr Urol Rep 2016;17:1–9.
Sonn GA, Natarajan S, Margolis DJA, MacAiran M, Lieu P, Huang J, et al. Targeted biopsy in the detection of prostate cancer using an office based magnetic resonance ultrasound fusion device. J Urol 2013;189:86–92.
Natarajan S, Marks LS, Margolis DJA, Huang J, MacAiran ML, Lieu P, et al. Clinical application of a 3D ultrasound-guided prostate biopsy system. Urol Oncol Semin Orig Investig 2011; 29:334–42.
Xu S, Kruecker J, Turkbey B, Glossop N, Singh AK, Choyke P, et al. Real-time MRI-TRUS fusion for guidance of targeted prostate biopsies. Comput Aided Surg 2008;13:255–64.
Hadaschik BA, Kuru TH, Tulea C, Rieker P, Popeneivuc IU, Simpfendorfer T, et al. A novel stereotactic prostate biopsy system integrating pre-interventional magnetic resonance imaging and live ultrasound fusion. J Urol 2011;186:2214–20.
Mendhiratta N, Rosenkrantz A, Meng X, Huang R, Taneja S. MP16-03 the impact of a learning curve in the performance of MRI-US fusion-targeted prostate biopsy: Improvements in cancer detection over time. J Urol 2016;195:e161. https://doi.org/10.1016/j.juro.2016.02.2568.
Verma S, Choyke PL, Eberhardt SC, Oto A, Tempany CM, Turkbey B, et al. The current state of MR imaging-targeted biopsy techniques for detection of prostate cancer. Radiology 2017;285:343–56.
Woodrum D, Gorny K, Greenwood B, Mynderse L. MRI-guided prostate biopsy of native and recurrent prostate cancer. Semin Interv Radiol 2016;33:196–205.
Puech P, Rouvière O, Renard-Penna R, Villers A, Devos P, Colombel M, et al. Prostate cancer diagnosis: multiparametric MR-targeted biopsy with cognitive and transrectal US-MR fusion guidance versus systematic biopsy-prospective multicenter study. Radiology 2013;268:461–9.
Wysock JS, Rosenkrantz AB, Huang WC, Stiftelman MD, Lepor H, Deng FM, et al. A prospective, blinded comparison of magnetic resonance (MR) imaging-ultrasound fusion and visual estimation in the performance of MR-targeted prostate biopsy: the PROFUS trial. Eur Urol 2014;66:343–51.
Cool DW, Zhang X, Romagnoli C, Izawa JI, Romano WM, Fenster A. Evaluation of MRI-TRUS fusion versus cognitive registration accuracy for MRI-targeted, TRUS-guided prostate biopsy. Am J Roentgenol 2015;204:83–91.
Venderink W, van der Leest M, van Luijtenaar A, van de Ven WJM, Futterer JJ, Sedelaar JPM, et al. Retrospective comparison of direct in-bore magnetic resonance imaging (MRI)-guided biopsy and fusion-guided biopsy in patients with MRI lesions which are likely or highly likely to be clinically significant prostate cancer. World J Urol 2017;35:1849–55.
Kwak JT, Hong CW, Pinto PA, Williams M, Xu S, Kruecker J, et al. Is visual registration equivalent to semiautomated registration in prostate biopsy? BioMed Res Int 2015;2015: 394742. https://doi.org/10.1155/2015/394742.
Arsov C, Rabenalt R, Blondin D, Quentin M, Hiester A, Godehardt E, et al. Prospective randomized trial comparing magnetic resonance imaging (MRI)-guided in-bore biopsy to MRI-ultrasound fusion and transrectal ultrasound-guided prostate biopsy in patients with prior negative biopsies. Eur Urol 2015;68:713–20.
Wegelin O, Exterkate L, van der Leest M, Kummer JA, Vreuls W, de Bruijn PC, et al. The FUTURE Trial: a multicenter randomised controlled trial on target biopsy techniques based on magnetic resonance imaging in the diagnosis of prostate cancer in patients with prior negative biopsies. Eur Urol 2019;75:582–90.
FDA Drug Safety Communication: FDA updates warnings for oral and injectable fluoroquinolone antibiotics due to disabling side effects | FDA. https://www.fda.gov/drugs/drug-safety-and-availability/fda-drug-safety-communication-fda-updates-warnings-oral-and-injectable-fluoroquinolone-antibiotics [accessed December 9, 2019].
Roth H, Millar JL, Cheng AC, Byrne A, Evans S, Grummet J. The state of TRUS biopsy sepsis: readmissions to Victorian hospitals with TRUS biopsy-related infection over 5 years. BJU Int 2015;116:49–53.
Lange D, Zappavigna C, Hamidizadeh R, Goldenberg SL, Paterson RF, Chew BH. Bacterial sepsis after prostate biopsy—a new perspective. Urology 2009;74:1200–5.
Tamma PD, Han JH, Rock C, Harris AD, Lautenbach E, Hsu AJ, et al. Carbapenem therapy is associated with improved survival compared with piperacillin-tazobactam for patients with extended-spectrum β-lactamase bacteremia. Clin Infect Dis 2015;60:1319–25.
Duplessis CA, Bavaro M, Simons MP, Marguet C, Santomauro M, Auge B, et al. Rectal cultures before transrectal ultrasound-guided prostate biopsy reduce post-prostatic biopsy infection rates. Urology 2012;79:556–63.
Chang DTS, Challacombe B, Lawrentschuk N. Transperineal prostate biopsy of native and recurrent prostate cancer. Clin Genitourin Cancer 2017;285:343–56. https://doi.org/10.1016/j.clgc.2016.07.007.
[65] Moran BJ, Braccioforte MH, Conterato DJ. Re-biopsy of the prostate using a stereotactic transperineal technique. J Urol 2006;176:1376–81.

[66] Pepdjonovic L, Tan GH, Huang S, Mann S, Frydenberg M, Moon D, et al. Zero hospital admissions for infection after 577 transperineal prostate biopsies using single-dose cephalozolin prophylaxis. World J Urol 2017;35:1199–203.

[67] Stefanova V, Buckley R, Flax S, Spevack L, Hajek D, Tunis A, et al. Transperineal prostate biopsies using local anesthesia: experience with 1,287 patients. Prostate cancer detection rate, complications and patient tolerability. J Urol 2019;201:1121–6.

[68] Gross MD, Shoag JE, Hu JC. Is in-office transperineal biopsy the future of prostate cancer diagnosis? Curr Opin Urol 2019; 29:25–6.

[69] Kubo Y, Kawakami S, Numao N, Takazawa R, Fujii Y, Masuda H, et al. Simple and effective local anesthesia for transperineal extended prostate biopsy: application to three-dimensional 26-core biopsy. Int J Urol 2009;16:420–3.

[70] Thurtle D, Starling L, Leonard K, Stone T, Gnanapragasam VJ. Improving the safety and tolerability of local anaesthetic outpatient transperineal prostate biopsies: a pilot study of the CAMbridge PROstate Biopsy (CAMPROBE) method. J Clin Urol 2018;11:192–9.

[71] Radtke JP, Kuru TH, Boxier S, Alt CD, Popencicu IV, Huettenbrink C, et al. Comparative analysis of transperineal template saturation prostate biopsy versus magnetic resonance imaging targeted biopsy with Magnetic Resonance Imaging-Ultrasound Fusion guidance. J Urol 2015;193:87–94.

[72] Hata N, Jinzaki M, Kacher D, Cormak R, Gering D, Nabavi A, et al. MR imaging-guided prostate biopsy with surgical navigation software: device validation and feasibility. Radiology 2001;220:263–8.

[73] Tilak G, Tuncali K, Song SE, Tokuda J, Olubiyi O, Fennessy F, et al. 3T MR-guided in-bore transperineal prostate biopsy: a comparison of robotic and manual needle-guidance templates. J Magn Reson Imaging 2015;42:63–71.

[74] Ho H, Yuen JSP, Mohan P, Lim EW, Cheng CWS. Robotic transperineal prostate biopsy: pilot clinical study. Urology 2011;78:1203–8.

[75] Altok M, Kim B, Patel BB, Shih YCT, Ward JF, McRae SE, et al. Cost and efficacy comparison of five prostate biopsy modalities: a platform for integrating cost into novel-platform comparative research. Prostate Cancer Prostatic Dis 2018;21:524–32.

[76] Venderink W, Govers TM, de Rooij M, Fütterer JJ, Sedelaar JPM. Cost-effectiveness comparison of imaging-guided prostate biopsy techniques: systematic transrectal ultrasound, direct in-bore MRI, and image fusion. Am J Roentgenol 2017;208:1058–63.

[77] De Rooij M, Crienen S, Witjes JA, Barentsz JO, Rovers MM, Grutters JPC. Cost-effectiveness of magnetic resonance (MR) imaging and MR-guided targeted biopsy versus systematic transrectal ultrasound-guided biopsy in diagnosing prostate cancer: a modelling study from a health care perspective. Eur Urol 2014;66:430–6.

[78] Boesen L, Nørgaard N, Løgager V, Balslev I, Bisbjerg R, Thestrup KC, et al. Assessment of the diagnostic accuracy of biparametric magnetic resonance imaging for prostate cancer in biopsy-naïve men: the biparametric MRI for detection of prostate cancer (BIDOC) study. JAMA Netw Open 2018;1:e180219. https://doi.org/10.1001/jamanetworkopen.2018.0219.

[79] Washino S, Okochi T, Saito K, Konishi T, Hirai M, Kobayashi Y, et al. Combination of prostate imaging reporting and data system (PI-RADS) score and prostate-specific antigen (PSA) density predicts biopsy outcome in prostate biopsy naïve patients. BJU Int 2017;119:225–33.

[80] Ullrich T, Quentin M, Arsov C, Schmaltz AK, Tschischka A, Laqua N, et al. Risk stratification of equivocal lesions on multiparametric magnetic resonance imaging of the prostate. J Urol 2018;199:691–8.

[81] Distler FA, Radtke JP, Bonekamp D, Kesch C, Schlemmer HP, Wieczorek K, et al. The value of PSA density in combination with PI-RADS™ for the accuracy of prostate cancer prediction. J Urol 2017;198:975–82.