



Prostate MR: pitfalls and benign lesions

Aritrick Chatterjee¹ · Stephen Thomas¹ · Aytekin Oto¹

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Abstract

Multiparametric MRI (mpMRI) of the prostate has evolved to be an integral component for the diagnosis, risk stratification, staging, and targeting of prostate cancer. However, anatomic and histologic mimics of prostate cancer on mpMRI exist. Anatomic feature that mimic prostate cancer on mpMRI include anterior fibromuscular stroma, normal central zone, periprostatic venous plexus, and thickened surgical capsule (transition zone pseudocapsule). Benign conditions such as post-biopsy hemorrhage, prostatitis or inflammation, focal prostate atrophy, benign prostatic hyperplasia nodules, and prostatic calcifications can also mimic prostate cancer on mpMRI. Technical challenges and other pitfalls such as image distortion, motion artifacts, and endorectal coil placements can also limit the efficacy of mpMRI. Knowledge of prostate anatomy, location of the lesion and its imaging features on different sequences, and being familiar with the common pitfalls are critical for the radiologists who interpret mpMRI. Therefore, this article reviews the pitfalls (anatomic structures and technical challenges) and benign lesions or abnormalities that may mimic prostate cancer on mpMRI and how to interpret them.

Keywords mpMRI · Prostate cancer · DWI · Pitfalls · Benign lesions

Introduction

Multiparametric MRI (mpMRI) of the prostate has evolved to be an integral component for the diagnosis, risk stratification, and staging of prostate cancer (PCa). Despite, the increasing use of mpMRI for prostate cancer diagnosis, about 15–30% of all clinically significant cancers can be missed by radiologists [1, 2]. Moreover, there is large inter-observer variability in the interpretation of mpMRI among radiologists [3]. In addition, there are several pitfalls which can mimic prostate cancer or hinder its diagnosis on mpMRI. Normal anatomical structures and benign processes may have similar imaging features to PCa and lead to false positive results. Some of these entities can be recognized by their imaging features, while others cannot be confidently distinguished from PCa. Knowledge of prostate anatomy,

location of the lesion and its imaging features on different sequences, and being familiar with the common pitfalls are critical for the radiologists who interpret mpMRI. In this article, we will review the pitfalls (anatomic structures and technical challenges) and benign lesions or abnormalities and that may mimic PCa on mpMRI.

Anatomic structures mimicking PCa

Anterior fibromuscular stroma

The anterior fibromuscular stroma (AFMS) is a band of fibromuscular tissue that forms the anterior surface of the prostate, and is located anterior to the transition zone. It is composed of connective tissue, smooth muscle, and some skeletal muscle. It extends from the apex to the base of the prostate. It is typically thicker at midline. The prostate capsule blends with the AFMS and in the majority of cases (89%), the AFMS forms the only anterior covering of the prostate [4]. Laterally, the AFMS fuses with the lateral pelvic fascia and covers outermost regions of the lateral and anterior surfaces of the prostate in the majority of cases (85%) [5]. AFMS has low signal on T2WI with low ADC values and therefore, can obscure an anterior tumor or mimic PCa when asymmetric (Fig. 1). Normal AFMS has a benign

✉ Aytekin Oto
aoto@radiology.bsd.uchicago.edu

Aritrick Chatterjee
aritrack@uchicago.edu

Stephen Thomas
stthomas@radiology.bsd.uchicago.edu

¹ Department of Radiology, University of Chicago, 5841 South Maryland Avenue, Chicago, IL 60637, USA

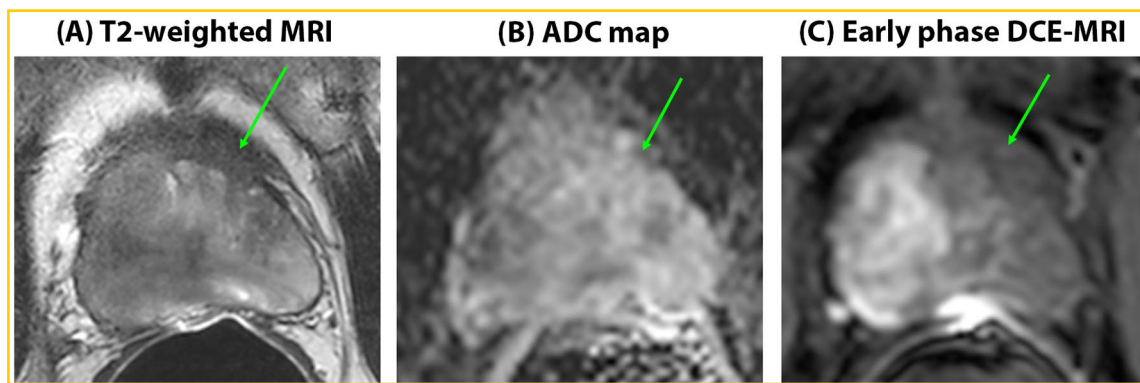


Fig. 1 57-year-old man with thickened AFMS that mimics prostate cancer. Axial T2WI (a) shows thickened AFMS (arrows) with low signal in the left side mimicking cancer (arrow), while the AFMS on the contralateral side appears normal. The left AFMS shows diffused low signal on ADC map (b), but shows no early enhancement

on DCE-MRI (c). The thickened AFMS was found to be benign on prostatectomy. The patient had multifocal, diffuse prostate cancer in rest of the prostate. The mpMRI images were taken on a 3T MRI system with an endorectal coil

progressive (type 1) curve at DCE [6]. Symmetric appearance and DCE-MRI characteristics are helpful in differentiating AFMS from anterior cancer.

Anterior cancers account for about 20% of all PCa and tend to have higher rates of extracapsular extension [7]. Most anterior tumors are located in the anterior third and inferior half of the TZ or in the AFMS [8]. In contrast to AFMS, anterior tumors are asymmetric and have a washout (type-3) enhancement curve with earlier enhancement than normal AFMS [6]. The consultation of T2W images in coronal and sagittal planes can confirm the continuity of the hypertrophic anterior fibromuscular stroma with the benign tissue [9].

Normal central zone

The currently utilized prostate zonal anatomy is based on the classical work done by McNeal, who proposed different prostate ‘zones’ based on anatomic location and histologic

and embryologic features [10]. Anatomically, the seminal vesicles are located posterosuperiorly and drain into the mid prostatic urethra via the ejaculatory ducts in the region of the verumontanum. The central zone surrounds the ejaculatory ducts and is located posterior to the transition zone and the urethra proximal to the verumontanum [11]. Typically, the central zone (CZ) is symmetric, is homogenous low signal on T2WI, has low uniform ADC and does not show focal early enhancement on DCE-MRI as seen in Fig. 2. However, the ADC of benign CZ is lower than the ADC of other zones of the prostate and has overlap even with significant PCa [12]. On DCE-MRI, the normal CZ has a progressive (type 1) or plateau (type 2) enhancement curve [13]. DCE-MRI and symmetric appearances at expected location are the key features differentiating normal CZ from cancer.

With age, the CZ undergoes distortion with hypertrophy of the transition zone (TZ) from prostatic hyperplasia, the CZ is compressed and displaced towards the base of the

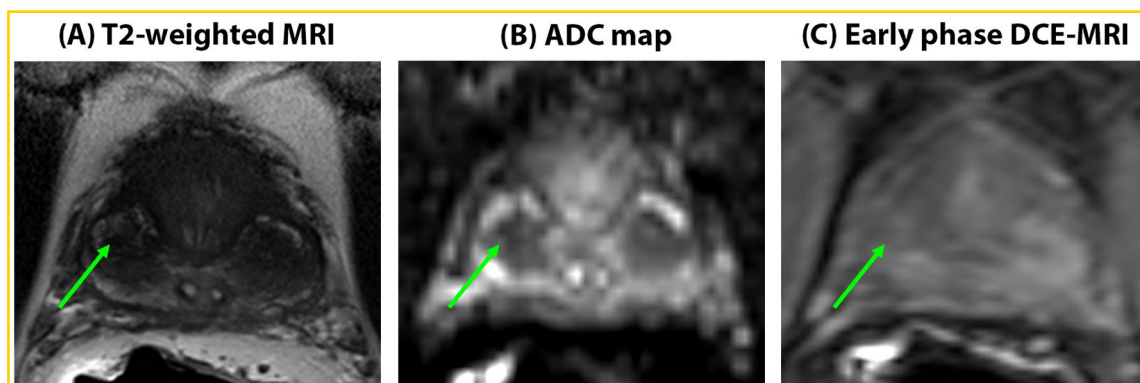


Fig. 2 68-year-old man with normal central zone. The right central zone (arrows) appears as a region of hypointensity on T2WI (a), restricted diffusion seen as low signal on ADC map (b). However, no

early focal enhancement on DCE-MRI (c) in central zone as seen here is indicative of benign tissue. The mpMRI images were taken on a 3T MRI system with an endorectal coil

prostate gland [14]. While central zone cancers account only for 0.5–2.5% of all PCa, they tend to have a higher Gleason grade, and more likely to invade the seminal vesicles, and have extracapsular extension. They are also prone to biochemical recurrence after prostatectomy [15]. Uneven hypertrophy of the TZ can cause asymmetric displacement of the CZ which can be seen as an area of focal signal abnormality near the peripheral zone (PZ) and present as a pseudolesion (asymmetric CZ). The asymmetric CZ will have a benign enhancement curve on DCE-MRI.

Another CZ related pitfall is the presence of focal hypointense region near the ejaculatory glands at the base of the prostate. This is related to the posterior bulging of the central zone over the verumontanum. This pitfall can show mildly restricted diffusion and early focal enhancement on DCE-MRI. The use of coronal and sagittal planes can be useful to demonstrate continuity and symmetry of the CZ.

Periprostatic venous plexus

The prostate is drained by the periprostatic venous plexus that lies along the posterolateral aspect of the gland. This plexus then communicates with the dorsal venous plexus (Santorini) and hemorrhoidal venous plexus posteriorly to drain into the internal iliac vein [16]. The diameter and number of the veins varies by age and is more pronounced in younger patients and patients with smaller glands [17].

The periprostatic veins are tubular structures and can be hypointense on T2W and ADC depending on blood flow velocity and turbulence (Fig. 3). Veins close to the peripheral zone may be mistaken for a peripheral zone PCa particularly in areas where the prostatic pseudocapsule is indistinct and partial volume averaging can hinder discerning the prostate boundary [18]. The periprostatic draining veins mostly enhance avidly on DCE-MRI (venous plexus

with low ADC). Following the veins as tubular structures on multiple planes and recognizing, the prostate boundary will help the reader identify the plexus. Layering signal intensities within the thrombosed or slow flowing vessels can be seen on DWI/ADC and T2W images. This feature can help to identify a vein of the periprostatic plexus. A hyperintense rim around the periprostatic venous plexus can be seen on T2W images (especially coronal view) [19].

Thick surgical capsule (transition zone pseudocapsule)

The transition zone and the peripheral zone both have similar embryological origin. The surgical capsule of the prostate separates the transition zone from the peripheral zone [20]. It is composed of fibromuscular stroma and compressed glandular tissue. This band forms a landmark for surgical enucleation or ablation of transitional zone benign prostate hyperplastic nodules [21]. Poor definition of the surgical capsule on T2-weighted images can sometimes be seen adjacent to transition zone cancers [22]. This band of tissue can also hypertrophy with age and enlargement of the transition zone [23].

The surgical capsule is hypointense on T2W with low ADC due to dense compressed fibromuscular tissue and does not demonstrate early focal enhancement (Fig. 4 shows thick pseudocapsule). Its symmetric crescentic band like shape and location should help identify this anatomic landmark.

Benign lesions mimicking PCa

Post-biopsy hemorrhage

Blood products from biopsy accumulate along the needle track and can be larger than the expected needle track. The prostate normally produces citrate which is an anti-coagulant

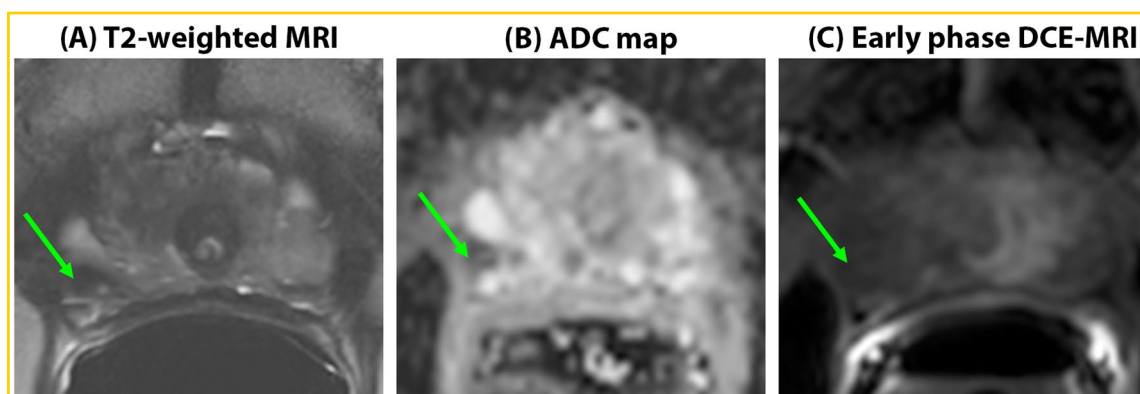


Fig. 3 57-year-old man with periprostatic vein in the right posterolateral side mimicking prostate cancer. The periprostatic vein in the right posterolateral side (arrows) in the peripheral zone appears as a region of hypointensity on T2WI (a), with restricted diffusion

(hypointensity) seen on ADC map (b) mimicking prostate cancer. However, periprostatic vein does not show early focal enhancement on DCE-MRI (c) in this case. The mpMRI images were taken on a 3T MRI system with an endorectal coil

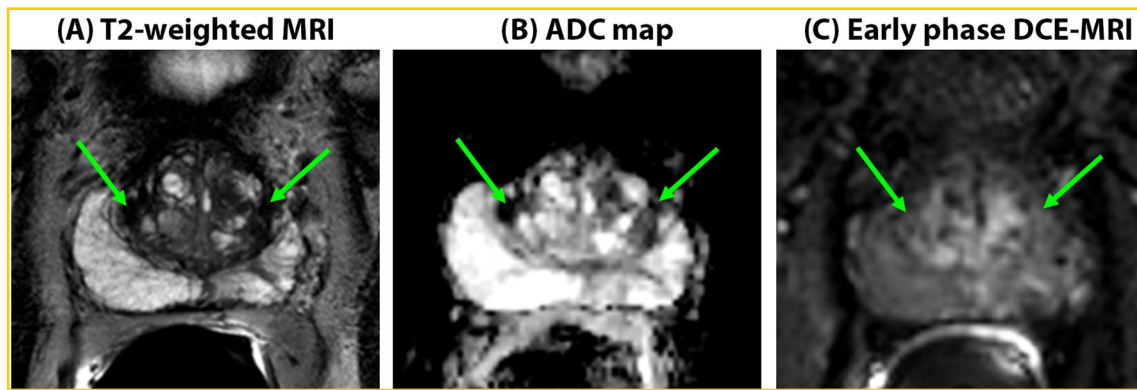


Fig. 4 71-year-old man with thickened surgical capsule mimicking prostate cancer. The surgical capsule (arrows) appears as a region of hypointensity on T2WI (a), with focal diffusion restriction (hypointensity) seen on ADC map (b) mimicking prostate cancer, but shows no early focal enhancement on DCE-MRI (c). The mpMRI images were taken on a 3T MRI system with an endorectal coil

tensity) seen on ADC map (b) mimicking prostate cancer, but shows no early focal enhancement on DCE-MRI (c). The mpMRI images were taken on a 3T MRI system with an endorectal coil

that causes protracted bleeding and after biopsy, therefore causing T2 signal intensity decrease. Similar to prostate cancer, hemorrhage is hypointense on T2W and show restricted diffusion (hypointense on ADC) on mpMRI (Fig. 5), and therefore can mimic or obscure focal PCa [24, 25]. However, a matching area of T1 hyperintensity is key to differentiating it from prostate cancer. Hemorrhage can demonstrate focal, diffuse, wedge shaped, or striated high signal on T1W. The intrinsic high T1 of hemorrhage also limits DCE evaluation without subtraction images.

Post-biopsy hemorrhage decreases with time with hemorrhagic changes seen in 72.2% patients at 4 weeks, 57.1% at 4–6 weeks and 52% after 6 weeks [26]. Post-biopsy hemorrhage can be detected up to 4 months after biopsy. A delay of 6–8 weeks after biopsy has been suggested to decrease the confounding imaging findings of prostate hemorrhage [27]. Hemorrhage can obscure underlying PCa [28].

The “T1 hemorrhage exclusion sign” is defined as a well-defined region of hypointensity on T1-weighted images that is completely surrounded by an area of high signal intensity.

The presence of the T1 hemorrhage exclusion sign with a corresponding area of low signal intensity on T2-weighted images has been shown to be strongly predictive of prostate cancer (95% positive predictive value) [29]. This can be seen in Fig. 5.

Prostatitis/inflammation

Acute and chronic prostatitis can cause signal changes in the PZ that may be indistinguishable from PCa. Acute bacterial infection is most commonly caused by *Escherichia coli* and accounts for up to 10% of all prostatitis diagnoses. Acute prostatitis has a bimodal age distribution affecting men aged 20 to 40 years of age and men older than 70 years [30, 31]. Acute prostatitis results in an influx of neutrophils into the prostate.

Chronic bacterial prostatitis is thought to result from undertreated acute prostatitis and may be caused by infection moving from the distal urethra to the prostate and with

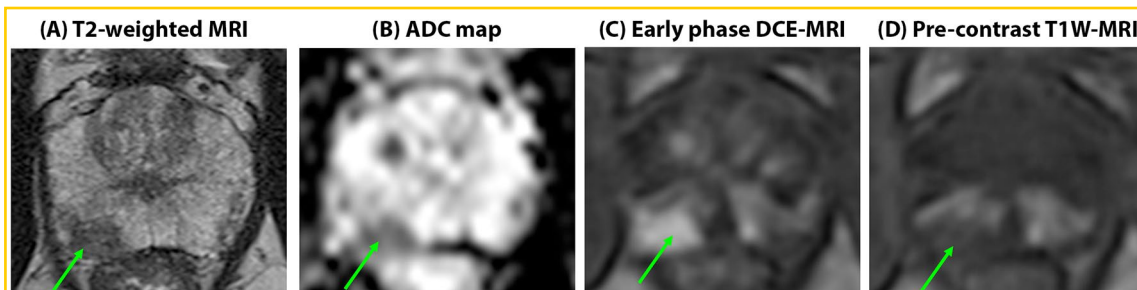


Fig. 5 61-year-old man with hemorrhage and biopsy confirmed prostate cancer undergoes prostate MRI for staging. Axial T2WI (a) shows a focal low signal area in the right peripheral zone (arrow). Axial ADC map (b) shows corresponding low ADC (arrow). Post-contrast T1 (c) show brisk enhancement of the lesion, biopsy confirmed PCa, Axial pre-contrast T1 (d) shows the area ‘spared’ (arrow) with surrounding hemorrhage, also known as T1 hemorrhage exclusion sign. The mpMRI images were taken on a 3T MRI system without an endorectal coil, using pelvic phased array coil

firmed PCa, Axial pre-contrast T1 (d) shows the area ‘spared’ (arrow) with surrounding hemorrhage, also known as T1 hemorrhage exclusion sign. The mpMRI images were taken on a 3T MRI system without an endorectal coil, using pelvic phased array coil

a lifetime prevalence of 1.8% to 8.2% [32]. The predominant cell type is a lymphocytic infiltrate.

Prostatitis can be focal or diffuse involving both the PZ and TZ. On MRI, the affected areas are hypointense on T2W and ADC with mild to moderate diffusion restriction due to the increased inflammatory cellular infiltrate (Fig. 6). The mean ADC of PCa was found to be significantly lower compared with non-cancerous tissue with prostatitis [33]. Similarly, wedge-shaped areas of prostatitis are mildly hypointense on ADC maps [25]. The clinical history of the patient plays a crucial role in distinguishing focal prostatitis from the presence of PCa.

Granulomatous prostatitis (GP) is an inflammatory condition characterized by the histologic presence of epithelioid granulomas with or without other inflammatory cells and is secondary to prior mycobacterial or fungal infection, prostate biopsy, or systemic granulomatous infection [34, 35]. The diagnosis of GP is on the rise due to increased transurethral resection of the prostate (TURP), needle biopsy procedures, and extensive use of intravesical Bacillus Calmette–Guerin (BCG) instillation for the treatment of non-muscle invasive bladder cancer (NMIBC) [35, 36]. Clinically, granulomatous prostatitis may present as a focal or diffuse area of induration, often giving a stony hard feel on rectal exam with normal to raised serum PSA levels and/or hematuria. It may co-exist with areas of PCa and diagnosis is based on needle biopsy and pathological analysis [35]. Granulomatous inflammation without a causative agent termed non-specific granulomatous prostatitis accounts for 75% of cases. On MRI, focal granulomatous prostatitis has low signal on T2W and low ADC making it difficult to discriminate from PCa with the lesions having lower mean ADC values than higher-grade PCa and are typically assigned higher PI-RADS scores of 4/5 [34, 37].

Focal prostate atrophy

Focal prostate atrophy (FPA) is a common histologic diagnosis found in 73% of biopsy specimens, and occurs most commonly in the PZ. The most common causes of atrophy are inflammation, androgen therapy, and irradiation. The histologic types are proliferative atrophy (PA) and proliferative inflammatory atrophy (PIA) being the most common type [38–40]. The theory of chronic inflammation from PIA leading to accumulated genetic mutations and cellular proliferation causing PCa has not been established. The histologic diagnosis of PIA in negative biopsies correlates with a decreased frequency of detecting PCa in men with persistent suspicion of PCa. There is a positive and significant association between the extent of atrophy and the total or free serum PSA elevation possibly due to damaged epithelial cells in atrophic acini [41]. The imaging features of FPA overlap with PCa on histology and mpMRI [39].

On MRI, FPA is seen as a focal area of volume loss usually with wedge-shaped appearance. It is hypointense on T2W, with moderate diffusion restriction and moderate early enhancement on DCE-MR images. Figure 7 shows a focal atrophy that appears as wedge-shaped lesion on T2W. About 26% of peripheral wedge-shaped lesions are associated with non-specific atrophy [25]. However, the diffusion restriction and tissue enhancement is usually less marked than PCa [22, 25]. Asymmetric volume loss due to atrophy can be a way to differentiate focal atrophy from PCa [22].

Benign prostatic hyperplasia nodules in the Transition zone

Benign prostatic hyperplasia (BPH) is a histologic diagnosis of cystic, stromal, and glandular epithelial hyperplasia that occurs in the transition zone (TZ) with a prevalence of 50 to 60% by age 60 years [42]. There are four

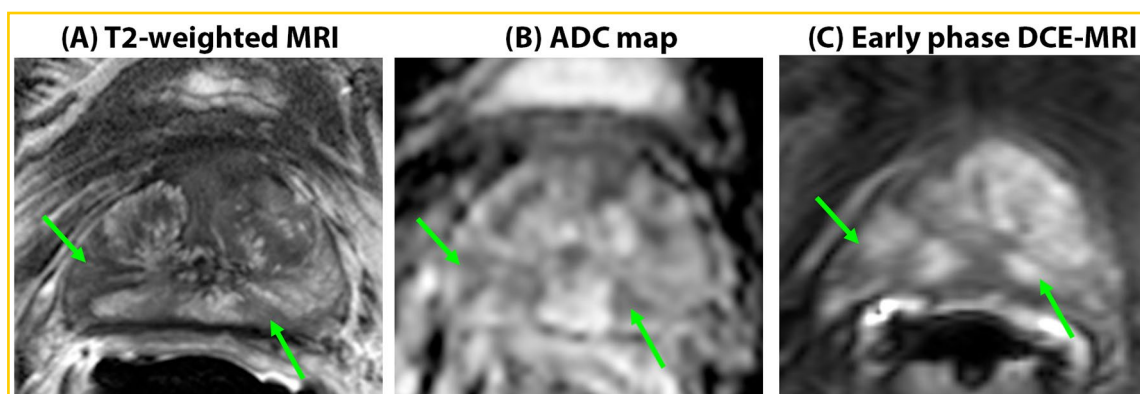


Fig. 6 49-year-old man with prostatitis mimicking prostate cancer. Prostatitis (arrows) appears as regions of hypointensity in the left peripheral zone on T2WI (a), with restricted diffusion (hypointensity) seen on ADC map (b) and early focal enhancement on DCE-MRI (c)

mimicking prostate cancer. Chronic inflammation was found on prostatectomy. The mpMRI images were taken on a 3T MRI system with an endorectal coil

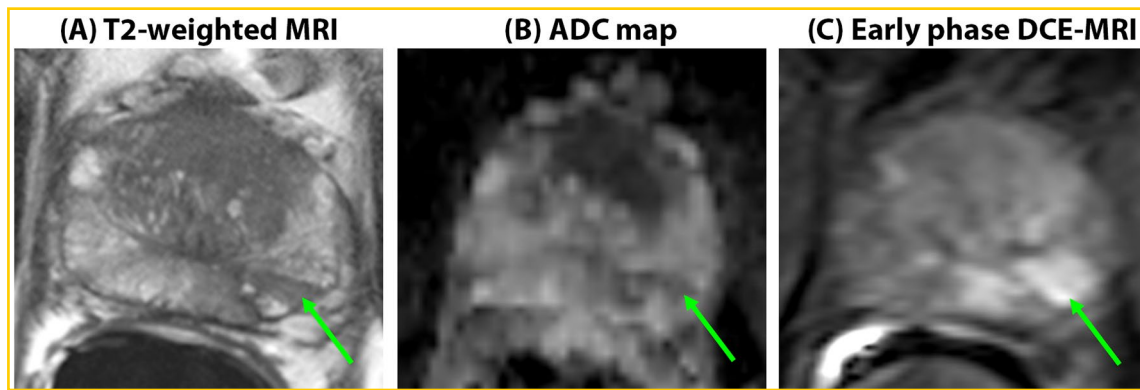


Fig. 7 59-year-old man with focal prostate atrophy mimicking prostate cancer. The focal prostate atrophy (arrows) appears as a wedge-shaped lesion in the left peripheral zone with hypointensity on T2WI (a), with mildly restricted diffusion (hypointensity) seen on ADC map (b) and early focal enhancement on DCE-MRI (c) mimicking

prostate cancer. On prostatectomy, the suspected lesion was found to have focal atrophy along with intraluminal blood and hemosiderin-laden macrophages. The mpMRI images were taken on a 3T MRI system with an endorectal coil

different types of BPH nodules: cystic, stromal, glandular, and mixed [43]. Stromal nodules are hypointense on T2-weighted images due to proliferation of fibromuscular components. They have low ADC signal and demonstrate early enhancement with washout on DCE-MR images mimicking PCa (Fig. 8) [43, 44]. Glandular BPH are not hyperintense on ADC and T2 as previously thought [44], and have similar appearance on mpMRI as stromal BPH [43]. Cystic BPH are well demarcated and, have higher fluid content due to cystic ectasia and hyperplastic glandular components. They tend to be hyperintense on T2 and ADC images and do not show any early enhancement of DCE-MRI, making it easy to distinguish from PCa. A recent study [43] showed that quantitative ADC values could be used to differentiate between PCa and BPH, as ADC for PCa was lower compared to all BPH types. T2

values were found to be significantly lower in PCa compared to cystic BPH only, while glandular and stromal BPH showed no significant difference from PCa. DCE-MRI was not found to be useful for distinguishing PCa from BPH.

About one quarter of all prostate cancers occur in the TZ, making it important to differentiate between benign BPH nodules and TZ cancers [45]. Transition zone cancers are larger, and have higher Gleason grade compared to peripheral zone lesions [46]. Morphology on T2WI can be used to differentiate PCa from BPH nodules. BPH nodules are rounded and better demarcated than PCa. Transition zone PCa have irregular margins, are lenticular shaped, and may show invasion of the AFMS. They have lower ADC values than BPH nodules, although the difference is less profound with the stromal nodules [14, 43, 44, 47].

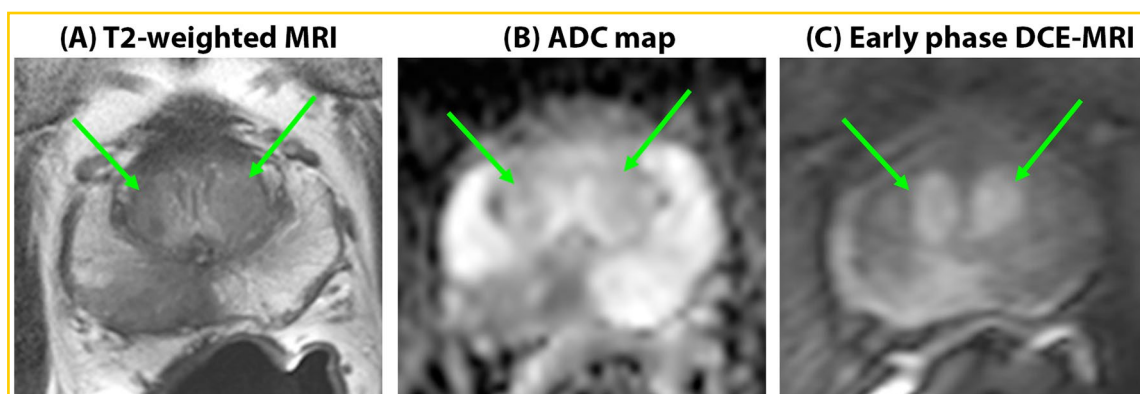


Fig. 8 53-year-old man with BPH mimicking prostate cancer. The stromal BPH (arrows) appears as a region of hypointensity on T2WI (a), with restricted diffusion (hypointensity) seen on ADC map (b), and shows early focal enhancement on DCE-MRI (c), mimicking

prostate cancer. Other clinical finding included a PI-RADS 5 lesion in the right peripheral zone that was found to be Gleason 4+3 cancer on prostatectomy. The mpMRI images were taken on a 3T MRI system with an endorectal coil

BPH nodules in the peripheral zone

Benign prostatic nodules can be found in the peripheral zone (PZ) of the prostate and tend to be glandular in etiology [48, 49]. Histologic tissue analysis suggests some of these nodules originate from the PZ and are not the extension or herniation of TZ nodules [50]. BPH nodules located in the PZ (Fig. 9) are usually well circumscribed, ovoid or round, and do not extend to the capsule [48]. They have low T2 and ADC signal with avid early enhancement and washout very similar to PZ cancers. Their well-defined borders, location (adjacent to TZ), and round shape are the important imaging findings in their differentiation from PZ cancer. In some cases, stromal tumors of uncertain malignant potential (STUMP) can appear like nodular region like BPH and extend into the PZ mimicking cancer [51].

Prostatic calcifications

Prostatic calcifications are relatively common radiologic findings on CT and increases with age over 40 years. They can be associated with disease processes such as BPH, infection, and metabolic abnormalities [52]. Corpora amylacea are presumably precursors to calcified “stones”. Analysis of the proteins in prostatic corpora amylacea and calculi found that the predominant proteins comprising these concretions are proteins involved in acute inflammation, and in particular proteins contained in neutrophil granules [53]. Prostate calcifications in the prostate are commonly seen at the junction between the transition zone and peripheral zone against the background of BPH. They are typically hypointense on T2W and ADC, with low signal intensity on DWI at all b values due to the dielectric effect of calcium in the calcification and do not show early enhance on DCE-MRI (Fig. 10). Complete lack of enhancement on MRI is important in their differentiation from PCa. Correlation with CT can be helpful in confirming focal calcification. While corpora amylacea

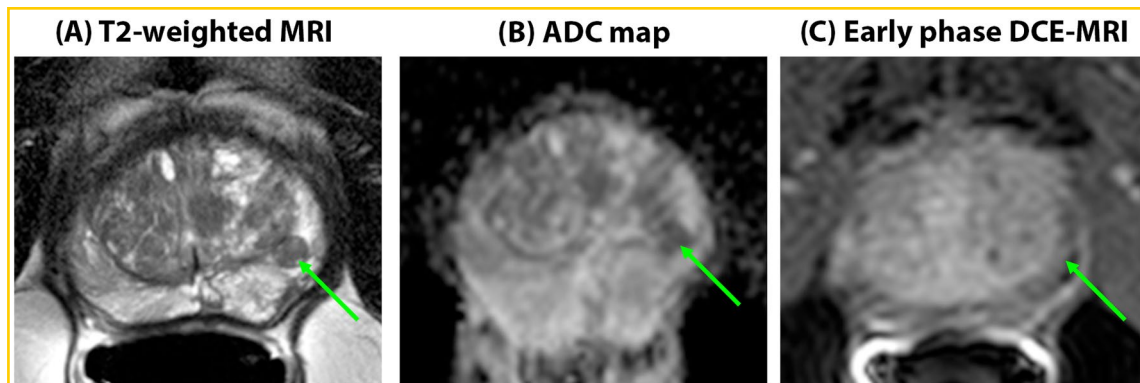


Fig. 9 53-year-old man with ectopic BPH in the peripheral zone mimicking prostate cancer. The BPH nodule in the left mid prostate (arrows) appears as a well-circumscribed round region of hypointensity on T2WI (a), with highly restricted diffusion (hypointensity) seen

on ADC map (b) mimicking prostate cancer. However, it does not show early focal enhancement on DCE-MRI (c) like other peripheral zone cancer. The mpMRI images were taken on a 3T MRI system with an endorectal coil

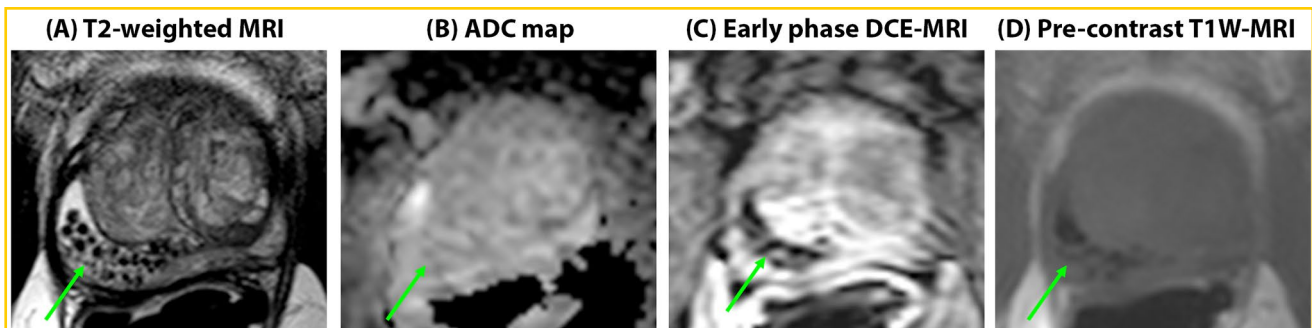


Fig. 10 84-year-old man with prostatic calcification mimicking prostate cancer. Prostatic calcification in the right peripheral zone (arrows) appears as a foci of low signal on axial T2WI, and shows some punctate foci of low signal that is subtle on ADC map (b).

However, post-contrast DCE-MRI (c) shows no enhancement (arrow). Pre-contrast T1W image (d) shows hypointense foci in the right peripheral zone. The mpMRI images were taken on a 3T MRI system with an endorectal coil

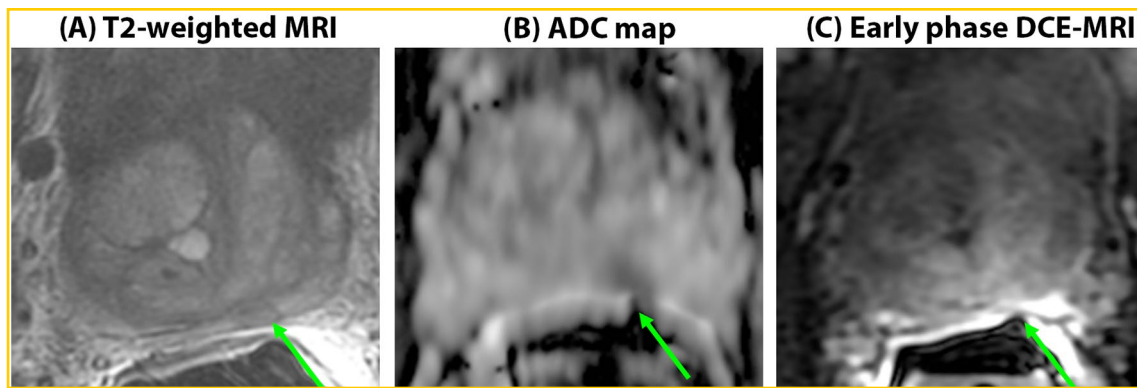


Fig. 11 65-year-old man undergoing prostate MRI found to have an endorectal coil artifact mimicking prostate cancer. Incompletely assessed mildly T2-hypointense focus within the left peripheral zone, in the area of artifact related to the endorectal coil seen on T2W

image (a), with corresponding area with restricted diffusion (hypointensity) seen on ADC map (b) and artifact on DCE-MRI (c) mimicking prostate cancer. The mpMRI images were taken on a 3T MRI system with an endorectal coil

can occur within foci of prostatic adenocarcinoma, their incidence is low and are more associated with low Gleason grade cancers [54].

Other pitfalls

Technical challenges can also limit the efficacy of mpMRI. Image distortion, motion artifacts, and endorectal coil placements can affect the PCa diagnosis results using mpMRI.

DWI especially is susceptible to image distortion. DWI uses echo-planar imaging, where gradients are switched to generate diffusion weighting (different b values). This changing magnetic field induces eddy currents, which in turn can create change in the main magnetic field, time-varying gradients, and also degrade the speed and efficiency of gradient switching. Therefore, this leads to a wide range of image artifacts, including shearing, shading, scaling, blurring, and spatial misregistration. High b value DWI is highly susceptible to anatomic distortions and warping, which can lead to obscurity of lesion and false positives. Figure 10 in addition to calcification also shows spatial misregistration between ADC and other mpMRI sequences due to artifacts from DWI echo-planar sequence.

The presence of stool and rectal gas can produce anatomic distortion/artifacts due to magnetic susceptibility. The use of enema to void the rectum of stool can be effective [55]. The use of left–right phase encoding instead of anterior–posterior can mitigate distortion due to stool and rectal gas. The presence of rectal gas and movement can affect ADC calculations and DCE-MRI images.

The use of an endorectal coil can induce artifacts. The antenna on the endorectal coil is recommended to be placed at 2 and 10 o'clock position. However, improper coil placement can lead to the artifacts from the antenna which can mimic cancer. Figure 11 shows artifact from an endorectal coil that mimics PCa.

Conclusion

While mpMRI is a robust application for the diagnosis, staging and targeting of prostate cancer, anatomic, and histologic mimics of prostate cancer exist. Some of these pitfalls can be recognized by evaluating all sequences (summarized in Table 1), while some can only be confirmed with histopathologic evaluation. It is important for the radiologists to recognize these pitfalls while interpreting prostate MRI.

Table 1 Differentiating prostate cancer from it mimics on mpMRI

<i>Anatomic structures mimicking prostate cancer</i>	
Anterior fibromuscular stroma	Symmetric appearance and DCE-MRI characteristics (benign progressive type 1 curve) are helpful in differentiating AFMS from anterior cancer. Consultation of T2W images in coronal and sagittal planes can confirm the continuity of the hypertrophic anterior fibromuscular stroma with the benign tissue
Normal central zone	DCE-MRI (progressive type 1 or plateau type 2 enhancement curve) and symmetric appearances at expected location are the key features differentiating normal CZ from cancer. The use of coronal and sagittal planes can be useful to demonstrate continuity and symmetry of the CZ
Periprostatic venous plexus	Periprostatic venous plexus enhance avidly on DCE-MRI. Following the veins as tubular structures on multiple planes and recognition of the prostate boundary will help the reader identify the plexus. Layering signal intensities within the thrombosed or slow flowing vessels can be seen on DWI/ADC and T2W images
Thick surgical capsule	Does not demonstrate early focal enhancement. Its symmetric crescentic band like shape and location should help identify this anatomic landmark
<i>Benign lesions mimicking PCa</i>	
Post-biopsy hemorrhage	T1 hyperintensity is key to differentiating hemorrhage from prostate cancer
Prostatitis/inflammation	ADC of prostate cancer is lower compared with non-cancerous tissue with prostatitis. The clinical history of the patient plays a crucial role in distinguishing focal prostatitis from the presence of PCa
Focal prostate atrophy	The diffusion restriction and tissue contrast enhancement on DCE-MRI in atrophy is usually less marked than prostate cancer. Asymmetric volume loss due to atrophy can be a way to differentiate focal atrophy from PCa
Benign prostatic hyperplasia nodules in the Transition zone	Morphology on T2W images can be used to differentiate prostate cancer from BPH nodules. BPH nodules are rounded and better demarcated than PCa. BPH nodules tend to have higher ADC values compared to cancer, although the difference is less profound with stromal nodules
BPH nodules in the peripheral zone	Their well-defined borders, location (adjacent to TZ and do not extend to the capsule) and shape (usually well circumscribed, ovoid or round) are the important imaging findings in their differentiation from PZ cancer
Prostatic calcifications	Complete lack of enhancement on MRI is important in their differentiation from PCa

Compliance with ethical standards

Conflict of interest Dr. Aritr Chatterjee and Dr. Stephen Thomas have no disclosures. Dr. Aytekin Oto has the following disclosures. Research Grant, Koninklijke Philips NV Research Grant, Guerbet SA Research Grant, Profound Medical Inc. Medical Advisory Board, Profound Medical Inc Speaker, Bracco Group.

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