



Biparametric MRI of the prostate

Prostatın Biparametrik MRG'si

Michele Scialpi¹, Alfredo D'Andrea², Eugenio Martorana³, Corrado Maria Malaspina¹, Maria Cristina Aisa⁴, Maria Napoletano¹, Emanuele Orlandi¹, Valeria Rondoni¹, Pietro Scialpi⁵, Diamante Pacchiarini⁶, Diego Palladino⁷, Michele Dragone⁷, Giancarlo Di Renzo^{4,8}, Annalisa Simeone⁷, Giampaolo Bianchi⁹, Luca Brunese¹⁰

Cite this article as: Scialpi M, D'Andrea A, Martorana E, Malaspina C, Aisa MC, Napoletano M, et al. Biparametric MRI of the prostate. Turk J Urol 2017; 43(4): 401-9

¹Department of Surgical and Biomedical Sciences, Division of Radiology 2, Santa Maria della Misericordia Hospital, Perugia University, Sant' Andrea delle Fratte, Perugia, Italy

²Department of Experimental Medicine, Magrassi Lanzara, Luigi Vanvitelli, Second University of Naples, Naples, Italy

³Department of Urology, Clinica Castelli, Bergamo, Italy

⁴Department of Surgical and Biomedical Sciences, Division of Gynaecology, Santa Maria della Misericordia Hospital, Perugia University, Sant' Andrea delle Fratte, Perugia, Italy

⁵Division of Urology, Portogruaro Hospital, Venice, Italy

⁶Health Management, S. Maria della Misericordia Hospital, Sant' Andrea delle Fratte, Perugia, Italy

⁷Department of Radiology, Casa Sollievo della Sofferenza Hospital, Foggia, Italy

⁸3DIFIC, Medical Area, University of Perugia, Perugia, Italy

⁹Division of Urology, University of Modena, Modena, Italy

¹⁰Department of Radiology, Campobasso University, C.da Tappino, Campobasso, Italy

Submitted:
13.10.2017

Accepted:
27.10.2017

Correspondence:
Michele Scialpi
E-mail:
michelescialpi1@gmail.com

©Copyright 2017 by Turkish Association of Urology

Available online at
www.turkishjournalofurology.com

ABSTRACT

Biparametric Magnetic Resonance Imaging (bpMRI) of the prostate combining both morphologic T2-weighted imaging (T2WI) and diffusion-weighted imaging (DWI) is emerging as an alternative to multiparametric MRI (mpMRI) to detect, to localize and to guide prostatic targeted biopsy in patients with suspicious prostate cancer (PCa). BpMRI overcomes some limitations of mpMRI such as the costs, the time required to perform the study, the use of gadolinium-based contrast agents and the lack of a guidance for management of score 3 lesions equivocal for significant PCa. In our experience the optimal and similar clinical results of the bpMRI in comparison to mpMRI are essentially related to the DWI that we consider the dominant sequence for detection suspicious PCa both in transition and in peripheral zone. In clinical practice, the adoption of bpMRI standardized scoring system, indicating the likelihood to diagnose a clinically significant PCa and establishing the management of each suspicious category (from 1 to 4), could represent the rationale to simplify and to improve the current interpretation of mpMRI based on Prostate Imaging and Reporting Archiving Data System version 2 (PI-RADS v2). In this review article we report and describe the current knowledge about bpMRI in the detection of suspicious PCa and a simplified PI-RADS based on bpMRI for management of each suspicious PCa categories to facilitate the communication between radiologists and urologists.

Keywords: Biparametric magnetic resonance imaging; prostate cancer; prostate imaging reporting and data system.

ÖZ

Prostatın biparametrik Manyetik Rezonans Görüntülemesi (bpMRG) hem morfolojik T2-ağırlıklı görüntüleme (T2W) hem de difüzyon ağırlıklı görüntülemeyi (DAG) kombine eden ve kuşku prostat kanseri (PKa) hastalarında, prostat kanserini belirleme, lokalize etme ve prostat hedefli biyopsiyi yönlendirmede multiparametrik MRG'ye (mpMRG) alternatif olarak ortaya çıkan bir yöntemdir. BpMRG, mpMRG'nin maliyet, çalışmayı uygulamak için gerekli süre, gadolinyum bazlı kontrast maddelerin kullanımı ve önemli PKa için kuşku skor 3 lezyonlarının tedavisini yönlendirememesi gibi bazı kısıtlamalarının üstesinden gelmektedir. Deneyimimize göre mpMR'e kıyasla bpMRG ile benzer ve optimal klinik sonuçlar esasen hem transiyonel hem de periferik zonda kuşku PKa'nın saptanmasında dominant sekans olduğunu düşündüğümüz DAG ile ilişkilidir. Klinik uygulamada, bpMRG ile standardize edilmiş skorlama sisteminin benimsenmesi, klinik açıdan önemli PKa'ya tanı koyma olasılığına işaret eder. Her bir kuşku kategorinin (1 ile 4 arası) tespiti Prostat Görüntüleme, Raporlama, Arşivleme Veri Sisteminin 2. sürümüne (PI-RADSV2) dayanarak mpMRG'nin güncel yorumlanmasını basitleştirme ve iyileştirme gereğini temsil edebilir. Bu derleme makalesinde radyologlar ve ürologlar arasındaki iletişimi kolaylaştırmak için kuşku PKa'yı tespit etme ve her bir kuşku PKa'nın tedavisi için bpMRG temelinde basitleştirilmiş PI-RADS'e ilişkin güncel bilgileri bildirmekte ve tanımlamaktayız.

Anahtar Kelimeler: Prostat kanseri; biparametrik manyetik rezonans görüntüleme; prostat görüntüleme raporlama ve veri sistemi.

Introduction

Prostate cancer (PCa) is the most common cancer among men and the second and third leading cause of cancer-related death in the United States and Europe, respectively.^[1-3] Accurate

detection and localization of the suspicious PCa is of paramount importance for appropriate management. Diagnostic algorithm based on several parameters [digital rectal examination, serum prostate-specific antigen (PSA) and transrectal ultrasound (TRUS)-guided prostate

biopsy and nomograms] provides limited information for this purpose.^[4]

Magnetic resonance imaging (MRI) of the prostate with T2-weighted imaging (T2WI) was first described in the mid-1980s.^[5,6] T2WI suffered from relatively poor sensitivity and specificity for detecting PCa and showed moderate accuracy in loco-regional staging.^[7] T2WI and dynamic contrast-enhanced (DCE) MRI protocol with endorectal coil (ERC) have not improved the diagnostic performance of MRI in the detection of PCa especially in the transition zone (TZ). A sensitivity of 55-88% and 46-90% and a specificity of 67-82% and 74-96% were reported by different authors for T2WI and DCE-MRI, respectively.^[8-10] The introduction of diffusion-weighted imaging (DWI) by measurement of the restriction of water diffusion of suspicious lesions has significantly improved the detection and localization of PCa in clinical practice.^[11-13]

State-of-the-art high field strength (1.5T and 3T) MRI technologies in multiparametric MRI (mpMRI) of the prostate (incorporating T2WI, DWI and DCE sequences) represents the most comprehensive diagnostic approach for PCa and it is actually considered the reference standard imaging modality in detecting, localizing and staging prostate cancer.^[14-17] MpMRI interpretation is based on Prostate Imaging and Reporting Archiving Data System version 2 (PI-RADS v2)^[18], introduced to improve standardization of MRI interpretation by reporting suspicion scores (1- to 5-point scale based on fixed criteria) to provide clinical management guideline for suspicious PCa.

MpMRI demonstrated that PCa is often multifocal and increased the sensitivity of conventional MRI in the detection of PCa, especially involving TZ and anterior fibromuscular stroma.^[19,20] MpMRI has some limitations such as its higher cost, longer time required to complete the study and the use of gadolinium-based contrast agents. Moreover, PI-RADS v2 does not offer a clear guidance for clinical management of PI-RADS score 3 lesions.^[18] Biparametric MRI (bpMRI), including T2W and DW MRI series, has been recently proposed to overcome these limitations and to offer a more appropriate management of PCa.^[21-28]

The benefits of bpMRI are represented by the reduced costs, shorter time required to complete the study, and the no-use of gadolinium. In addition, bpMRI provides similar results to those of mpMRI for the detection and localization of PCa.^[24,29]

Different studies have also demonstrated the validity of the lesion volume as predictor of PCa and aggressiveness of the lesion.^[30] Several authors have also reported a good correlation between radiologic estimation of the lesion volume and histology of the tumor volume.^[19,25,30]

A potential of bpMRI is represented by the management (biopsy or clinical surveillance) of the score 3 lesions through the calcu-

lation of their volume (cut-off of 0.5 cm³) which is consequently a simplification of PI-RADS v2. In this article we review the bpMRI of the prostate and its performance in the detection, localization and management of PCa.

Clinical history

The following information should be available before MRI acquisition and interpretation in men with suspicious PCa: PSA values (recent, and previous), family history for PCa, persistently, and highly suspicious prostatic lesion despite negative standard 12-core biopsy, previous diagnosis of High Grade Prostatic Intraepithelial Neoplasia (HGPIN) or Atypical Small Acinar Proliferation (ASAP).

Biparametric MRI implementation and technical requirements

A diagnostic prostate bpMRI is achievable at 1.5T field strength, even when not using ERC or preferentially by 3T system if available. In our 6 years' experience, bpMRI has been performed using 3T scanner (Achieva, Philips Medical Systems, Healthcare, Eindhoven, the Netherlands) with 16-channels torso phased-pelvic array. Imaging with high signal to noise ratio (SNR) of the entire gland, capsule and neurovascular bundles is obtained, and lesions have been accurately detected, and localized. Patients with contraindications for MRI have to be excluded. When possible, patients should evacuate the rectum just prior to the MRI exam. Administration of antispasmodics such as butylscopolamine may be helpful to reduce intestinal peristalsis.

T2-weighted imaging

High-resolution T2WI turbo-spin echo (TSE) allows morphological informations depicting prostate gland anatomy [peripheral zone (PZ), TZ, capsule and neurovascular bundle] and any suspicious lesions. The T2WI-TSE sequence is acquired in the axial, sagittal and coronal planes. It allows measurement of both prostate and lesion volume, localizes suspicious PCa within the gland and guides targeted biopsy. In addition, sagittal and coronal planes are useful for evaluating extracapsular extension (ECE) of PCa (e.g. seminal vesicle invasion).

On T2WI, PCa localized in PZ appears predominantly and distinctly hypointense compared to the hyperintense glandular tissue of the PZ^[31], and when ECE is recognizable on T2WI, it must be interpreted as a reliable sign of malignancy. In TZ different but morphologically similar entities (e.g. post-inflammatory or post-biopsy scars, atrophic changes, prostatitis, granulomatous prostatitis, intraepithelial neoplasia (PIN) or post-treatment lesions and often benign prostatic hyperplasia [BPH])^[32-34] that appear as focal hypointense areas, may mimic PCa.

Post-biopsy hemorrhages (generally persisting for several months after biopsy, so it is suggested an interval of at least six weeks before performing MRI)^[18] also appear hypointense on T2WI

but hyperintense on fat-suppressed T1-weighted gradient-echo sequence with High-Resolution Isotropic Volume Excitation (THRIVE). In addition, an axial TSE T2W high resolution spectral pre-saturation inversion recovery (HR SPIR) sequence, using an extended field-of-view (FOV), allows the detection of enlarged parailiac and locoregional lymph nodes and/or skeletal metastases.

PI-RADS v2 considers T2WI the dominant sequence in the detection of PCa localized in TZ. T2WI is performed using an echo time (TE) of 100-120 ms and a long repetition time (TR) of 4000 – 8000 ms (depending on the equipment and B0 field strength), a minimum slice thickness of 3 mm at 1.5T or at 3T with no gap, and a minimum in-plane resolution of 0.7×0.7 mm for both field strengths. In practice we perform T2WI acquisitions on axial, coronal and sagittal planes to detect more rapidly the lesions and to assess extraglandular extension of disease. On our 3T system a FOV approximately of 18 cm, with matrices of 212×212 (frequency \times phase) and slice thicknesses of 2 mm with no gap is used.

Diffusion-Weighted Imaging

DWI analyzes the movement (diffusion) of water molecules and expresses it by a parameter known as the apparent diffusion coefficient (ADC). Restriction of water diffusion in PCa, due to numerous intact cell membranes and scarce extracellular spaces, with a resultant decrease in the ADC value, is measured. PCa shows lower ADC or restricted diffusion surrounding healthy prostate tissue and appears hyperintense on the diffusion-weighted (DWI) images with high b-value and hypointense on ADC maps.^[35,36] The different numbers and magnitudes of the selected b-values may determine variations in the ADC results related to different field strengths.^[37]

Although several studies showed the potential of quantitative ADC values for differentiating high Gleason score (GS) from low GS PCa, an overlapping between benign and malignant lesions exists and ADC values for this differentiation vary among different MR systems. PI-RADS v2^[18] considers DWI the dominant sequence to identify tumors in PZ, while in our experience we

have considered DWI the dominant sequence for detection of PCa in both TZ and PZ and only visual analysis of lesions on ADC map has been performed.

DWI is generally performed with a single-shot echo-planar (SS-EPI) sequence in the same axial orientation of the T2WI. Diffusion gradients should be applied in 3 orthogonal spatial directions. A minimum of 3, ideally 5, b-values > 1000 s/mm² should be used. TE should be as short as possible (typically <90 ms). The sequence is prone to susceptibility artefacts that may lead to distortions of the DW images due to adjacent bowel gas. The measurement of restricted diffusion in tumor tissue using high b-values improves diagnostic accuracy of MRI for PCa.

Minimum slice thickness of 3 mm with no gap and 2 number of excitations (NEX) should be used. B-values 0-2000 s/mm² with ADC map are generated. According to PI-RADS v2, a maximum b-value of 2000 s/mm² may be used. DWI with very high b-values may be challenging in some systems due to excessive loss of SNR, particularly if imaging at 1.5T or without using ERC is obtained.^[38]

On our 3T system we perform DWI acquisitions on axial plane, slice thicknesses of 2 mm with no gap using 124×100 (frequency \times phase) matrices, a FOV of 28-32 cm, and 4 b-values 0-2000 s/mm² with reconstruction of ADC maps. BpMRI acquisition protocol and sequence parameters are presented in Table 1.

Biparametric vs multiparametric MRI in the detection of prostate cancer

PI-RADS v2 and several studies^[18,20,27,28,39] recognized that the data supporting the role of DCE are limited and the added value provided by DCE to T2WI and DWI appears to be modest. Costs, time required to perform the study, gadolinium-based contrast media potential risk in patients with impaired renal function and its accumulation in the central nervous system represent the major drawbacks of DCE.^[40]

The value of bpMRI in the detection of PCa using 1.5T MRI with endorectal coil has been reported.^[23,41] Recently, it has been dem-

Table 1. Biparametric MRI of the prostate at 3T: acquisition protocol and sequence parameters

	Sequences	ST (mm)	Gap	TR (ms)	TE (ms)	FOV (8 mm)	Voxel size
DWI	2D-EPI	2	0	3700	67	280x320	1.46x1.45x2
T2W Axial	TSE	2	0	12700	90	180x180	0.28x0.28x2
T2W Coronal	TSE	2	0	8400	90	180x180	0.28x0.28x2
T2W Sagitta	TSE	2	0	8400	90	180x180	0.28x 0.28x2
T1W Axial	THRIVE	1.5	0	3.0	1.43	300x370	1.46x1.46x1.5

DWI: diffusion-weighted imaging; EPI: echo planar imaging; TSE: turbo spin echo; THRIVE: T1 High-Resolution Isotropic Volume Excitation; ST: slice thickness; TR: time of repetition; TE: time of echo; FOV: field of view

Table 2. Diagnostic accuracy and sensitivity of biparametric and multiparametric MRI in the detection prostate cancer at 3T without endorectal coil: literature review

Authors	Year	bpMRI accuracy	mpMRI accuracy	bpMRI sensitivity	mpMRI sensitivity
Radtke et al. ^{[20]*}	2015	-	-	91.9%	86.4-88.5%
Fascelli et al. ^[28]	2016	81.4%**	-	95.5%	-
Thestrup et al. ^[42]	2016	41.5%	39%	94.6%	93-100%
Stanzione et al. ^[44]	2016	92.7%	93.9%	83.5%	91.1%
Scialpi et al. ^{[29]**}	2017	99.4%	99.4%	98.2%	98.2%
De Visschere et al. ^[27]	2017	72.2-74.7%	72.9%	61.8-72.2%	72.9%
Kuhl et al. ^[43]	2017	89.1%	87.2%	93.9%	84.6%

* Anterior lesions GS \geq 3+4 and GS \geq 4+3; **Overall accuracy (bpMRI and PSA); index lesions \geq 10mm.

onstrated that the sensitivity of bpMRI at 3T without endorectal coil in the detection PCa is similar to that of mpMRI.^[20,24,27,28,42-44] The diagnostic value of bpMRI in men with or without prior biopsy and combined with PSA has been validated, resulting in an improved accuracy for detecting clinically significant PCa (sPCa) and representing an effective technique to direct biopsy needles under TRUS guidance, after MRI-ultrasonography fusion.^[20,23,28]

For diagnosis of clinically sPCa in patients with elevated PSA before biopsy, DCE was found to be an unnecessary imaging technique in the majority of patients evaluated by PI-RADS v2.^[27,42] In our experience, the high sensitivity of DWI/ADC associated with T2WI, corresponding to the performance of DWI alone (97.6%, 100% and 94.7% for, respectively, all lesions, PZ and TZ), was equal to that of mpMRI for lesions in PZ, TZ and in the anterior fibromuscular stroma.^[24] The sensitivity rates of bpMRI and mpMRI at 3T without ERC in the detection of PCa are 61.8-97.6%, and 72.9-97.6%, respectively (Table 2).^[20,24,27,28,42-44]

Biparametric MRI: lesion detection and localization

According to the criteria and lexicon of the PI-RADS v2 guidelines^[18], the bpMRI images' analysis is based on the recognition of lesion patterns on T2WI and DWI and corresponding ADC maps. Because in our experience DWI represents the predominant sequence to detect lesions both in PZ and TZ with confirmed image quality, for routine clinical work we recommend starting viewing DWI with high b-values (and the same sequence inverted) on ADC maps and subsequently at T2WI.

The index lesion is detected as a focal area with hyperintensity on DWI with high b-values and mild/moderate or marked hypointensity on ADC maps, and moderate or marked hypointensity on T2WI. A standard bpMRI layout of our prostate image display is shown in Figure 1.

Prostate is analyzed from the seminal vesicles to the apex for the detection of focal lesions localized in the PZ and TZ. Findings

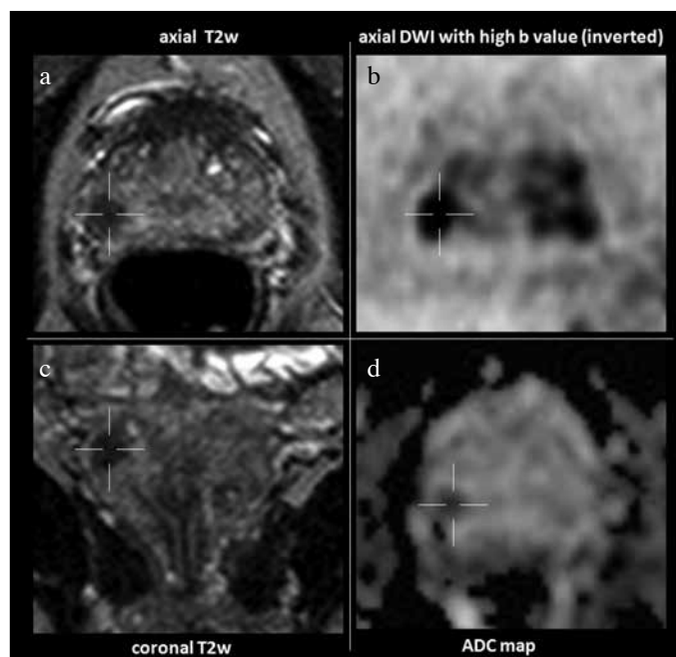


Figure 1. a-d. Standard layout of prostate image display in a one screen setting. Double clicking in one of the image allows to display this image in full screen mode. On the left side the morphologic axial and sagittal T2W images and on the right side DWI with high b-values (inverted) and ADC map are displayed. Ideally the morphologic T2W and the DW images should be linked at the same table position. With this described display mode, detection and localization of the suspected lesions (hypointense on T2W, hyperintense on DWI with high b-values or hypointense on inverted with corresponding decrease on ADC map), can already be carried out.

based on bpMRI are described and at maximum 4 index lesions are identified and localized according to the 39 sector/region- prostate map model reported in PI-RADS v2^[18] within approximately 15 minutes. In cases with significant enlargement of TZ and presence of median lobe, to localize lesions in this side, we propose a 41

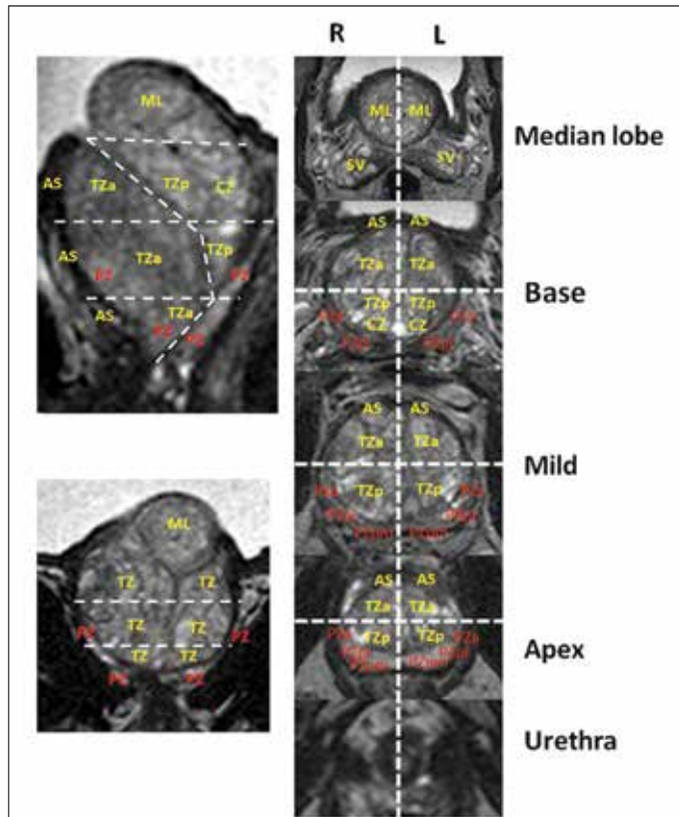


Figure 2. The 41 segmentation model proposed for prostate with enlargement of transition zone and presence of median intravesical lobe: fourthy-one sectors/regions: thirty-eight for the prostate, two for the seminal vesicles and one for the external urethral sphincter are identified. In comparison to 39 segmentation model proposed by PI-RADS v2, the 41 segmentation model adds two segments (right and left) of an intravesical median lobe.

TZ: transition zone; CZ: central zone; PZ: peripheral zone; AS: anterior stroma; SV: seminal vesicle; ML: median lobe.

sector/region- prostate map model, by adding two segments (right and left) of an intravesical median lobe (Figure 2).

Biparametric MRI volume calculation for the management of score 3 lesions

One of the major drawback of PI-RADS v2 is that it does not offer a precise guidance on clinical management (biopsy or clinical surveillance) of score 3 lesions (equivocal for clinical sPCa). Some authors biopsise all PI-RADS score 3 lesions^[44] leading to increased number of unnecessary biopsies, on the other hand, others have demonstrated that PI-RADS score 3 lesions are associated with low likelihood of clinically sPCa, concluding that these lesions should not be sampled but monitored only.^[45] Several studies^[25,29] have demonstrated the adequacy of MRI lesion volume estimation compared to histopathological tumor

volume. According to the Epstein criteria^[46], the presence of sPCa should be predicted by lesion volume measurement, using a cut-off value of 0.5 cm³.

We discriminated two subgroups as 3a (<0.5 cm³) and 3b (>0.5 cm³) PI-RADS v2 3 score lesions (lesions both in PZ and in TZ, moderately hypointense on T2, hyperintense on high b-value DWI and mildly/moderately hypointense on ADC map)^[47], proposed a simplified PI-RADS (S-PI-RADS) score based on bpMRI and focused on the clinical management of PI-RADS v2 score 3 lesions. S-PI-RADS scoring system consists of 4 lesion categories (from 1 to 4), considers volumes of all score 3 lesions and includes all PI-RADS v2 score 4 and 5 lesions within a single risk category (score 4).

Lesion volume for score 3 lesions should be calculated using the ellipsoidal formula (width x height x length x 0.52) or alternatively using a software (Mimics Innovation Suite, Materialise NV, Leuven, Belgium) with three-dimensional (3D) reconstruction, drawing a freehand region of interest (ROI) around the discernible index lesion on the inverted DWI with a high b-value (Figure 3).

S-PI-RADS: communication of findings and management

In our experience, the adoption of S-PI-RADS based on bpMRI of suspicious lesions offers the advantage of an easier communication of findings between radiologists and urologists and a precise indication for the management of each risk category. The S-PI-RADS scheme adapted to bpMRI and clinical management of each category is presented in Figure 4.

S-PI-RADS category 1 corresponds to PI-RADS v2 score 1 lesion: moderately homogeneous signal intensity (normal) on T2WI and no abnormality on DWI/ADC. Follow-up with PSA is indicated.

S-PI-RADS category 2 corresponds to PI-RADS v2 score 2 lesion: circumscribed hypointense area on T2WI without restriction on diffusion on DWI/ADC. Follow-up with PSA and bpMRI eventually within 2 years is indicated.

S-PI-RADS category 3 includes PI-RADS v2 score 3 lesion: focal rounded, lenticular or irregular lesion moderate heterogeneous or homogeneous hypointense on T2WI, mild/moderate hyperintense on DWI with high b-values and marked hypointense on ADC map. For lesions with volume <0.5 cm³ (subgroup a) accurate evaluation of age and clinical information, periodic monitoring of PSA value and repetition of bpMRI 1 year later is suggested. This approach is related to the fact that indolent PCa remains stable over time from diagnosis^[48] and lesions with volume <0.5 cm³ include a minimal percentage of GS score 7 adenocarcinoma (about 2%).^[30] A score 3 lesion becomes suitable of bioptic check if its volume reaches 0.5 cm³ during the follow-up, shifting from category 3 to 4. For lesions with volume >0.5 cm³ targeted biopsy is indicated.

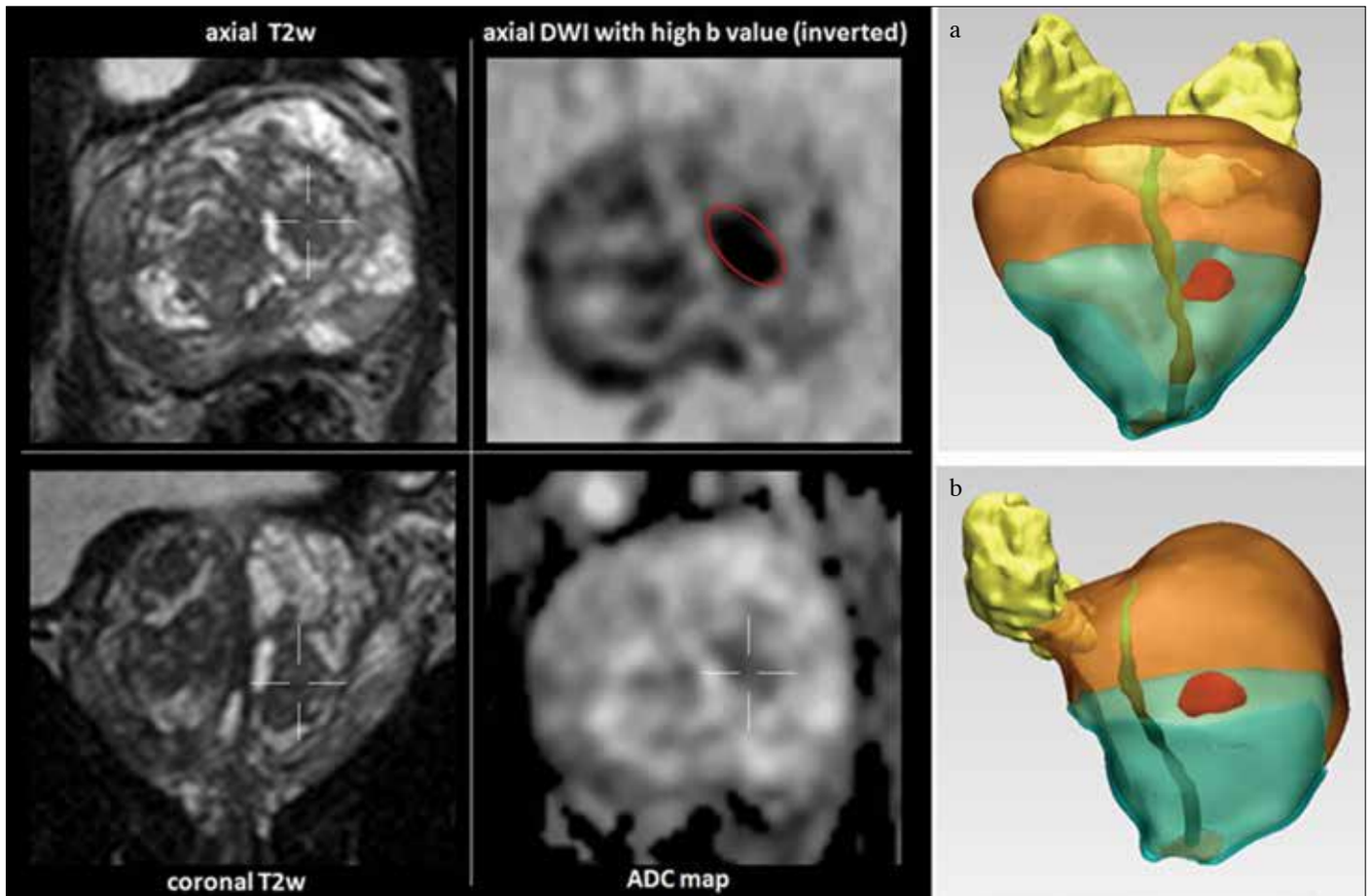


Figure 3. a, b. Lesion score 3 lesions detected in the transition zone at mild of the prostate on the left side on morphologic axial and coronal T2W images and DWI with high b-values and ADC map. Three-dimensional (3D) reconstruction coronal (a) and oblique (b) is obtained by drawing a freehand region of interest (ROI) around the discernible index lesion on the inverted DWI with high b-value

S-PI-RADS category 4 includes PI-RADS v2 score 4 and 5 lesions. Heterogeneous or homogeneous mild/moderate or markedly hypointense focal rounded, lenticular or irregular lesions on T2WI, hyperintense on high b-value DWI and marked hypointense on ADC map. PI-RADS v2 score 4 are intraglandular lesions and PI-RADS v2 score 5 are lesions extending outside the prostate or bulging the capsule and/or invading the seminal vesicles. Targeted biopsy is indicated.

Thereafter, the entire pelvis should be analyzed for lymph node involvement, bone metastases and other findings. As reported, especially DW sequences of the entire pelvis in combination with meticulous analysis of T2W STIR sequence help to detect lymph node metastases.^[49]

At our institution we formulate the report in a structured way on a free text basis. The bpMRI report includes clinical notes, technical details, findings [prostate volume, detection and localization of the lesion (at maximum 4 lesions are indicated), volume calculation for score 3 lesions, assessment of the lesion's category

and its management]. Besides possible ECE and infiltration of the neurovascular bundle and seminal vesicles as well as infiltration of adjacent organs, exact location, number and size of possible lymph node metastases in the entire pelvis, and bone metastases are also recorded.

The knowledge of already categorized and reported pitfalls (normal anatomic structures, noncancerous abnormalities and technical challenges related to DW sequences) in PCa,^[50] may aid the radiologist to reduce false-positive results obtained with bpMRI. These pitfalls, of which the radiologist have to be aware of, are presented in Table 3.

Conclusion

BpMRI (morphological T2WI and DWI) allows an accurate detection and localization of suspicious PCa, in addition to reduction of the time required to complete the study, lower costs without using gadolinium. BpMRI and lesion volume measurement for PI-RADS score 3 lesions discriminate 4 categories of lesions

Category	T2WI	DWI with high b value	DWI with high b value (inverted)	ADC	Management
1					Follow-up by PSA
2					Follow-up by PSA and bpMRI eventually within 2 years
3a					Lesion volume <math>< 0.5 \text{ cm}^3</math>: monitoring of PSA value and repeat bpMRI 1 year later
3b					Lesion volume $\geq 0.5 \text{ cm}^3$: targeted biopsy
4					Targeted biopsy

Figure 4. Simplified PI-RADS (S-PI-RADS) based on biparametric MRI

S-PI-RADS category 1: homogeneous intermediate signal intensity (normal) on T2WI and no abnormality on DWI/ADC. Follow-up by PSA is indicated.

S-PI-RADS category 2: focal rounded, lenticular or irregular mild/moderately or markedly hypointensity area on T2WI without restriction on diffusion on DWI/ADC. Follow-up by PSA and biparametric MRI eventually within 2 years is indicated.

S-PI-RADS category 3: includes 1): subgroup a (lesion volume: $< 0.5 \text{ cm}^3$: round lesion moderately hypointense on T2WI (arrow), hyperintense on high b-value DWI (arrow) and inverted (arrow), and moderately hypointense on ADC map (arrow): accurate evaluation of age, clinical informations, periodic monitoring of PSA value and repetition of bpMRI 1 year later is suggested; 2) subgroup b (lesion volume: $> 0.5 \text{ cm}^3$): lenticular lesion moderately hypointense on T2WI (arrow), hyperintense on high b-value DWI (arrow) and inverted (arrow), and moderately hypointense on ADC map (arrow). Targeted biopsy is indicated.

S-PI-RADS category 4: includes PI-RADS v2 score 4 (intraglandular lesions) and 5 (lesions extending to an area outside of the prostate or bulging the capsule of the prostate and/or invasion of the seminal vesicles lesions). Focal lenticular lesion markedly hypointense on T2WI (arrow), hyperintense on high b-value DWI (arrow) and inverted (arrow), and markedly hypointense on ADC map (arrow). Targeted biopsy is indicated.

Table 3. Pitfalls that can mimic prostate cancer on biparametric MRI

Normal anatomic structures	Normal central zone or periprostatic venous plexus or neurovascular bundle can mimic peripheral zone lesions. Bilateral T2w image hypointensities at the base and/or median posterior T2w image hypointense area in the middle third of the gland can mimic peripheral zone lesions. Asymmetric thickening of the surgical capsule can be difficult to distinguish from a focal lesion.
Noncancerous abnormalities	Post-biopsy haemorrhage. Differentiation between stromal benign prostatic hyperplasia and a transition zone tumour. Benign prostatic hyperplasia (BPH) nodule protrusion (can mimic a peripheral zone lesion). Acute and chronic prostatitis, post-inflammatory scars, granulomatous prostatitis and atrophy.
Technical challenges related to DWI	Anatomic distortion or lack of suppression of benign prostate tissue on standard high b-value DWI can lead to false positive findings. Suboptimal windowing of ADC maps can determine false negative results.

for each of which a corresponding decision-making is indicated. In clinical practice the adoption of S-PI-RADS based on bpMRI represents a potential valid system that facilitates lesion management and communication with other professional colleagues.

Peer-review: This manuscript was prepared by the invitation of the Editorial Board and its scientific evaluation was carried out by the Editorial Board.

Author Contributions: Concept – M.S., A.D.A., E.M., C.M.M., A.S., G.B., L.B.; Design – M.S., A.D.A., E.M., C.M.M.; Supervision – M.S., A.D.A., E.M., C.M.M., A.S., D.P.; Data Collection and/or Processing – M.S., A.D.A., E.M., C.M.M., A.S., D.P., M.D.; Analysis and/or Interpretation – M.S., A.D.A., E.M., C.M.M.; Literature Search – M.S., C.M.M., M.N., E.O., V.R., M.C.A., D.P.; Writing Manuscript – M.S., A.D.A., E.M., C.M.M., L.B.; Critical Review – M.S., A.D.A., E.M., C.M.M., P.S., M.N., V.R., G.D.R., M.C.A., D.P., G.B., L.B.

Acknowledgements: The Authors thank for 3DIFIC the Mr Marco Pergentini for his contribute in 3D reconstruction, and the technologist Domenico Mezzasoma for his assistance in the evaluation of MRI sequence parameters.

Conflict of Interest: No conflict of interest was declared by the authors.

Financial Disclosure: The authors declared that this study has received no financial support.

Hakem Değerlendirmesi: Bu makale Editörler Kurulu'nun davetiyle hazırlandığından bilimsel değerlendirme Editörler Kurulu tarafından yapılmıştır.

Yazar Katkıları: Fikir – M.S., A.D.A., E.M., C.M.M., A.S., G.B., L.B.; Tasarım – M.S., A.D.A., E.M., C.M.M.; Denetleme – M.S., A.D.A., E.M., C.M.M., A.S., D.P.; Veri Toplanması ve/veya İşlemesi – M.S., A.D.A., E.M., C.M.M., A.S., D.P., M.D.; Analiz ve/veya Yorum – M.S., A.D.A., E.M., C.M.M.; Literatür Taraması – M.S., C.M.M., M.N., E.O., V.R., M.C.A., D.P.; Yazıyı Yazan – M.S., A.D.A., E.M., C.M.M., L.B.; Eleştirel İnceleme – M.S., A.D.A., E.M., C.M.M., P.S., M.N., V.R., G.D.R., M.C.A., D.P., G.B., L.B.

Teşekkür: Yazarlar 3DIFIC ve 3 Boyutlu rekonstrüsyona katkısı için Bay Marco Pergentini'ye, MRG dizin parametrelerinin değerlendirmesinde yardımcı için teknolog Domenico Mezzasoma'ya teşekkür ederiz.

Çıkar Çatışması: Yazarlar çıkar çatışması bildirmemişlerdir.

Finansal Destek: Yazarlar bu çalışma için finansal destek almadıklarını beyan etmişlerdir.

References

1. Siegel R, Naishadham D, Jemal A. Cancer statistics, 2013. *CA Cancer J Clin* 2013;63:11-30. [Crossref]
2. Jemal A, Center MM, DeSantis C, Ward EM. Global patterns of cancer incidence and mortality rates and trends. *Cancer Epidemiol Biomarkers Prev* 2010;19:1893-907. [Crossref]
3. Ferlay J, Autier P, Boniol M, Heanue M, Colombet M, Boyle P, et al. Estimates of the cancer incidence and mortality in Europe in 2006. *Ann Oncol* 2007;18:581-92. [Crossref]
4. Fine SW, Amin MB, Berney DM, Bjartell A, Egevad L, Epstein JI, et al. A contemporary update on pathology reporting for prostate cancer: biopsy and radical prostatectomy specimens. *Eur Urol* 2012;62:20-39. [Crossref]
5. Poon PY, McCallum RW, Henkelman MM, Bronskill MJ, Sutcliffe SB, Jewett MA, et al. Magnetic resonance imaging of the prostate. *Radiology* 1985;154:143-9. [Crossref]
6. Hricak H, Dooks GC, Jeffrey RB, Avallone A, Jacobs D, Benton WK, et al. Prostatic carcinoma: staging by clinical assessment, CT, and MR imaging. *Radiology* 1987;162:331-6. [Crossref]
7. Tempny CM, Rahmouni AD, Epstein JI, Walsh PC, Zerhouni EA. Invasion of the neurovascular bundle by prostate cancer: evaluation with MR imaging. *Radiology* 1991;181:107-12. [Crossref]
8. Yoo S, Kim JK, Jeong IG. Multiparametric magnetic resonance imaging for prostate cancer: A review and update for urologists. *Korean J Urol* 2015;56:487-97. [Crossref]
9. Wu LM, Xu JR, Ye YQ, Lu Q, Hu JN. The clinical value of diffusion-weighted imaging in combination with T2-weighted imaging in diagnosing prostate carcinoma: a systematic review and meta-analysis. *AJR Am J Roentgenol* 2012;199:103-10. [Crossref]
10. Verma S, Turkbey B, Muradyan N, Rajesh A, Cornud F, Haider MA, et al. Overview of dynamic contrast-enhanced MRI in prostate cancer diagnosis and management. *AJR Am J Roentgenol* 2012;198:1277-88. [Crossref]
11. Bammer R. Basic principles of diffusion-weighted imaging. *Eur J Radiol* 2003;45:169-84. [Crossref]
12. Tan CH, Wang J, Kundra V. Diffusion weighted imaging in prostate cancer. *Eur Radiol* 2011;21:593-603. [Crossref]
13. Haider MA, van der Kwast TH, Tanguay J, Evans AJ, Hashmi AT, Lockwood G, et al. Combined T2-weighted and diffusion-weighted MRI for localization of prostate cancer. *AJR Am J Roentgenol* 2007;189:323-8. [Crossref]
14. Rais-Bahrami S, Siddiqui MM, Turkbey B, Stamatakis L, Logan J, Hoang AN, et al. Utility of multiparametric magnetic resonance imaging suspicion levels for detecting prostate cancer. *J Urol* 2013;190:1721-8. [Crossref]
15. Siddiqui MM, Rais-Bahrami S, Truong H, Stamatakis L, Vourganti S, Nix J, et al. Magnetic resonance imaging/ultrasound-fusion biopsy significantly upgrades prostate cancer versus systematic 12-core transrectal ultrasound biopsy. *Eur Urol* 2013;64:713-9. [Crossref]
16. Bloch BN, Genega EM, Costa DN, edrosa I, Smith MP, Kressel HY, et al. Prediction of prostate cancer extracapsular extension with high spatial resolution dynamic contrast-enhanced 3-T MRI. *Eur Radiol* 2012;22:2201-10. [Crossref]
17. de Rooij M, Hamoen EH, Fütterer JJ, Barentsz JO, Rovers MM. Accuracy of multiparametric MRI for prostate cancer detection: a meta-analysis. *AJR Am J Roentgenol* 2014;202:343-51. [Crossref]
18. Prostate Imaging Reporting and Data System (PI-RADS) Reston (VA): American College of Radiology. Available at: <http://www.acr.org/Quality-Safety/Resources/PIRADS/> Accessed March 5, 2015.
19. Radtke JP, Schwab C, Wolf MB, Freitag MT, Alt CD, Kesck C, et al. Multiparametric Magnetic Resonance Imaging (MRI) and MRI-Transrectal Ultrasound Fusion Biopsy for Index Tumor Detection: Correlation with Radical Prostatectomy Specimen. *Eur Urol* 2016;70:846-53. [Crossref]

20. Radtke JP, Boxler S, Kuru TH, Wolf MB, Alt CD, Popeneciu IV, et al. Improved detection of anterior fibromuscular stroma and transition zone prostate cancer using biparametric and multiparametric MRI with MRI-targeted biopsy and MRI-US fusion guidance. *Prostate Cancer Prostatic Dis* 2015;18:288-96. [\[Crossref\]](#)
21. Scialpi M, Falcone G, Scialpi P, D'Andrea A. Biparametric MRI: a further improvement to PIRADS 2.0? *Diagn Interv Radiol* 2016;22:297-8. [\[Crossref\]](#)
22. Scialpi M, Martorana E, D'Andrea A. Standardizing biparametric MRI to simplify and improve Prostate Imaging Reporting and Data System, version 2, in prostate cancer management. *AJR Am J Roentgenol* 2016;6:W1-2. [\[Crossref\]](#)
23. Rais-Bahrami S, Siddiqui MM, Vourganti S, Turkbey B, Rastinehad AR, Stamatakis L, et al. Diagnostic value of biparametric magnetic resonance imaging (MRI) as an adjunct to prostate-specific antigen (PSA)-based detection of prostate cancer in men without prior biopsies. *BJU Int* 2015;115:381-8. [\[Crossref\]](#)
24. Scialpi M, Prosperi E, D'Andrea A, Martorana E, Malaspina C, Palumbo B, et al. Biparametric versus Multiparametric MRI with Non-endorectal Coil at 3T in the Detection and Localization of Prostate Cancer. *Anticancer Res* 2017;37:1263-71. [\[Crossref\]](#)
25. Baco E, Ukimura O, Rud E, Vlatkovic L, Svindland A, Aron M, et al. Magnetic resonance imaging-transectal ultrasound image-fusion biopsies accurately characterize the index tumor: correlation with step-sectioned radical prostatectomy specimens in 135 patients. *Eur Urol* 2015;67:787-94. [\[Crossref\]](#)
26. Radtke JP, Schwab C, Wolf MB, reitag MT, Alt CD, Kesch C, et al. Multiparametric magnetic resonance imaging (MRI) and MRI-transrectal ultrasound fusion biopsy for index tumor detection: correlation. *Eur Urol* 2016;70:846-53. [\[Crossref\]](#)
27. De Visschere P, Lumen N, Ost P, Decaestecker K, Pattyn E, Villeirs G. Dynamic contrast-enhanced imaging has limited added value over T2-weighted imaging and diffusion-weighted imaging when using PI-RADSV2 for diagnosis of clinically significant prostate cancer in patients with elevated PSA. *Clin Radiol* 2017;72:23-32. [\[Crossref\]](#)
28. Fascelli M, Rais-Bahrami S, Sankineni S, Brown AM, George AK, Ho R, et al. Combined biparametric prostate magnetic resonance imaging and prostate-specific antigen in the detection of prostate cancer: a validation study in a biopsy-naive patient population. *Urology* 2016;88:125-34. [\[Crossref\]](#)
29. Scialpi M, Rondoni V, Aisa MC, Martorana E, D'Andrea A, Malaspina CM, et al. Is contrast enhancement needed for diagnostic prostate MRI? *Transl Androl Urol* 2017;6:499-509. [\[Crossref\]](#)
30. Martorana E, Pirola GM, Scialpi M, Micali S, Iseppi A, Bonetti LR, et al. Lesion volume predicts prostate cancer risk and aggressiveness: validation of its value alone and matched with prostate imaging reporting and data system score. *BJU Int* 2017;120:92-103. [\[Crossref\]](#)
31. Hricak H. Imaging prostate cancer. *J Urol* 1999;162:1329-30. [\[Crossref\]](#)
32. Barentsz JO, Richenberg J, Clements R, Choyke P, Verma S, Villeirs G, et al. ESUR prostate MR guidelines 2012. *Eur Radiol* 2012;22:746-57. [\[Crossref\]](#)
33. Oto A, Kayhan A, Jiang Y, Tretiakova M, Yang C, Antic T, et al. Prostate cancer: differentiation of central gland cancer from benign prostatic hyperplasia by using diffusion-weighted and dynamic contrast-enhanced MR imaging. *Radiology* 2010;257:715-23. [\[Crossref\]](#)
34. Kim CK, Park BK, Kim B. Localization of prostate cancer using 3T MRI: comparison of T2-weighted and dynamic contrast-enhanced imaging. *J Comput Assist Tomogr* 2006;30:7-11. [\[Crossref\]](#)
35. Zelhof B, Pickles M, Liney G, Gibbs P, Rodrigues G, Kraus S, et al. Correlation of diffusion-weighted magnetic resonance data with cellularity in prostate cancer. *BJU Int* 2009;103:883-8. [\[Crossref\]](#)
36. Sato C, Naganawa S, Nakamura T, Kumada H, Miura S, Takizawa O, et al. Differentiation of noncancerous tissue and cancer lesions by apparent diffusion coefficient values in transition and peripheral zones of the prostate. *J Magn Reson Imaging* 2005;21:258-62. [\[Crossref\]](#)
37. Mueller-Lisse UG, Mueller-Lisse UL, Zamecnik P, Schlemmer HP, Scherr MK. Diffusion-weighted MRI of the prostate. *Radiologe* 2011;51:205-14. [\[Crossref\]](#)
38. Wang X, Qian Y, Liu B, Cao L, Fan Y, Zhang JJ, et al. High-b-value diffusion-weighted MRI for the detection of prostate cancer at 3 T. *Clin Radiol* 2014;69:1165-70. [\[Crossref\]](#)
39. Barentsz JO, Weinreb JC, Verma S, Thoeny HC, Tempany CM, Shtern F, et al. Synopsis of the PI-RADS v2 Guidelines for Multiparametric Prostate Magnetic Resonance Imaging and Recommendations for Use. *Eur Urol* 2016;69:41-9. [\[Crossref\]](#)
40. Olchowy C, Cebulski K, Łasecki M, Chaber R, Olchowy A, Kawak K, et al. The presence of the gadolinium-based contrast agent depositions in the brain and symptoms of gadolinium neurotoxicity - A systematic review. *PLoS One* 2017;12:e0171704. [\[Crossref\]](#)
41. Delongchamps NB, Rouanne M, Flam T, euvon F, Liberatore M, Zerbib M, et al. Multiparametric magnetic resonance imaging for the detection and localization of prostate cancer: combination of T2-weighted, dynamic contrast-enhanced and diffusion-weighted imaging. *BJU Int* 2011;107:1411-8. [\[Crossref\]](#)
42. Thestrup KC, Logager V, Baslev I, Møller JM, Hansen RH, Thomsen HS. Biparametric versus multiparametric MRI in the diagnosis of prostate cancer. *Acta Radiol Open* 2016;5:20-5. [\[Crossref\]](#)
43. Kuhl CK, Bruhn R, Krämer N, Nebelung S, Heidenreich A, Schrading S. Abbreviated Biparametric Prostate MR Imaging in Men with Elevated Prostate-specific Antigen. *Radiology* 2017;285:493-505. [\[Crossref\]](#)
44. Stanzione A, Imbriaco M, Coccozza S, Fusco F, Rusconi G, Nappi C, Biparametric 3T Magnetic Resonance Imaging for prostatic cancer detection in a biopsy-naive patient population: a further improvement of PI-RADS v2? *Eur J Radiol* 2016;85:2269-74. [\[Crossref\]](#)
45. Liddell H, Jyoti R, Haxhimolla HZ. mp-MRI Prostate Characterised PIRADS 3 Lesions are Associated with a Low Risk of Clinically Significant Prostate Cancer - A Retrospective Review of 92 Biopsied PIRADS 3 Lesions. *Curr Urol* 2015;8:96-100. [\[Crossref\]](#)
46. Epstein JI, Walsh PC, Carmichael M, Brendler CB. Pathologic and clinical findings to predict tumor extent of nonpalpable (stage T1c) prostate cancer. *JAMA* 1994;271:368-74. [\[Crossref\]](#)
47. Scialpi M, Martorana E, Aisa MC, Rondoni V, D'Andrea A, Bianchi G. Score 3 prostate lesions: a gray zone for PI-RADS v2. *Turk J Urol* 2017;43:237-40. [\[Crossref\]](#)
48. Albertsen PC, Hanley JA, Fine J. 20-year outcomes following conservative management of clinically localized prostate cancer. *JAMA* 2005;293:2095-101. [\[Crossref\]](#)
49. Thoeny HC, Froehlich JM, Triantafyllou M, Huesler J, Bains LJ, Vermathen P, et al. Metastases in Normal-sized Pelvic Lymph Nodes: Detection with Diffusion-weighted MR Imaging. *Radiology* 2014;273:125-35. [\[Crossref\]](#)
50. Rosenkrantz AB, Taneja SS. Radiologist, be aware: ten pitfalls that confound the interpretation of multiparametric prostate MRI. *Am J Roentgenol* 2014;202:109-20. [\[Crossref\]](#)