Pitfalls in Prostate Cancer Magnetic Resonance Imaging

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Abstract
Image-guided prostate biopsies are changing the outlook of prostate cancer (PCa) diagnosis, with the degree of suspicion on multiparametric magnetic resonance imaging (mp-MRI) being a strong predictor of targeted biopsy outcome. It is important not only to detect these suspicious lesions but also to be aware of the potential pitfalls in mp-MRI prostate imaging. The aim of this pictorial essay is to show a wide spectrum of representative cases, which are frequently misdiagnosed as PIRADS ⅘ while reporting mp-MRI of the prostate. We provide some valuable recommendations to avoid these fallacies and improve mp-MRI of prostate evaluation.

Keywords
► Cancer
► magnetic resonance imaging
► pitfall
► prostate

Introduction
In the past few years, multiparametric magnetic resonance imaging (mp-MRI) of the prostate has undergone many technical advances, related to scientific research, in the diagnosis of prostate cancer (PCa). At present, mp-MRI prostate has become the imaging modality of choice for the detection, localization, staging and grading of PCa, and response assessment to therapy.¹,² Mp-MRI has been proved to be an imaging modality superior to any other imaging technique in evaluating prostate pathologies for several reasons such as detailed anatomical and functional information of the prostate gland. The detection of PCa on T2-weighted imaging (T2WI) can be confounded by false-positive findings such as prostatitis, calcification, postbiopsy hemorrhage, benign prostatic hyperplasia, fibrosis, radiation, and hormonal tissue changes. The mp-MRI combines anatomic details with functional information while evaluating tissues by diffusion-weighted imaging (DWI) and perfusion dynamic contrast-enhanced (DCE) sequences. The multiparameter concept refers to the fact that magnetic resonance sequences in isolation have limited accuracy in detecting and classifying prostate neoplasms; however, a combination of several sequences has demonstrated high accuracy in the detection of PCa.³ The positivity rates for the biopsy is around 70 to 90% for MRI findings highly suspicious for PCa; however, true data regarding false-negative rates of MRI-transrectal ultrasound (TRUS) fusion-targeted biopsies are lacking.⁴ Predicting final histopathological Gleason score on mp-MRI still remains a challenge, which has not been taken into account with the current prostate imaging reporting and data system (PIRADS) scoring system. However, various models are used, mainly taking into consideration clinical factors such as age, preoperative prostate-specific antigen (PSA) level, PIRADS score ≥ 4, and the number of positive biopsy cores involved with the cancer to predict the final Gleason score.⁵

The term "pitfall" means a hidden or unsuspected danger or difficulty. It is mandatory to know the pitfalls that can be encountered while reporting mp-MRI prostate. To standardize the reporting of mp-MRI prostate, the PIRADS was developed by the American College of Radiology and European
Society of Urogenital Radiology in 2012. At present, the PIRADS version 2.1 is effective since March 2019.

A sound knowledge of the pelvic anatomy and pathology, with emphasis on that of the prostate and PI-RADS v2.1 chalk out the “dominant” sequences needed to adequately evaluate the anatomical zones of the prostate, with emphasis on the utilization of diffusion/apparent diffusion coefficient (ADC) map for the peripheral zone (PZ) and T2 sequences for the central zone (CZ).3

This pictorial essay aims to give special importance to the cases of pitfalls of PCa, and suggestions are given to avoid them in daily clinical practice. Pitfalls can be divided into those conforming to the anatomical zones of the prostate gland, including the anterior fibromuscular stroma (AFMS), surgical capsule, posterior pseudolesion of the PZ (teardrop sign), periprostatic venous plexus, and neurovascular bundle, which can simulate cancer and benign conditions such as benign prostate hyperplasia and inflammatory/infectious pathology, and other conditions that can mimic malignant lesions of prostate. Various pitfalls in prostate imaging are highlighted in Table 1, and various MRI recommendations to avoid these pitfalls are tabulated in Table 2.

### Magnetic Resonance Anatomy of Prostate

Prostate gland consists of the base, the midgland, and the apex. It is divided into four histologic zones: (a) the AFMS which contains no glandular tissue; (b) the transition zone (TZ), surrounding the urethra proximal to the verumontanum, which contains 5% of the glandular tissue; (c) the CZ, surrounding the ejaculatory ducts, which contains approximately 20% of the glandular tissue; and (d) the outer PZ, containing 70 to 80% of the glandular tissue (Fig. 1).

On mp-MRI, T1-weighted (T1W) images are used primarily to determine the presence of hemorrhage within the prostate and delineate the outline of the gland. T2W images are used to interpret zonal anatomy and abnormalities within the gland and evaluate extraprostatic invasion. On T2W images, clinically significant cancers in the PZ usually appear as round or ill-defined hypointense focal lesions. DWI reflects the random motion of water molecules and is a key component of the prostate mp-MRI examination. It should include an ADC map and high b-value images.

Most clinically significant cancers have restricted/impeded diffusion compared with normal tissues and thus appear hypointense on ADC maps. DCE MRI is defined as the acquisition of rapid T1W gradient echo scans before, during, and after the intravenous (IV) administration of a low-molecular weight gadolinium-based contrast agent. As with many other malignancies following a bolus injection of contrast, PCas often demonstrate early enhancement compared with normal tissue.

### Anatomic Pitfalls

**Hypertrophic Anterior Fibromuscular Stroma**

According to the McNeal zonal anatomy distinction, AFMS is the most anterior portion of the gland, situated between the two lobes that constitute the TZ, which often appears to be flattened by them.6 This zone corresponds to a band of tissue formed by muscle cells and connective tissue, which is devoid of glandular tissue. There is variable prominence of AFMS among individuals. Due to the compressive effect of benign prostatic hyperplasia (BPH), it is less prominent in men with increased age and gland size.7 Owing to its histological composition (muscle cells and connective tissue), it can be mistaken for PCa and also tend to be hypointense (Fig. 2). In some cases where hypertrophy of AFMS may mimic PCa on T2W images, signal characteristics on DWI and enhancement on DCE-MRI should be correlated.8

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Abbreviations: BPH, benign prostatic hyperplasia; PCa, prostate cancer; mp-MRI, multiparametric magnetic resonance; MR, magnetic resonance.
mass-like configuration (►Fig. 3). In some cases, asymmetric thickening of the capsule may simulate a suspicious lesion. Multiplanar, high-resolution T2W images and DCE-MRI help us to differentiate it from the malignant lesion.9,10

### Table 2 Fallacies and recommendations to avoid pitfalls

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Abbreviations: DWI, diffusion-weighted imaging; DCE, dynamic contrast-enhanced; MR, magnetic resonance; T2WI, T2-weighted image; T1WI, T1-weighted image.

### Fig. 1 Line diagram in different planes showing normal anatomy of prostate (AFS—anterior fibromuscular stroma, TZ—transitional zone, CZ—central zone; PZ—peripheral zone; ED—ejaculatory ducts; PUT—prostatic urethra; NVB—neurovascular bundle; SV—seminal vesicles; U—urethra; V—verumontanum).

### Fig. 2 Axial T2-weighted imaging (T2WI) showing triangular area of T2-hypointensity (arrows) in midlower prostate, corresponding to hypertrophic anterior fibromuscular stroma (AFS).

### Fig. 3 Axial T2-weighted imaging (T2WI) showing T2-hypointense linear thickening (arrows) of surgical capsule, bulging into the bright peripheral zone (PZ).

### Periprostatic Venous Plexus

The location of the periprostatic venous plexus at the anterolateral margin of the prostate, near the prostate capsule, presents a challenge in assessing lesions of the PZ. Also known as Santorini’s plexus, it is in close proximity to the prostate capsule, and the size of the vessels is variable among patients.11 Patients with local inflammatory processes often show asymmetric dilation of the venous plexus. It appears rounded on axial images, with variable signal intensity on T2W images (depending on the blood flow inside the
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vessel and flow voids) and low value on ADC map (►Fig. 4). To avoid this pitfall, we must know the location of the periprostatic venous plexus, which is found in the anterolateral aspect of the PZ.

**Neurovascular Bundle**

The neurovascular bundle that innervates the corpora cavernosa is located at the posterolateral margin of the prostate, near the capsule, usually at 5 and 7 o’clock position. The preservation of this plexus is critical in postprostatectomy sexual function; however, in infiltrative PCa patients, surgeons are bounded to perform a wider resection. The neurovascular bundle may show a rounded appearance on axial images, with a low signal on T2W and ADC maps (►Fig. 5). In distorted ADC images, the neurovascular bundle may appear to be in the PZ and mimic a nodule. It is mandatory to evaluate these suspicious nodules on multiplanar, high-resolution T2W images.

**Benign Prostatic Hyperplasia**

**Moustache Sign (Bilateral Hyperplasia with Compression of Peripheral Zone)**

It is the presence of bilateral symmetrical small adenomatous nodules in the TZ, which determines compression of the peripheral and CZ at the base/middle of the prostate gland.

**Fig. 4** Axial T1-weighted (T1W) dynamic contrast-enhanced MR images (a) showing dot like enhancement in the periphery of the prostate, (b) corresponding to perfusion map images showing increased perfusion suggestive of periprostatic venous plexus.

**Fig. 5** Apparent diffusion coefficient (ADC) map showing hypointense dots in posterolateral region of prostate at 5 to 7 o’clock position, which could be erroneously diagnosed as peripheral zone (PZ) cancer, but location of these hypointense dots clinches the diagnosis of neurovascular bundle.

**Fig. 6** Axial and coronal T2-weighted image (T2WI) of the prostate base (a and b) showing bilateral symmetrical oval-shaped areas of low-signal intensity on T2WI (arrows) at the base of the prostate lateral to the opening of ejaculatory ducts. (c) No diffusion restriction is seen in these nodules. (d) Line diagram showing resemblance with a moustache.
It is also referred to as “moustache sign”, which is the presence of bilateral symmetrical homogenous oval-shaped areas of low-signal intensity on T2W images at the base of prostate, lateral to the opening of ejaculatory ducts, which appears like moustache (►Fig. 6). Presence of symmetry is a crucial feature to determine whether you are dealing with a true “moustache sign” or a clinically significant suspicious lesion in this same area, especially when evaluating on coronal T2 sequence.12

**Moustache-Like Sign (Larger Adenoma Against the Peripheral Zone)**

Protrusion of large adenomatous nodules in the PZ at the base of the gland shows moustache sign-like appearance (►Fig. 7).

**Cancer in Moustache Sign**

The key feature to differentiate a pitfall from PCa is the symmetry of the finding. Asymmetry in the signal intensity with marked T2 hypointensity together with significant early focal enhancement and high grade of diffusion restriction points toward a malignant change—cancer in moustache sign (►Fig. 8).

**Teardrop Sign (Median Posterior Compressed Central Zone)**

This corresponds to the fusion and thickening of the prostate-fascia capsule in the midline at the junction of both

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**Fig. 7** Axial T2-weighted image (T2WI) of the prostate base, (a) showing a large adenoma bulging into the peripheral zone (PZ) (arrows) and (b) corresponding line diagram showing resemblance to a moustache, (c) with restricted diffusion in the apparent diffusion coefficient (ADC) map images.

**Fig. 8** MRI prostate showing asymmetrical oval area of low signal on T2-weighted imaging (T2WI) (a) depicting early enhancement on dynamic contrast-enhanced MRI (b), and diffusion restriction (c and d) in keeping with prostate cancer.

**Fig. 9** (a) Axial T2-weighted imaging (T2WI) showing triangular area of hypointensity (arrows) in the peripheral zone (PZ) in the prostate base. (b) Coronal T2WI showing teardrop configuration (arrows) of the compressed central zone (CZ) between the transitional zone (TZ) and PZ. (c) Line diagram showing resemblance with a teardrop.
the lobes surrounding the ejaculatory ducts, also called “tear-
drop sign” because of its shape that resembles a teardrop. This
pseudolesion is found as a focal nodular hypointense area in
the midline at the level of the base-middle third of the gland,
as the CZ is compressed between the TZ and PZ (►Fig. 9).13

Teardrop-like Sign (Protruding Benign Prostatic
Hyperplasia Above the Verumontanum)
Verumontanum is an important landmark in distinguish-
ing compressed CZ and protruding BPH nodule. Teardrop
sign is due to bulging of CZ above the verumontanum, while
teadrop-like sign is due to bulging of BPH nodule above the
verumontanum.14

Ectopic/Extruded Benign Prostatic Hyperplasia Nodule
Ectopic adenomatous nodules are frequently misdiagnosed
by inexperienced radiologists, interpreting it as mp-MRI
prostate. These can mimic malignant lesions due to their
rounded contour with low signal on T2WI; in some cases,
these can show diffusion restriction with early enhancement
pattern in perfusion study (DCE). Quantitative analysis of
ADC values could improve the diagnostic accuracy in differ-
entiating ectopic BPH nodule from peripheral PCa: the higher
the ADC values, the higher the likelihood of ectopic BPH nod-
ule. The crucial step in identification of these nodules is the
demonstration of well-defined T2 hypointense capsule. Some
of the ectopic BPH nodules may show tiny T2 hyperintense
spots, corresponding to dilated acini, and this is a feature that
can be used to distinguish them from PZ PCa (►Fig. 10).

Bacterial Prostatitis (Acute and Chronic)
It comprises the most common category of benign pathologies
mimicking PCa. Acute bacterial prostatitis commonly affects
young men; however, chronic prostatitis is commonly seen in

Fig. 10 Multiparametric MRI (mp-MRI) prostate showing well-defined encapsulated oval area of T2-hypointensity with internal bright spots
within the nodule on axial and coronal T2-weighted imaging (T2WI) (a and b). Histopathology (c) showing stromal nodule with slit-like vessels,
H and E, ×2 consistent with benign prostatic hyperplasia (BPH) nodule.

Fig. 11 Multiparametric MRI (mp-MRI) prostate showing altered area of signal with fuzzy margins and bulging contour (a), patchy enhance-
ment on postcontrast images (b), and diffusion restriction (c). Histopathological images (d) depicting acute prostatitis with karyorrhectic
debris and necrosis, H and E, ×10.
older patients. Chronic prostatitis tends to be more indolent, and patients are often asymptomatic; however, acute prostatitis patients presents with fever, urinary tract symptoms, and often frequently associated with abscess.\(^\text{15}\) Although prostatitis reveals similar signal characteristics as PCa on mp-MRI, the morphology and degree of altered signal intensity with quantitative analysis of perfusion kinetics provide a clue to clinch the diagnosis. Inflammatory lesions exhibit nonmass-like morphology with ill-defined linear rather than rounded margins and diffuse lobar distribution (\(\text{Fig. } 11\)).\(^\text{16}\)

**Granulomatous Prostatitis**

Most of the cases are idiopathic; however, prior history of intravesical bacillus Calmette–Guerin (BCG) therapy of the urinary bladder, tuberculous prostatitis, and transurethral resection of the prostate is frequently encountered. On

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**Fig. 12** MRI prostate showing an area of mild, low-signal intensity on T2-weighted imaging (T2WI) (a), with corresponding hyperintense area on precontrast T1-weighted imaging (T1WI) (b). Mild restricted diffusion on diffusion-weighted imaging (DWI) and apparent diffusion coefficient (ADC) maps is seen (c and d). This is consistent with the products from the hemoglobin degradation after biopsy.

**Fig. 13** MRI prostate showing hypointense foci on T2-weighted imaging (T2WI) (a), T1-weighted imaging (T1WI) (b), without any diffusion restriction on diffusion-weighted imaging (DWI) and apparent diffusion coefficient (ADC) maps (c and d), suggestive of calcification within the prostate gland.
mp-MRI, it may appear as mild-to-markedly T2 hypointense lesion, which shows diffusion restriction and early enhancement on DCE-MRI. After antimicrobial therapy, short-term follow-up MRI may be helpful to assess therapeutic response.

Iatrogenic Changes
Biopsy needle-induced damage may determine a transient irregularity of the periprostatic capsule, leading to an overestimation of extraprostatic tumor extension.

Postbiopsy Hemorrhage
Hemorrhage is seen more frequently in the PZ than TZ. The signal intensity of postbiopsy hemorrhage depends on the state of hemoglobin degradation. In some metabolite states, it can mimic the suspicious lesion due to low signal on T2W and ADC images; however, it characteristically shows a high signal on the T1W sequence (Fig. 12). As most of the mp-MRI studies are performed before prostate biopsies, the incidence of this pitfall is low, and if it is performed later, most of the experienced radiologists’ advice to wait for at least 6 to 8 weeks postbiopsy. This delay is more advisable if it is required for staging purpose, as staging is more dependent on T2WI. If the primary aim is a diagnosis, then this delay can be avoided, as the likelihood of clinically significant PCs at the site of postbiopsy hemorrhage without a corresponding suspicious finding on MRI is low.

Posttreatment Changes
Treatment commonly includes radical prostatectomy, prostate-sparing focal therapy, or radiation therapy (RT), which can include external-beam RT or brachytherapy. There are some common mimics that should be identified in posttreatment cases. Granulation tissue and hemorrhage both will show early enhancement on DCE phase; however, on DWI, they will appear benign with no notable signal abnormalities. Posttreatment appears more hypointense than a recurrent tumor with the absence of early enhancement and presence of delayed enhancement during the venous phase. If metallic clips are used during surgery, the DCE sequence is more reliable than DWI. Early enhancement in DCE is highly suggestive of recurrence. All these changes should be correlated with clinical setting, type of treatment received, and PSA levels.

Artifacts

Mispositioned Endorectal Coil
Use of endorectal is mandatory for the good-quality MRI scanning of prostate on 1.5T scanner. Accurate positioning of a coil is essential; otherwise, it creates artifacts affecting image quality, thus making the diagnosis difficult. Sometimes, it involves adjacent PZ and mimics PCs.

Calcifications
Incidence of prostate calcification is high and found basically due to concreted secretions or corpora amylacea. They are generally detected in the TZ (central gland). Most of the time, they are indolent. They are characteristically hypointense in all sequences like T1, T2, and ADC, with blooming on susceptibility-weighted imaging (SWI), and do not show diffusion restriction (Fig. 13). CT is helpful to demonstrate the calcification.

Foley’s Catheter Tip
Foley’s catheter tip is hypointense on all sequences (Fig. 14). Coronal images will show its entire urethral course and its extension into the urinary bladder.

Periprostatic Lymph Nodes
Periprostatic lymph nodes are encountered rarely while interpreting mp-MRI prostate. These are commonly located...
at the base of the prostate gland laterally or posterolaterally. On mp-MRI imaging, these show well-defined rounded morphology with a hypointense signal on T2W sequence and marked diffusion restriction; however, the degree of diffusion restriction does not suggest metastatic involvement. Sometimes on low-resolution DWI/ADC images, periprostatic lymph nodes appeared to be intraprostatic in location. Detailed evaluation on high-resolution, multiplanar T2W images reveals its periprostatic location.11,23

**Conclusion**

All oncologists and radiologists involved in treating and interpreting mp-MRI prostate should be aware of these fallacies described above. Key points to avoid these pitfalls are illustrated in Table 3. Despite the pitfalls described above, biopsy should be considered in men with an equivocal MRI and a PSA density of > 0.15.

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**Conflicts of Interest**

There are no conflicts of interest.

**References**