Computer-Aided Detection of Prostate Cancer with MRI:
Technology and Applications

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Abstract

One in six men will develop prostate cancer in his life time. Early detection and accurate diagnosis of the disease can improve cancer survival and reduce treatment costs. Recently, imaging of prostate cancer has greatly advanced since the introduction of multi-parametric magnetic resonance imaging (mp-MRI). Mp-MRI consists of T2-weighted sequences combined with functional sequences including dynamic contrast-enhanced MRI, diffusion-weighted MRI, and MR spectroscopy imaging. Due to the big data and variations in imaging sequences, detection can be affected by multiple factors such as observer variability and visibility and complexity of the lesions. In order to improve quantitative assessment of the disease, various computer-aided detection systems have been designed to help radiologists in their clinical practice. This review paper presents an overview of literatures on computer-aided detection of prostate cancer with mp-MRI, which include the technology and its applications. The aim of the survey is threefold: an introduction for those new to the field, an overview for those working in the field, and a reference for those searching for literature on a specific application.

Keywords

Prostate cancer; MR imaging; Image Quantification; Computer-aided Detection

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INTRODUCTION

Prostate cancer (PCa) is currently the most common cancer in men and the second leading cause of cancer-related deaths among men in the United States [1]. In 2015, it is estimated that the number of estimated new cases and deaths will be 220,800 and 27,540, respectively, accounting for 26.0% of new cancer cases and 8.8% of cancer deaths for American men [1].

The prostate is subdivided into the base, mid-gland, and apex from superior to inferior. The prostate also has four anatomic zones: the transition zone (TZ), which contains 5% of the glandular tissue and accounts for around 25% of PCa; the central zone (CZ), which contains 20% of the glandular tissue and accounts for around 5% of PCa; the peripheral zone (PZ), which contains 70–80% of the glandular tissue and accounts for about 70% of PCa; and the non-glandular anterior fibromuscular stroma. Accurate localization of PCa within the TZ or the PZ is extremely important as TZ prostate cancer is associated with favorable pathologic features and better recurrence-free survival [2].

At present, the clinical standard for definitive diagnosis of prostate cancer is transrectal ultrasound (TRUS)-guided sextant or systematic biopsy. The PSA blood test and DRE results are considered to identify patients who need biopsy. The actual impact of MRI for prostate cancer management is through guided biopsies and improved cancer diagnosis and staging yield. In recent years, magnetic resonance imaging (MRI) targeted prostate biopsies have been showing better disease localization and more accurate sampling than conventional TRUS-guided biopsy in various studies [3–6]. MRI-based computer-assisted sophisticated imaging for individual patients would offer such a significant role in defining an optimal targeted biopsy and interventional approach. Several approaches have been explored to improve the accuracy of image-guided targeted prostate biopsy, including in-bore MRI-guided, cognitive fusion, and MRI/transrectal ultrasound fusion-guided biopsy [7].

MR imaging provides excellent soft-tissue contrast and has become an imaging modality of choice for localization of prostate tumors. Multiparametric MRI (mp-MRI) includes high-resolution T2-weighted (T2W) MRI, diffusion-weighted imaging (DWI), dynamic contrast-enhanced imaging (DCE-MR), and MR spectroscopy (MRS). The mp-MRI has proven to be an effective technique to localize high-risk prostate cancer [8, 9]. The combined use of anatomic and functional information provided by the multiparametric approach increases the accuracy of MR imaging in detecting and staging prostate cancer [8, 9]. It can also help guide biopsies to achieve a higher tumor detection rate and better reflect the true Gleason grade. The European Society of Urogenital Radiology (ESUR) in 2012 established the Prostate Imaging-Reporting and Data System (PI-RADS) scoring system for multiparametric MRI of the prostate [10]. The MR PI-RADS aims to enable consistent interpretation, communication and reporting of prostate mp-MRI findings [10, 11]. A joint steering committee formed by the American College of Radiology, ESUR, and the AdMeTech Foundation have recently announced an updated version of the proposals of PI-RADS Version 2 [12]. Prostate mp-MRI at 3 T had been recommended in PI-RADS Version 2. Generally, CAD systems are classified into two categories: computer-aided detection (CADe) and computer-aided diagnosis (CADx) systems. Currently, most CAD systems in prostate MRI focus on local suspicious lesions and discrimination between benign and
malignant lesions; most of them are computer-aided diagnosis systems. As the combination of various MR images creates large amounts of data, supportive techniques or tools, such as computer-aided diagnosis (CADx), are needed in order to make a clinical decision in a fast, effective, and reliable way.

In the past 10 years, computer-aided techniques have developed rapidly. Automated computer-aided detection and diagnosis may help improve diagnostic accuracy of PCa, and reduce interpretation variation between and within observers [13, 14]. Prostate cancer diagnosis requires an experienced radiologist to read prostate MRI, and such expertise is not widely available. Addition of CADx may significantly improve the performance of less-experienced observers in prostate cancer diagnosis. When less-experienced observers used CADx, they reached similar performance as experienced observers [13]. In a more recent study, the use of CAD can also improve prostate mp-MRI study interpretation in experienced readers [15]. For cases in which radiologists are less confident, they can get higher performance by using the computer output. A recent study showed a pattern recognition system enables radiologists to have a lower variability in diagnosis, decreases false negative rates, and reduces the time to recognize and delineate structures in the prostate [16]. The benefit of CADx also includes guiding biopsy using cancer location information from MRI [14]. Therefore, along with rapid development of MR technique, CADx of prostate cancer has become an active field of research in the last five years.

This paper starts with the review of MR image acquisition technology and then focuses on a comprehensive review of the state-of-the-art image quantification methods. The part on validation and clinical applications is a reference of the literatures available in the clinical management of the disease. The paper closes with a discussion and future perspectives.

A PubMed electronic database search for the terms "computer-aided," "CAD," "prostate," and "MRI" was completed for articles about CAD of prostate cancer up to September 11, 2015.

**MR IMAGE ACQUISITIONS**

Contemporary MR imaging of the prostate combines anatomic images from high-resolution T1W and T2W sequences and functional information obtained from DWI, DCEI, and MRS. The combination of conventional anatomical and functional MRI is known as multiparametric MRI. The PI-RADS Prostate MR Guidelines published in 2012 suggest the use of T2W images plus 2 functional techniques [10]. The anatomy of the prostate gland is visualized with T2W images; DWI and MRS add specificity to lesion characterization, while DCE-MRI has a high sensitivity in cancer detection. In the PI-RADS ™ v2, the essential components of the mp-MRI prostate examination are T2W, DWI, and DCE [12]. For the PZ, DWI is the primary determining sequence. For the TZ, T2W is the primary determining sequence. In order to obtain high and stable accuracy, a combination of anatomical and functional imaging is necessary in clinical practice. Recent studies showed an increasing interest in developing CADx systems to detect and characterize prostate cancer on the basis of an mp-MRI imaging approach [14, 15, 17, 18]. T2W MR images are frequently used in
mp-MRI CADx systems. T2W plus DWI and DCE-MRI are also commonly used among the combinations.

**T2WI and T2 mapping**

The anatomy of the prostate gland is best visualized with T2W images. The acquisition of high-resolution T2W images of the prostate is the first and most important step in an mp-MR imaging protocol. In T2W images, the peripheral zone of the prostate has hyperintense signal, whereas the central and transition zones have low signal, allowing the zonal anatomy of the prostate to be clearly delineated (Figure 1). In T2W images (Figure 2), PCa in the peripheral zone is usually depicted as a low-signal area. However, the growth pattern and the aggressiveness of the tumor can alter its appearance. T2W MR imaging has been advocated as an accurate technique in the detection of PCa in the transition zone [19, 20]. The value of T2W MR images is also in predicting pathological stage and extra-capsular extension of PCa [21].

Because T2W MR images play an important role in both location and staging of PCa, T2W MRI is the basis and important sequence in CADx systems for PCa. In T2W MR images, the tumor region of interest (ROI) has more dark pixels than bright pixels, whereas the normal tissue ROI has more bright pixels than dark pixels. Different features, including fractal features, textural features and signal intensity can be used by CADx. Because prostate cancers at the central gland and peripheral zone usually have significantly different texture on T2W MR images [22], and because the use of mp-MRI may have challenges for detecting cancer at the transition zone [23], a CADx system that can analyze features based on the lesion’s location may be able to aid in the detection of suspicious lesions.

T2 maps offer quantitative T2 values. As the standard T2 mapping approach of performing multiple single spin-echo acquisitions with a range of TE settings requires excessive scan times, the T2 mapping is not include in most clinical applications. Recently, some new sequences can provide an effective approach to speed up T2 quantification [24, 25]. T2 values of histologically proven malignant tumor areas were significantly lower than the suspicious lesions but nonmalignant lesions or normal areas [26]. The use of quantitative T2 measurement improves the specificity and/or sensitivity of prostate cancer detection [27] and aggressiveness assessment [28, 29]. There is a potential benefit of incorporating quantitative T2 values into CADx systems.

**Dynamic contrast-enhanced MRI (DCE-MRI)**

DCE-MRI, which enables visualization of vascular permeability and perfusion, is an important tool in oncology to define tumor. DCE-MRI is sensitive to alterations in vascular permeability, extracellular space, and blood flow. The clinical application of DCE-MRI for prostate cancer is based on data showing that malignant lesions show earlier and faster enhancement and earlier contrast agent washout compared with healthy prostate tissues (Figure 3) [30].

The DCE-MRI data can be analyzed with various semiquantitative or quantitative models to extract parameters related to vascular permeability, extracellular space, blood flow, and water exchange [31]. Semi-quantitative DCE-MRI analysis: As semi-quantitative DCE-MRI
data are only relative to the patient, the baseline intensity is highly variable depending on the patient and the MRI protocol. It is necessary to use indicators relating to signal amplitude. The most commonly used quantitative approach of analyzing DCE-MRI is two-compartment pharmacokinetic (PK) models that can be used to generate pharmacokinetic parameters such as Ktrans (transfer of gadolinium contrast from the vasculature to the tumor, representing forward vascular perfusion and permeability) and Kep (reverse transfer of contrast agent from the extracellular space back to the plasma, representing backward leakage) in order to quantify tumor enhancement and the contrast uptake and wash-out [32]. However, pharmacokinetic model implementation typically involves assuming some prior knowledge; and the arterial input function (AIF) estimation methodology can have significant effects on the parameters estimated by PK modeling [33]. The empirical approach based on phenomenological universalities (PUN) is able to reproduce experimental data from a DCE-MRI acquisition [34] [35].

Different CADx systems have been developed to analyze the DCE-MRI data. Vos et al. developed a CADx system capable of discriminating PCa from non-malignant disorders in the peripheral zone and achieved a diagnostic accuracy of 0.83 (0.75–0.92) [36]. They also developed an automated segmentation per patient calibration method to improve the diagnostic accuracy of CADx [37]. Puech et al. designed a prostate CADx software to provide a 5-level cancer suspicion score for suspicious foci detected in DCE-MRI and T1-weighted images [38, 39].

DCE-MRI usually has lower spatial resolution than other sequences, especially when DCE-MRI is performed rapidly in a short period of time. Limitations in the interpretation of DCE-MRI data include overlap in enhancement properties between benign and malignant regions in the transition zone. Benign prostatic hyperplasia and other benign inflammatory conditions within the transition zone also exhibit substantial hypervascularity [40]. Diagnostic models containing contrast enhancement parameters have reduced performance when applied across zones, so zone-specific models can improve classification of prostate cancer on multi-parametric MRI [41].

**Diffusion-weighted MR imaging (DWI)**

The diffusion properties of tissue are related to the amount of interstitial free water and permeability. In general, cancer tends to have more restricted diffusion than normal tissue, because of the higher cell densities and abundance of intra- and intercellular membranes in cancer [42]. Diffusion weighted MRI images can be used to detect prostate cancer from differences in the diffusion of water molecules of the normal and tumor tissues (Figure 4) [42]. The diffusion-weighted image is usually generated with different b values which can be used to calculate the apparent diffusion coefficient (ADC), and the ADC for each pixel of the image is displayed as ADC map. Diffusion of water molecules in tumor tissue is thought to reflect tissue architecture such as cell density and nucleus/cytoplasm ratio, and reductions in ADC values. For these reasons, ADC values have received the attention as a predictor of Gleason score in prostate cancer [43, 44]. Studies show that DWI findings may indicate tumor aggressiveness [27, 45, 46].
Technologic advances enable performance of diffusion-weighted imaging (DWI) at high b- or ultrahigh b-values (greater than 1,000 s/mm$^2$). High b-value images can be obtained in one of two ways: either directly by acquiring a high b-value DWI sequence, or by calculating (synthesizing) the high b-value image by extrapolation from the acquired lower b-value data. Previous research has shown that high b-value DWI images allow for increased delineation between tumors and healthy tissue which makes the prostate cancer detection more robust [47, 48]. Whereas contrast in ADC maps does not significantly change with different b values, contrast ratios of DWI images are significantly higher at b-values of 1500 and 2000 s/mm$^2$ in comparison to b values of 800 and 1000 s/mm$^2$ [49]. Wang et al. have reported that DWI images and ADC maps using b = 1500 s/mm$^2$ should be considered more effective than those at b = 2000 s/mm$^2$ or b = 1000 s/mm$^2$ for detecting prostate cancer at 3 T MRI [50].

DWI images and ADC maps are the key component of the prostate mp-MRI exam. Several CADx systems adopting DWI images or ADC maps have been developed. DWI was mostly often combined with T2W in these CADx systems. Peng et al. demonstrated that the combination of 10th percentile ADC, average ADC, and T2-weighted skewness with CADx is promising in the differentiation of prostate cancer from normal tissue [27]. Niaf et al. presented a CADx system based on T2W, DWI and DCE for assisting cancer identification in the PZ [18]. Stember et al. develop a software system that identifies suspicious regions at the prostate transition zone (TZ) using signal and textural features on T2W and ADC maps, free of user input [51]. Kwak et al. recently designed a prostate CADx combined T2W and high b-value (b = 2000 s/mm$^2$) DWI. They obtained an AUC of 0.89 [52].

**MR Spectroscopy (MRS)**

In MRS, the position of each metabolite peak in the output graph reflects the resonant frequencies or chemical shifts of its hydrogen protons, and the area of each peak reflects the relative concentration of that metabolite [53]. The dominant peaks observed in prostate MRS are from protons in citrate (2.60 ppm), creatine (3.04 ppm), and choline compounds (3.20 ppm) (Figure 5) [53].

As a metabolic biomarker for PCa, MRS has not gained wide acceptance in routine clinical practice due to a variety of factors including the length and complexity of data acquisition, zonal anatomy, processing, and analysis. Visual interpretation of the spectra by a trained spectroscopist is time-consuming and requires accurate knowledge of prostate anatomy. Therefore, a method for automated analysis of prostate MRS data is necessary.

Over the last decade, with a view to assisting radiologists in interpretation and analysis of MRS data, several researchers have begun to develop CADx schemes for PCa identification from spectroscopy. Tiwari et al. developed an approach that integrated a manifold learning scheme (spectral clustering) with an unsupervised hierarchical clustering algorithm to identify spectra corresponding to cancer on prostate MRS [54]. The scheme successfully identified MRS cancer voxels with a sensitivity of 77.8%, a false positive rate of 28.92%, and a false negative rate of 20.88% [54]. They also presented a CADx scheme that integrated nonlinear dimensionality reduction (NLDR) with an unsupervised hierarchical clustering algorithm to automatically identify suspicious regions on the prostate using MRS [55]. They
introduced the use of wavelet embedding to map MRS and T2-W texture features into a common space to identify the voxels that are affected by prostate cancer [56]. They recently presented a computerized decision support system called Semi Supervised Multi Kernel Graph Embedding (SeSMiK-GE) that may be developed into a powerful diagnostic and prognostic tool for distinguishing high and low grade PCa \textit{in vivo}. Matulewicz et al. used an artificial neural network (ANN) model to automatically detect cancerous voxels from prostate MRS datasets and found that the additional information concerning the prostate’s zonal anatomy can improve the performance of the detection [57].

**Other imaging methods**

Although T2W, DWI, DCE-MRI and MRS are more commonly used in mp-MRI, some MRI methods, including diffusion tensor imaging (DTI), diffusion kurtosis imaging (DKI), and MR elastography (MRE), have been investigated for characterizing prostate cancer [58–61]. Other MRI methods, including proton density-weighted (PD-W) image [62] and T1 map[14], had also been added for feature calculation purposes in some CADx system.

DTI has been widely used in clinical applications, especially in neuro- and musculoskeletal imaging. Fractional anisotropy (FA) and apparent diffusion coefficient (ADC) values provided from DTI data reflect the degree of water diffusion restriction in different tissue. Pathological processes may cause change in normative FA values and disruption of fibers in tractography. The feasibility of performing DTI of the prostate had been demonstrated by some studies; and DTI tractography can successfully visualize fiber tracts around the prostate [58]. DTI tractography might be applicable to the estimation of structures of the prostate [59], the characterization of prostate cancer [60], and monitoring prostatic structural changes under radiotherapy [61].

The novel technique, diffusion kurtosis imaging, enables characterization of non-Gaussian water diffusion behavior. DK model may add value in PCa detection and diagnosis, and DKI potentially offers a new metric for assessment of PCa[63]. A recent study demonstrated no significant benefit of DKI for detection and grading of PCa as compared with standard ADC in the peripheral zone determined from b values of 0 and 800 s/mm [64]. The mechanical properties of the tissue of interest are calculated from the wave fields and displayed as an image, commonly referred as an elastogram. In MR elastography (MRE), an external mechanical excitation is applied to the tissue of interest to induce tissue vibrations [65]. MRE has been shown to be of clinical value in MRI for its ability to detect tissue abnormalities in organs such as the liver [66], brain [67] and breast [68–70]. More recently, researchers have also focused on the development of MRE methods to detect prostate cancer [71–73]. The resulting wave fields are imaged using a motion-sensitized MRI pulse sequence. Elastograms may add another dimension to current mp-MRI techniques for diagnosing prostate cancer, and may further increase the sensitivity and specificity of such techniques.

T1 maps offer quantitative T1 values and can be produced by a variety of methods, such as multiple inversion or multiple repetition time acquisitions, typically requiring lengthy acquisition times. Another approach taken in the context of the prostate has been to employ spoiled gradient-echo (SPGR) sequences where it is possible to obtain T1 estimates in...
relatively short acquisition times by varying the RF flip angle [74]. The T1 mapping is not including in most CADx systems. Vos, et al. [14] had presented a fully automatic CADx by combining a histogram analysis on mpMR images including T1, pharmacokinetic, T2 and ADC maps.

MR lymphography has been used for the investigation of the lymphatic channels and lymph glands. Different imaging techniques, including nanoparticle-enhanced [75, 76] and non-contrast MR lymphography [77, 78], had been developed for detection of nodal metastases. MR lymphography is a noninvasive technique that is well suited for the examination of regional (intrapelvic) lymph node metastases in PCa.

### MR IMAGE QUANTIFICATION METHODS

#### General framework

Development of computer-aided detection systems includes several aspects: image preprocessing, algorithm development, methodology for assessing CADx performance, validation using appropriate cases to measure performance and robustness, observer performance studies, assessing performance with a clinical trial, and ultimately commercialization. The development must confront several challenges. Computerized image procedure may cover different aspects of segmentation, registration, feature extraction, and classifiers. A computer algorithm should be developed based on the understanding of image reading by radiologists, such as how radiologists detect certain lesions, why they may miss some abnormalities, and how they can distinguish between benign and malignant lesions. It is important to develop CADx systems that extracts quantitative data in a more accurate and automated fashion.

Many different types of CADx systems are produced to locate/diagnose prostate cancer in MR imaging, including T2W, DWI, DCE-MRI and MRS. Considering the particularity of prostate cancer in anatomy, pathology and clinic, the core of a CADx system for the detection of prostate cancer is associated with its computerized algorithms. In general, the pipeline of the CADx system for prostate cancer is visualized schematically in Figure 6. In the initial stage, lesion candidates are selected within a likelihood map that is generated by a voxel classification of one or more images. Hereafter, the lesion candidates are segmented into a region of interest from which region-based features are extracted. Finally, the extracted information is fused by a classifier into malignancy likelihood. The following sections describe each step in detail.

#### Preprocessing

The purpose of preprocessing is to normalize the MR data or to transform the MR data to a domain in which prostate lesions can be easily detected.

For T2W-MRI, the image intensities can vary, even using the same protocol and the same scanner. The quality of images depends on the acquisition conditions such as temperature, calibration adjustment, B0 intensity, coil position, and the receiver gain value. In addition, the intensity variation will increase when different scanners are used. This relationship must be taken into account for MR image analysis. Collewet et al. [79] proposed four schemes for
the intensity normalization. The most used method is that intensities are proportionally normalized by defining the median+2*(inter-quartile range).

ADC maps calculated from DWI are useful for detecting prostate cancer with a relatively high specificity. However, it has lower resolution than T2W-MRI and is subject to magnetic susceptibility artifacts [17]. ADC represents a quantitative assessment of water diffusion. Lower ADC value is associated with higher rate of malignancy. Prostate cancer can be identified as a low signal region on ADC maps against a background of normal tissue with higher signal intensity [17].

Intensity inhomogeneity arises from the imperfections of the image acquisition, which can reduce the accuracy of segmentation, classification, and registration. The most intuitive method to correct intensity inhomogeneity is image smoothing or homomorphic filtering [80]. Vovk et al. [81] classify inhomogeneity correction methods into two categories, which are prospective and retrospective. Prospective methods aim at the calibration and improvement of image acquisition processes. Retrospective methods rely exclusively on the information of the acquired images or on a priori knowledge. Sled et al. [82] proposed a nonparametric non-uniform intensity normalization (N3) method for inhomogeneity correction, which is independent of pulse sequence. Tustison et al. [83] proposed a variant of N3 for bias field correction. Similar to the N3, the source code, testing, and technical documentation are publically available and the package is "N4ITK". This algorithm is available to the public through the Insight Toolkit of the National Institutes of Health.

Segmentation

The segmentation aims to reduce the burden of the classifier in the later stages. Therefore, the classifiers only focus on the prostate region obtained by segmentation methods. T2W imaging provides the best resolution and contrast to show the anatomy of the prostate and has a very high sensitivity for prostate cancer. Therefore, T2W-MRI is the most useful image sequence in determining the contours of prostate.

Extensive studies were developed to segment the prostate from MR images [62, 84–90]. It can be a challenging task to obtain accurate prostate volume in T2W-MRI. Firstly, the contrast between the prostate and the surrounding tissues can be low. Therefore, it may be difficult to accurately segment the boundary of the prostate. Secondly, the prostate shapes of different patients can be significantly different. Even for the same patient, the prostate motion at different patient positions can be large, which results in a shape difference on MR images. Thirdly, MR image appearance, quality, and the presence of artifacts can be affected by different scanners, which in turn can have a large influence on the performance of computerized algorithms. All these aspects need to be considered when developing a robust and accurate segmentation method for prostate MR images.

Contour and shape based methods [91–95]exploit edge and shape features to segment the prostate, which contains two categories. The first category is edge based segmentation methods. The edge detection operators are used to produce edges on MR images. The candidate edges are picked up and then connected to obtain the prostate boundary. Zwiggelaar et al [91] developed a semi-automatic method to segment the prostate in MRI
data. Their method exploits the characteristics of the anatomical shape of the prostate when represented in a polar transform space. The edge detection and non-maximum suppression are used to track the boundary of the prostate.

The second category is deformable model based segmentation methods. Kass et al [96] proposed an active contour model and used the image gradient to evolve a curve. The internal spline force pushes the curve toward the salient image feature, while external force is responsible for putting the curve near the object. Chan and Vese [97] proposed a level set algorithm of the piecewise constant variant of the Mumford-Shah model [98] for segmentation.

Atlas based methods are also used to segment the prostate in MR images [99]. An atlas consists of original image data and its corresponding manual segmentation. The atlas can be used as a reference to segment the prostate of a new patient. Klein et al. [99] proposed an automatic method for segmenting the prostate in 3D MR images. Their method is based on non-rigid registration of a set of pre-labeled atlas images. The label images of the deformed atlas are fused to yield a segmentation of images from a new patient.

Besides the above methods, a global optimization algorithm called graph cut [100, 101] is becoming more and more popular due to its efficient global minimization. The segmentation problem can be formulated as a minimization of an energy minimization. Egger [102] proposed a graph-based approach to automatically segment prostate based on a spherical template. The minimal cost on the graph is optimized by a graph cut algorithm, which can get the segmentation of the prostate volume. Mahapatra and Buhmann [85] proposed a fully automatic method for prostate segmentation using random forests classifiers and graph cuts. The prostate probability map was generated based on a random forests classifier. The negative log-likelihood of the probability maps was used as the penalty cost in an energy function, which was minimized by graph cuts. Tian et al. [103] proposed a supervoxel-based segmentation method for the prostate. The prostate segmentation problem was considered as assigning labels to supervoxels. An energy function with both data and smoothness terms was used to model the labels, which was minimized using graph cuts. The segmentation results are shown in Figure 7. Other segmentation methods were also developed for the prostate [104, 105]. Ghose et al. [105] reviewed segmentation methods for the prostate in TRUS, MR and CT images. They studied the similarities and differences among the different methods, highlighted their advantages and disadvantages in order to assist in the choice of an appropriate segmentation methods. They also showed a comprehensive description of the existing methods in all TRUS, MR and CT images, and highlighted their key-points and features. They provided a strategy for choosing segmentation method for a given image modality.

A publicly available data set called MICCAI Challenge Prostate MR Image Segmentation (PROMISE12) [104] can be used to evaluate the performances of the new proposed methods. This data set contains 50 cases with ground truths for training, and 30 cases without ground truths for test, which are 3D T2w MR transverse images of the prostate. The MR images were obtained from multicenter, multivendor, and different acquisition protocols (i.e., with/without endorectal coil, differences in slice thickness).
Registration

Image registration is a process of aligning two or more images, which aims to find the optimal transformation that best aligns the structures of interest in the input images. Image registration is needed in order to integrate the features from different images of mp-MRI such as DCE-MRI and T2W MRI. The registration of images requires the selection of the feature space, a similarity measure, a transformation type, and a search strategy [106]. The DICOM header of MR images can provide coordination and orientation information that are useful for registering T2W, ADC, and $K^{\text{trans}}$ maps. T2W-MRI is considered as the reference. Other modalities can be registered to T2W-MRI by aligning the coordinates of their origins, which are obtained from the DICOM header. If necessary, resolution adjustment is also performed after the alignment.

Registration is also used to validate in vivo MR imaging using ex vivo histologic images [107, 108]. To obtain the reliable ground truth of the prostate cancer region, whole-mount histology is performed on ex vivo prostate. The pathologist labels the cancer region in the histology images. Based on the registration between the whole-mount histology and T2W MRI, the labelling of the cancer in histology can be mapped to T2W MRI for validation [107, 108]. Kalavagunta et al. [108] proposed a method to register MRI and histology using local affine transformations guided by internal structures. First, the histologic and MR images are first segmented, scaled, and translated. Second, the prostate capsule and internal structure masks are identified to constrain the pathology transformation. A transformation matrix is obtained by registering two images based on capsule and internal structure masks. Third, the pathology images are warped using a computed transformation matrix. Fourth, a transformation matrix is applied for each annotated cancer region. The warped cancer regions are superposed on registered pathology images. Last, the cancer regions in MRI can be obtained by mapping the cancer regions of pathologic images to MR images. In another study, Chappelow et al. [107] presented a new registration method that maximizes the combined mutual information shared by the intensity of the reference image and multiple representations of the floating images in multiple feature spaces. The method provides enhanced registration performance by combining the intensity information with transformed feature images from the images. These feature images are not as susceptible to intensity artifacts and provide additional similarity information regarding the reference image but not contained in the floating image. This method is particularly useful for registering MRI and histology.

Feature extraction

Feature extraction plays an important role in prostate MRI CADx systems. Classic features for medical images include intensity, shape, texture, and statistical features. For medical image classification, choosing the right features for a classifier is more important than choosing the classifier itself [62].

Litjens et al. classified the features into five types: intensity, pharmacokinetic, texture, blobness, and anatomical features [62]. For the intensity feature, a T2-estimate map is generated by using the MR signal equation, the proton density image and a reference tissue [88]. Anatomical features include the relative distance to the prostate boundary and the
relative position feature. Both the relative distance and relative position features are calculated with respect to the prostate surface obtained by segmentation methods. For the pharmacokinetic feature, the traditional analysis is incorporated in their CADx system by using a curve fitting technique to fit a bi-exponential curve to the time data, as presented in [109]. For the texture feature, a Gaussian texture bank was used to capture the textural distortions [22]. For the blobness feature, it was found that prostate cancer tends to appear as a blob-like lesion in DWI and DCE-MRI. The blobness-filter presented by Li et al. was chosen as a blobness measure [110]. Blobness is calculated on the ADC, tau and LateWash images, and on the $K^{\text{trans}}$ and $K^{\text{ep}}$ images as well [110].

Shah et al. [17] created an mp-MRI feature set for CADx systems (Figure 8). First, in order to reduce interpatient variability, normalized T2W maps were calculated from the transversal T2W intensities using the average fat signal adjacent to the prostate as a reference. Second, quantitative ADC maps were computed from the transversal DWI by fitting the dependence of the signal intensity in each pixel. Third, each dynamic curve was de-noised by using a wavelet filter for DCE-MRI. The pharmacokinetic parameters were extracted by using the generalized kinetic model (GKM) [111, 112]. Then, the GKM was fitted to the measured concentration time curves, using the linear least-squared method [112] to yield the volume transfer constant $K^{\text{trans}}$, and the rate constant $k^{\text{ep}}$. Finally, the normalized T2W and ADC maps were resized to have a pixel resolution equal to the T1 and $K^{\text{trans}}$ and $k^{\text{ep}}$ maps in order to form the final feature set for the CADx system.

Niaf et al. extracted about 140 kinds of features for a CADx system [18]. Most of these features were chosen based on their proven efficiency between cancer and non-cancer. Two categories of features were proposed: image features and functional features. For image features, there were three types: grey-level features, texture features, and gradient features. The image intensity values of T2, DCE and ADC maps were used as grey-level features. First order texture measurements were computed for each pixel over a local window, which includes mean, median, standard deviation and average deviation. Second order texture features were computed based on two neighboring pixels, which includes co-occurrence matrix. The Sobel and Kirsch filters and numerical gradient operators were used to compute gradient features (Figure 9).

Radiomics is an emerging field for the quantification of tumor phenotypes by applying a large number of quantitative image features [113, 114]. Radiomics can provide complementary and interchangeable information to improve individualized treatment selection and monitoring. Since medical imaging technology is routinely used in clinical practice worldwide, radiomics may have a high clinical impact on future patient management. The workflow of radiomics consists of three steps [113]. The first step is the acquisition of standardized images for diagnostic or planning purposes. On the images, the tumor regions are extracted by an algorithm or an experienced radiologist. Second, quantitative imaging features are extracted from the tumor regions. These features involve tumor image intensity, texture, and shape and size of the tumor. Last, all the extracted features are analyzed and selected by a model. The most informative features are identified and incorporated into predictive models for treatment outcome. Radiomics, as a high dimensional mineable feature space, can be used for prostate cancer. Cameron et al. had
constructs a comprehensive radiomics feature model to detect tumorous regions using mp-MRI [115]. New radiomics-driven texture feature models had been developed for the detection of prostate cancer and for the classification of prostate cancer Gleason scores by utilizing mp-MRI data [116–118].

**Classification**

Image classification involves training and testing with features extracted from image data and its corresponding labels [62]. A classifier is usually trained by using the labeled image data set and applied to unseen image data sets. Several classification techniques from the machine learning field have been developed for picking up discriminative features. Support vector machines (SVMs) and random forests could achieve good performance based on the positive and negative training samples [17, 119]. A pixel classification provides a likelihood between 0 and 1 for each pixel, with 0 indicating no suspicion of prostate cancer and 1 indicating high suspicion of cancer.

Litjens et al. [62] experimented with three different classifiers: a linear discriminate classifier, a Gentle Boost classifier [120], and a random forests classifier [119] with regression trees. Shah et al. used SVM to create a classifier model [17]. Because real data are not linearly separable, the SVM implementation was used to allow relaxed constraint for misclassified points. SVMs “kernel trick” was also implemented to enable operations to be performed in the input space rather than the potentially high-dimensional feature space [121].

Chan et al. [122] investigated the use of a statistical classifier for detecting prostate cancer by combining information from MR images. SVM is used to predict the tumor likelihood in the peripheral zone using the derived features. For SVM training, they randomly sampled 10% of the PZ data and retained all the tumor data to confine the training dataset to a manageable size for SVM training convergence. The radial basis function kernel was used for SVM. These works indicate that the SVM classifiers and random forests work well on the problem of classifying prostate tumors on mp-MRI.

**VALIDATION**

When developing a CADx system for prostate MRI, the accuracy of the “gold standard” is important. Histopathology, as the ground truth, usually includes findings from prostatectomy specimens or biopsy specimens. The validation of CADx systems is summarized in Table 2. In order to transfer the labels from pathology to MR images, MR images usually need to be registered with pathological sections of the prostate. An accurate registration of histologic and MR images serves as the bridge between *in vivo* anatomical information and *ex vivo* pathologic information, which is valuable in developing a CADx system.

Whole-mount sections are generated from tissue slices and microscopic slices are stained with hematoxylin-eosin staining after being embedded in paraffin [111, 123]. Pathologists outline each lesion on the microscopic slices. Gleason scores of different regions may also be provided on the microscopic slices. For correlation between MR images and histopathologic images, the corresponding anatomical landmarks and cancerous regions are
manually labeled by an expert. The urethra may serve as a guide for correlating the images. In order to improve the accuracy and efficiency of the correlation, some automatic methods have been developed [111, 124].

There are several challenges in establishing automatic correlation between in vivo MR images and histopathologic images. The orientation of the specimen and its sections may be different from that of in vivo MR imaging. There are mismatches between MR imaging and histopathology, which make it difficult to assess the true accuracy of MRI. Once the anatomic orientation in the body is lost, it may be difficult to section the prostate in the same plane as that of in vivo MR images. The specimen can be marked with separate colors on the left, right, and anterior aspects for anatomic orientation [111]. Using image processing, computer-aided design, and rapid prototyping technology, a customized mold has been used to process prostatectomy specimens for each patient [124]. The customized mold holds the prostate in the same position and the same shape as those of in vivo MR images and guides the cutting knife to obtain tissue blocks that correspond to the image slices.

The prostate is an easily deformable organ, hence, the gland deforms during and after prostatectomy. Additionally, prostate MRI is often performed by using an endorectal coil, which further deforms the gland. Specimen formalin fixation and paraffin embedding also induce variable tissue shrinkage. Deformable image registration provides a high degree of flexibility for registration of histologic images with in-vivo/ex-vivo MR images, and can assist in more accurate evaluation of MRI findings. Boundary landmarks and internal landmarks of the same prostate have been used in a deformable registration algorithm. Mazaheri et al. describe a semi-automatic method by using a free-form deformation (FFD) algorithm based on B-splines [125]. This method enabled successful registration of anatomical prostate MR images to pathologic slices. Jacobs et al.[126] proposed a method for the registration and warping of MR images to histologic sections. This method consists of a modified surface-based registration algorithm followed by an automated warping approach using nonlinear thin plate splines to compensate for the distortions between the datasets.

There are two general approaches to map ex vivo histological PCa extent to pre-operative MR images. The first method, perhaps the more intuitive approach, is to reconstruct the 3D histologic volume, and then register the 3D histologic volume with the 3D MR volume [127, 128]. The second approach is to register each 2D histology slice to its corresponding 2D MRI slice separately [107, 129]. In the first approach, one critical prerequisite was the accurate reconstruction of the histologic volume; while in the second approach, the prerequisite was to determine the histology-MRI slice correspondence. In some cases, the former prerequisite may not be achievable hence the only solution is to take the second approach. There is an increasing interest in the registration of 3D histopathology with prostate MR imaging. Three-dimensional reconstruction of prostate histology facilitates these registration-based evaluations by reintroducing 3D spatial information lost during histology processing [130, 131]. Patel et al. [132] presented a scheme for the registration of digitally reconstructed whole mount histology to pre-operative in vivo mp-MRI using spatially weighted mutual information. McGrath et al. [133] used reference landmarks that are visible in both data sets to assist 3D histopathology reconstruction and thus can provide
important information on the deformation effects of fixation, and hence improved registration accuracy. Histostitcher, a software system designed to create a pseudo whole mount histology section from a stitching of four individual histology quadrant images, is another alternative for reconstructing pseudo whole-mount prostate images [134].

Registering pathologic information to mp-MRI is a challenging problem in developing a CADx system for mp-MRI (Figure 10). Chappelow et al. [135] described a method based on mutual information that registers T2W, DCE-MRI, and ADC. However, this method is based on 2D histology and requires considerable expertise to determine the correspondence between histologic and MR images. Orczyk et al. [136] described a method based on the present registration method and were the first to create a 3D counterpart within the same reference space between histology and both anatomical and functional sequences provided by prostate mp-MRI (Figure 11). The method enables a true, deformable transformation and achieves an accuracy of 1–2 mm. The registration of different MR images is critical considering prostate motion, especially related to rectal peristalsis. Orczyk et al. [136] used rigid registration to correct motion between difference sequences.

Although whole mount prostate histological analysis provides accurate label information for training a CADx system, whole mount histology is expensive and registering whole mount histologic slices with 3D mp-MRI is a challenging problem. Therefore, histologic interpretations from biopsy specimens are used to determine the ground truth in some studies [14, 62, 122, 137, 138]. In vivo biopsy can only label the pathology of the core inside the prostate. Radiologists must manually define lesion boundaries on mp-MRI retrospectively based on the biopsy results.

Meyer et al. [139] reviewed the registration methods of 3-D medical images and histopathology of the prostate. They examined the registration process and techniques for registering MRI or PET with whole-mounted prostatectomy specimens.

**CLINICAL APPLICATIONS**

**Diagnosis**

The functional MR imaging data, like DCE-MRI and MRS, are more complex and larger in amounts than anatomic MR imaging. There are clinical needs to develop fast, cost-effective, supportive techniques, such as computer-aided analysis tools, for easy and more reproducible diagnosis of prostate cancer. Researchers have focused on developing CADx methodology for automated prostate MRS classification and DCE-MRI analysis. Because all functional MR imaging techniques have their strengths and shortcomings, single technique cannot adequately detect and characterize PCa. The combination of anatomic (T2W) images and functional techniques has been shown to increase the accuracy of MR imaging for diagnosis of PCa. Table 1 compares the performance of the major published prostate CADx systems [13, 14, 16–18, 22, 26, 27, 36, 37, 39, 51, 52, 54–57, 62, 122, 129, 140–151]. Chan et al. were one of the first groups who implemented an mp-MRI CADx system for the diagnosis of prostate cancer [122]. In their approach they used line-scan diffusion, T2 and T2-weighted images to identify predefined areas of the peripheral zone of the prostate for the presence of prostate cancer. Viswanath et al. [129] present an mp-MRI CADx system for
PCa detection by integrating functional and structural information obtained via DCE and T2W MRI. Liu et al. [141] present fuzzy MRF models for prostate cancer detection of multispectral MR prostate images. Tiwari et al. [55] investigated the use of MR spectroscopy in combination with T2W MRI to identify the voxels that are affected by prostate cancer. They also introduced the use of wavelet embedding to map MRS and T2-W texture features into a common space. In a study by Peng et al. [27], the combination of 10th percentile ADC, average ADC, and T2-weighted skewness with CADx yielded an AUC value of 0.95 in differentiating prostate cancer from normal tissue. The combination achieved higher accuracy than any MR parameter alone. In a more recent study by Litjens et al [62], they developed a fully automated computer-aided detection system which consists of two stages. The first (detection) stage consists of segmentation of the prostate on the transversal T2W MRI, extraction of voxel features from the image volumes, classification of the voxels and candidate selection. The second (diagnosis) stage consists of candidate segmentation, candidate feature extraction and candidate classification. The system was evaluated on a large consecutive cohort of 347 patients, and yielded an AUC value of 0.889.

**Aggressiveness**

Treatment choice for prostate cancer is based on initial PSA level, clinical stage of disease, and Gleason score, together with baseline urinary function, comorbidities, and patient age [152, 153]. Therefore, there is an urgent clinical need to detect high-grade cancers and to differentiate them from the indolent, slow-growing tumors. The Gleason system, using a rating system to determine the grade of prostate cancer, remains one of the widely used prognostic factors in prostate cancer. The higher grade tumors have a tendency to grow quickly and to spread faster than lower grade tumors.

DWI, DCE-MRI, and MRS are noninvasive assessment methods of PCa aggressiveness. The Gleason grading system is a fundamental indicator of the aggressive nature of prostate cancer. Studies found that ADC image features correlate with Gleason scores [27, 28, 46, 154–156]. A study by Yamamura et al. found a highly significant negative correlation between ADC-value and the Gleason score, while MRS did not show a significant correlation [157]. Recently, Zhang et al. found that transrectal ultrasound (TRUS)-guided, MRI-directed biopsies improved the prediction of PCa aggressiveness in comparison with 12-core TRUS-guided biopsies. DWI directed biopsies had a superior performance when compared with MRS directed biopsies in the peripheral zone [6]. Diffusion of water molecules in tumor tissue was thought to reflect tissue architecture such as cell density and nucleus/cytoplasm ratio, and reductions in ADC values in tumor tissue in fact correlates well with increases in cellular density [158–160]. For these reasons, ADC value has received more attention as a predictor of Gleason score in prostate cancer.

DCE-MRI is based on the permeability of blood vessels and extravasation of contrast agent into the surrounding tissue. Investigators have observed that quantitative parameters (K\text{trans} and K\text{ep}) and semi-quantitative parameters (wash-in and wash-out) derived from DCE-MRI have the potential to assess the aggressiveness of PCa. Oto et al. found a moderate correlation between k\text{ep} and microvessel density of prostate cancer [154]. Peng et al. found K\text{trans} moderately correlate with Gleason scores [27].
*In vivo* MRS imaging has revealed a trend towards an increased (choline + creatine)/citrate (CC/C) ratio with increased Gleason score [161, 162]. This relationship has also been demonstrated by *ex vivo* HR-MAS MRS [163]. However, other *in vivo* MRS imaging studies have found no correlation between metabolite ratios and aggressiveness [164, 165].

On T2W MRI, changes in signal intensity for prostate cancer detection have been associated with its aggressiveness [166]. In a large retrospective study with 220 patients [166], T2W MRI and MRS imaging scores based on a three-point scale for clinical prostate cancer aggressiveness were significantly correlated to biologic markers such as androgen receptor levels, which were associated with prostate cancer progression. In that study, the combination of biomarkers with T2W MRI and MRS imaging results can discriminate clinically unimportant prostate cancer. If mp-MRI can potentially aid in identifying low-grade disease *in vivo*, this might allow PCa patients to opt for active surveillance rather than immediately opting for aggressive therapy. Lee et al. demonstrated that the simple measurement of the diameter of suspicious tumor lesions on DWI could improve the prediction of insignificant prostate cancer in candidates for active surveillance therapy [167].

Although these MRI metrics are related to Gleason score, the power and threshold value of each metric are different and how to combine these anatomic and functional MRI information is still a problem. Developing a computerized decision support system may help in noninvasive assessment of PCa aggressiveness. Recently, a system called Semi-Supervised MultiKernel Graph Embedding (SeSMiK-GE), was developed to quantitatively combine T2WI and MRS data for distinguishing benign versus cancerous, and high- versus low-Gleason grade PCa regions *in vivo* [148].

**Biopsy guidance**

Transrectal ultrasound (TRUS)-guided sextant or systematic prostate biopsy is the clinical standard for definitive diagnosis of prostate cancer. The Gleason score derived from biopsy specimens is important for appropriate treatment selection. However, PCa is often heterogeneous and multicentric [168]. In addition, the biopsy, which samples a small portion of the prostate, might not represent the whole gland efficiently. Traditionally, it is believed that Gleason score in systematic random TRUS-guided biopsy tends to downgrade the surgical specimen, because a less differentiated pattern may not have been sampled in the biopsy [169, 170]. Systematic random TRUS-guided biopsies often require repeated biopsy procedures, which are associated with discomfort and potential morbidity [171]. In order to reduce the overtreatment and the number of biopsies, lesions must be accurately detected, characterized and targeted during biopsy. More effective imaging-guided targeted biopsy techniques are under investigation in order to improve the detection rate of prostate biopsies.

Optimization of prostate biopsy requires addressing the shortcomings of standard systematic TRUS-guided biopsy, including false-negative rates, incorrect risk stratification, detection of clinically insignificant disease and the need for repeat biopsy. MRI is an evolving noninvasive imaging modality that increases the accurate localization of prostate cancer at the time of biopsy, and thereby enhances clinical risk assessment and improves the ability to appropriately counsel patients regarding therapy.
Use of mp-MRI for targeted prostate biopsies has the potential to reduce the sampling error associated with conventional biopsy by providing better disease localization and sampling, and also has a potential role in avoiding biopsy and reducing over detection/overtreatment. MRI-compatible biopsy systems were developed for this purpose [172]. More accurate risk stratification through improved cancer sampling may impact therapeutic decision making. Optimal clinical application of MRI targeted biopsy remains under investigation.

There are three different manners in which an MRI-detected lesion can be targeted for biopsy: (1) Direct targeting within the magnet using MR-compatible devices, also called in-bore MRI guided biopsy; (2) Use of fusion software to allow an MRI-defined lesion to be identified on ultrasound during a TRUS-guided biopsy procedure (Figure 12); or (3) Cognitive targeting, in which the physician reviews the MRI data before the procedure and attempts to target the suspected area during the TRUS-guided biopsy using anatomic landmarks as reference [173]. An MRI-guided robotic prostate biopsy system, named APT-MRI robotic biopsy system, has been reported with an accuracy within 2 mm [174]. A real-time phase-only cross correlation (POCC) algorithm-based sequence has been used in transrectal 3T in-bore MR-guided prostate biopsies [175]. Fusion of pre-biopsy MR images onto interventional TRUS images might increase the overall biopsy accuracy [176, 177]. A novel method to identify the 2D axial MR slice from a pre-acquired MR prostate volume that closely corresponds to the 2D axial TRUS slice obtained during prostate biopsy has been reported by Mitra et al. [178].

**Treatment planning and therapeutic response assessment**

MRI-based techniques are used for computer-aided treatment procedures such as treatment planning of radiotherapy, MRI-guided radioactive seeds placement in prostate brachytherapy, and MRI-guided local ablation procedures [179–190].

The excellent soft-tissue contrast of MRI means that the technique is having an increasing role in contouring the gross tumor volume (GTV) and organs at risk (OAR) in radiation therapy treatment planning systems (TPS). MRI-planning scans from diagnostic MRI scanners are currently incorporated into the planning process by being registered to CT data. The soft-tissue data from the MRI provides target outline guidance and the CT provides a solid geometric and electron density map for accurate dose calculation on the TPS computer [191].

A number of minimally invasive, focal, organ-preserving methods have been used in recent years as further alternatives to the radical treatment of prostate cancer [170]. The focal therapy methods used to date for the prostate include cryotherapy, high-intensity focused ultrasound, laser-induced thermal ablation, and radioactive seed placement. Mp-MRI makes it possible to determine the exact location of tumor foci that are generally accessible for ablation or radioactive seed placement. Moreover, mp-MRI can also monitor treatment during and after minimally invasive therapy. A CADx system for the prostate may have potential value in helping clinicians to target tumor foci during treatment.

Mp-MRI can also be used as an imaging biomarker for monitoring therapeutic response, including radiotherapy of localized prostate cancer [191] and systemic therapy for metastatic...
disease. Successful treatment response to therapy is usually depicted by reductions in signal intensity accompanied by ADC increases [192–194]. There are clinical needs to develop mp-MRI-based CADx systems for monitoring therapeutic response of the prostate in the future.

**DISCUSSION AND FUTURE DIRECTIONS**

Unlike breast and lung cancer, prostate cancer CADx systems for MR images have not been widely used in daily clinical work for detection or diagnosis. The majority of the prostate CADx systems reported the AUC in the range from 0.80 to 0.89 [179], while one reported AUCs of 0.96 [46], which represented a high performance. However, most systems generated lesion candidates based on manually selected ROIs, which may be data set dependent, and employed a relatively small data set. Validation on a large-scale data set with several hundred patients is required. A prostate CADx system should be tested in multicenter trials to make the systems widely usable in clinical work.

One challenge of prostate CAD is related to mp-MRI protocols. Both 3T protocols and endorectal coils have the advantage of increasing the signal-to-noise ratio (SNR). At 3T without the use of an endorectal coil (ERC), image quality can be comparable with that obtained at 1.5T with an endorectal coils [195]. Turkbey et al. found that dual-coil prostate MRI detected more cancer foci than non-endorectal coil MRI at 3T on T2W and DWI [196]. At 3 T MRI, DWI images and ADC maps using b = 1500 s/mm² should be considered more effective than those at b = 2000 s/mm² or b = 1000 s/mm² for detecting prostate cancer [50]. Most members of the PI-RADS steering committee recommend 3 T for prostate MRI. There is no consensus among experts concerning the potential benefits of the use of endorectal coils [12]. The impact of the mp-MRI protocol on CADx systems should been considered and researched in the future. The combination of T2W, DWI, and DCE-MRI is the most commonly used set of parameters for the detection or diagnosis of prostate cancer. MRS with other parameters is also used in some research. The introduction of new imaging modalities or new modality combinations for mp-MRI may lead to better CADx systems. Combining CAD prediction and PI-RADS into a combination score has the potential to improve diagnostic accuracy [197]. The MR PI-RADS system may provide a platform for CAD system development in the future.

The diagnostic value of these parameters for discrimination between benign and malignant tissue depend on the lesion’s location. The parameter values of PCa are in the range of those of nonmalignant diseases or conditions such as prostatitis, fibromuscular benign prostatic hyperplasia (BPH), post-biopsy hemorrhagic change, making for poor diagnostic value, especially in the transition zone. TZ and PZ cancer possess distinct quantitative imaging features on MRI. Computer-extracted parameters may be useful for cancer detection in the PZ, but are not suited in the TZ. In recent years, research focus has shifted from PZ prostate cancer to whole prostate cancer. There are more challenges in developing a CADx system for both PZ and TZ lesions than for PZ lesions only. Applications of anatomical segmentation from MRI as an additional input to ANN improve the accuracy of detecting cancerous voxels from MRSI [198]. A CAD system, utilizing two MRI sequences, such as T2-MRI and high-b-value (b = 2000 s/mm²) DWI, and texture features based on local binary
patterns, is able to detect the discriminative texture features for cancer detection and localization, and the performance of the CADx system was not dependent on the specific regions of the prostate [52]. Future direction should also include whether zonal segmentation of the prostate is necessary when some new imaging sequences being used.

*Ex vivo* whole mount prostate histological analysis provides more accurate label information for training a CADx system. However, whole mount histology is expensive, and registering whole mount histological slices with 3D mp-MRI is a challenging problem. This is especially true during the preparation of the prostate histological data for training a CADx system. Pathologists must collect a large amount of training data from many patients, apply reliable biomarkers for each patient, prepare blocks, scan a large number of histological slices, and manually define lesion boundaries on histological slices. However, these are laborious and time-consuming procedures. Therefore, the histological image preparation procedures need to be performed by some automatic methods to improve efficacy. A software system has been designed to create a pseudo whole mount histology section [134]. A computer-aided system to automatically grade pathological images according to the Gleason grading system has also been investigated [199]. A scheme, including automatic diagnosis from histologic images, 3D histologic reconstruction and registration, should be developed for ground truth definition in the future.

Image quantification methods, such as accurate image registration for motion correction, compartment modeling for functional parameters estimation, feature extraction in high dimensional data, automatic image classification for differentiating cancer from normal tissue, and correlation analyses among radiological data and genomic information, will play key roles in the future development of intelligent CAD systems.

Radiomics, as a high dimensional extraction of large amounts of image features with high throughput from radiographic images, can provide valuable diagnostic, prognostic or predictive information. Cameron et al. had developed a quantitative radiomics feature model for performing prostate cancer detection using mp-MRI [115]. Khalvati et al. [118] present new texture feature models for radiomics-driven detection of prostate cancer utilizing mp-MRI data. Radiomics are emerging as a useful tool for prostate cancer detection. Further work is needed to build radiomics-based CAD systems for prostate cancer diagnosis, treatment planning, treatment prediction and treatment response evaluation.

The Gleason grade of PCa is the most widely used prognostic factor for prostate cancer. MR metrics on T2W, DWI, DCE-MRI and MRS imaging relate to microenvironment and microstructure. Therefore these MR metrics can predict the Gleason grade of the cancer. Building a CAD system based on mp-MRI and Gleason score is feasible. It can play a significant role in prognostic prediction, guiding biopsy, identifying suitable patients under active surveillance and making a decision of appropriate treatment. CAD systems for prediction of Gleason score should be developed in the future.

As the anatomic information is important when analyzing functional data, T2W images are frequently used in mp-MRI CADx systems. T2W plus DWI and DCE-MRI are commonly used as the combinations. Chan et al. constructed a summary statistical map of the
Peripheral zone based on the utility of multichannel statistical classifiers by combining textural and anatomical features in PCA areas from T2W, DWI, proton density maps, and T2 maps [122]. Langer et al. included DCE-MRI and pharmacokinetic parameter maps as extra features to a CADx system for detecting prostate cancer at the peripheral zone [111]. They evaluated their system in predefined regions of interest, but on a per voxel basis. Vos et al. implemented a two-stage CADx system for prostate cancer using an initial blob detection approach combined with a candidate segmentation and classification using statistical region features [14]. Litjens et al. recently investigated a fully automated computer-aided detection system including a novel combination of segmentation, voxel classification, candidate extraction and classification [62].

Promising preliminary results have been obtained with CADx systems that combine the analysis of statistical, structural, and functional MR imaging features and the use of an adapted classification scheme. Likelihood maps have been obtained by combining information from mp-MRI using mathematical descriptors. These studies showed the discrimination between benign and malignant tissues is feasible with good performances [62, 111].

**CONCLUSION**

We comprehensively reviewed mp-MRI based, computer aided technology for prostate cancer detection. Prostate CADx systems are a complicated composition of preprocessing, segmentation, registration, feature extraction, and classification modules. There are some challenges in accurate registration of MRI and histopathology, which is important for ground truth definition. Clinical applications of computer aided systems include localization, diagnosis, staging, aggressiveness assessment, guiding biopsy, treatment planning, and therapeutic response assessment. Although the performance of some CADx systems is good, there is no such a system that has been wildly used in clinic. It is likely that more improvements in quantitative image analysis and computer-aided methods would need to be made in order to meet the clinical needs in near future work.

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**REFERENCES**

4. Pokorny MR, de Rooij M, Duncan E, Schroder FH, Parkinson R, Barentsz JO, Thompson LC. Prospective study of diagnostic accuracy comparing prostate cancer detection by transrectal


85. Mahapatra D, Buhmann JM. Prostate MRI segmentation using learned semantic knowledge and graph cuts. Biomedical Engineering, IEEE Transactions on. 2014; 61


92. Flores-Tapia, D., Thomas, G., Venugopa, N., McCurdy, B., Pistorius, S. Semi automatic MRI prostate segmentation based on wavelet multiscale products. In Engineering in Medicine and


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Abbreviation

T2W  T2-weighted
ADC  Apparent diffusion coefficient
DCE  Dynamic contrast-enhanced
MRS  Magnetic resonance Spectroscopy
DWI  Diffusion-weighted imaging
T1-PC (principal component of T1-weighted dynamic series)
T1  T1 mapping
T2  T2 mapping
SVM  Support vector machine
P-SVM  Probabilistic SVM
FLD  Fisher linear discriminant
MRFs  Markov random fields
NLDR  Nonlinear dimensionality reduction
CRF  Conditional random fields
EMPrAvISE Enhanced Multi-Protocol Analysis via Intelligent Supervised Embedding
QDA  Quadratic Discriminant Analysis
ANN  Artificial neural network
SeSMiK-GE  Semi Supervised Multi Kernel Graph Embedding
LDA  Linear discriminant analysis
PCA  Principal component analysis
PCA-VIP  Variable importance on projection measure for PCA
LLE  Locally linear embedding

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<th>Abbreviation</th>
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<tr>
<td>AUC</td>
<td>Area under a receiver operating characteristic curve</td>
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<tr>
<td>Se</td>
<td>Sensitivity</td>
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<tr>
<td>Sp</td>
<td>Specificity</td>
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<tr>
<td>FP</td>
<td>False positive</td>
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<td>FN</td>
<td>False negative</td>
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<tr>
<td>TZ</td>
<td>Transition zone</td>
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<td>PZ</td>
<td>Peripheral zone</td>
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<tr>
<td>CG</td>
<td>Central gland</td>
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<tr>
<td>WP</td>
<td>Whole prostate</td>
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<td>WMHS</td>
<td>Whole mount histological sections</td>
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Figure 1.
High-resolution T2-weighted MRI. T2-weighted MR images can differentiate the normal intermediate- to high-signal-intensity peripheral zone (Region 1) from the low-signal-intensity central and transition zones (Region 2).
Figure 2.
High-resolution T2-weighted MR images of prostate cancer. (A) There is a low–signal intensity lesion on the right peripheral zone (white arrows) at the mid-gland of the prostate. At prostatectomy, the lesion was classified as a Gleason grade 7 (4+3) prostate adenocarcinoma. (B) An ill-defined homogeneous low–signal intensity area at the left transition zone (white arrows) at mid-gland of the prostate in another patient. TRUS-guided biopsy showed a Gleason grade 8 (4+4) prostate adenocarcinoma on the corresponding position (Images from Neto JA, Parente DB: Multiparametric magnetic resonance imaging of the prostate. Magn Reson Imaging Clin N Am 2013, 21:409–426).
Figure 3.
Dynamic contrast enhanced MRI (DCE-MRI) of the prostate. (a) Axial T1 GRE unenhanced image. After contrast agent administration, an area with early enhancement is seen on the right in the peripheral zone (b, ROI1) with significant washout in the late-phase image (c). The curve (red) with early enhancement is a typical finding in the case of prostate cancer, while healthy prostate tissue is characterized by a steady slow enhancement (green). High transport constants Ktrans (e) and kep (f) can confirm suspicion of prostate cancer. Prostate adenocarcinoma with a Gleason score of 4+5=9 was diagnosed after prostatectomy (Image from Durmus, T, Baur, A, Hamm, B: Multiparametric magnetic resonance imaging in the detection of prostate cancer. Aktuelle Urol 2014, 45:119–126).
Figure 4.
Multiparametric MRI (mp-MRI) of the prostate. Axial T2 TSE (A) and coronal T2 TSE (B) images show a well-defined T2 hypointense lesion in the peripheral zone (arrow) with corresponding high signal on DWI (C) and low signal on the ADC map (D). Biopsy of this region was positive for Gleason 4 + 3 prostate cancer (Images from Yacoub, JH, Oto, A, Miller, FH: MR Imaging of the Prostate. Radiologic Clinics of North America 2014, 52:811–837).
Figure 5.
MR spectroscopy (MRS) of prostate cancer. (A) Axial T2-weighted MR images at the level of the prostate mid-gland to apex, shows a large hypointense lesion on the left peripheral zone. (B) A 3D MRS shows a normal spectrum on the right peripheral zone (red box) with normal choline plus creatine-to-citrate ratio of 0.48. In the voxel placed over the lesion on the left peripheral zone (blue box), the curve shows an increased choline peak and the citrate peak is markedly reduced. Random systematic biopsy showed a Gleason grade 9 (4 + 5) prostate adenocarcinoma on the left apex (Images from Neto JA, Parente DB: Multiparametric magnetic resonance imaging of the prostate. Magn Reson Imaging Clin N Am 2013, 21:409–426).
Figure 6.
Flowchart for computer aided detection of prostate cancer in mp-MRI.
Figure 7.
Prostate segmentation on MR images. Left: 2D MR image and segmentation results where the red curve represents the segmentation from a computer algorithm while the blue curve is the ground truth labeled by a radiologist. Right: 3D visualization after segmentation. The gold region is the prostate surface obtained by the computer algorithm while the red region is the ground truth.
Figure 8.
Figure 9.
Figure 10.
Registration between multiparametric MRI and histology.
Figure 11.
Figure 12.
MRI and ultrasound fusion for targeted biopsy of the prostate. (A and B) Anterior lesion of the high suspicious lesion identified on mp-MRI. (C) Real-time ultrasound targeting the corresponding lesion. (D and E) 3D models demonstrate the target (blue), prostate (brown), and biopsy cores (tan cylinders). (F) Radical prostatectomy pathology confirmed a 2.3 cm Gleason 8 (4+4) cancer centered in the right anterior prostate (Images from Sonn, GA, Margolis, DJ, Marks, LS: Target detection: Magnetic resonance imaging-ultrasound fusion–guided prostate biopsy. Urologic Oncology: Seminars and Original Investigations 2014, 32:903–911).
### TABLE 1

Summary of representative studies in the literature.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Modality</th>
<th>Validation</th>
<th>Region</th>
<th>Classifier</th>
<th>Data size</th>
<th>Performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chan et al. (2003)[122]</td>
<td>T2WI, ADC, T2</td>
<td>Biopsy</td>
<td>PZ</td>
<td>SVM, FLD</td>
<td>15</td>
<td>FLD, AUC=0.839; SVM, AUC=0.761</td>
</tr>
<tr>
<td>Puech et al. (2007)[39]</td>
<td>DCE</td>
<td>prostatectomy</td>
<td>PZ &amp; TZ</td>
<td>Software titled &quot;ProCAD&quot;</td>
<td>100</td>
<td>PZ, Se/Sp=100/49%; TZ, Se/Sp=100/40%</td>
</tr>
<tr>
<td>Tiwari et al. (2007)[54]</td>
<td>MRS</td>
<td>Biopsy</td>
<td>WP</td>
<td>Spectral clustering</td>
<td>14</td>
<td>Se=77.8%, FP=28.92%, and FN=20.88%</td>
</tr>
<tr>
<td>Vos et al. (2008)[36]</td>
<td>DCE</td>
<td>WMHS</td>
<td>PZ</td>
<td>SVM</td>
<td>34</td>
<td>AUC = 0.83</td>
</tr>
<tr>
<td>Viswanath et al. (2008)[140]</td>
<td>DCE</td>
<td>WMHS</td>
<td>WP</td>
<td>LLE and Consensus Clustering</td>
<td>6</td>
<td>Se=60.72%, Sp=83.24%</td>
</tr>
<tr>
<td>Viswanath et al. (2009)[129]</td>
<td>T2WI, DCE</td>
<td>WMHS</td>
<td>WP</td>
<td>Random forest</td>
<td>6</td>
<td>AUC = 0.815</td>
</tr>
<tr>
<td>Vos et al. (2009)[37]</td>
<td>DCE</td>
<td>WMHS</td>
<td>PZ</td>
<td>SVM</td>
<td>38</td>
<td>AUC = 0.80</td>
</tr>
<tr>
<td>Liu et al. (2009)[141]</td>
<td>T2W, T2,ADC, DCE</td>
<td>WMHS</td>
<td>PZ</td>
<td>Fuzzy MRF Model</td>
<td>11</td>
<td>Se=89.58%, Sp=87.50%</td>
</tr>
<tr>
<td>Tiwari et al. (2009)[55]</td>
<td>MRS</td>
<td>prostatectomy</td>
<td>WP</td>
<td>NLDR</td>
<td>18</td>
<td>Se=89.33%, Sp=79.79%</td>
</tr>
<tr>
<td>Artan et al. (2010)[142]</td>
<td>T2, ADC, DCE</td>
<td>Biopsy</td>
<td>PZ</td>
<td>cost-sensitive CRF</td>
<td>21</td>
<td>AUC = 0.79</td>
</tr>
<tr>
<td>Vos et al. (2010)[143]</td>
<td>T2WI, DCE</td>
<td>WMHS</td>
<td>PZ</td>
<td>SVM</td>
<td>29</td>
<td>AUC = 0.89</td>
</tr>
<tr>
<td>Viswanath et al. (2011)[144]</td>
<td>T2W, DWI, DCE</td>
<td>WMHS</td>
<td>WP</td>
<td>EMPrAvisE</td>
<td>12</td>
<td>AUC=0.77</td>
</tr>
<tr>
<td>Lopes et al. (2011)[145]</td>
<td>T2WI</td>
<td>WMHS</td>
<td>WP</td>
<td>SVM, AdaBoost</td>
<td>17</td>
<td>SVM, Se/Sp=83.91%; AdaBoost, Se/Sp=83.93%</td>
</tr>
<tr>
<td>Liu and Yetik (2011)[26]</td>
<td>T2W, DWI, DCE</td>
<td>WMHS</td>
<td>WP</td>
<td>SVM</td>
<td>20</td>
<td>AUC = 0.89</td>
</tr>
<tr>
<td>Sung et al. (2011)[146]</td>
<td>DCE</td>
<td>prostatectomy</td>
<td>PZ &amp; TZ</td>
<td>SVM</td>
<td>42</td>
<td>PZ, Se/Sp=89/89%; TZ, Se/Sp=91.64%</td>
</tr>
<tr>
<td>Tiwari et al. (2012)[56]</td>
<td>T2WI, MRS</td>
<td>WMHS</td>
<td>WP</td>
<td>Random forest</td>
<td>36</td>
<td>AUC = 0.89</td>
</tr>
<tr>
<td>Reference</td>
<td>Modality</td>
<td>Validation</td>
<td>Region</td>
<td>Classifier</td>
<td>Data size</td>
<td>Performance</td>
</tr>
<tr>
<td>-----------</td>
<td>----------</td>
<td>------------</td>
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<td>-------------</td>
</tr>
<tr>
<td>Viswanath et al.(2012)[22]</td>
<td>T2WI</td>
<td>WMHS</td>
<td>PZ &amp; CG</td>
<td>QDA</td>
<td>22</td>
<td>CG, AUC=0.86; PZ, AUC=0.73</td>
</tr>
<tr>
<td>Vos et al.(2012)[14]</td>
<td>T1, T2, ADC, DCE</td>
<td>Biopsy</td>
<td>WP</td>
<td>LDA</td>
<td>200</td>
<td>Se=0.74, at a FP level of 5 per patient.</td>
</tr>
<tr>
<td>Niaf et al.(2012)[18]</td>
<td>T2W, DWL, DCE</td>
<td>WMHS</td>
<td>PZ</td>
<td>SVM</td>
<td>30</td>
<td>AUC = 0.89</td>
</tr>
<tr>
<td>Arjan et al.(2012)[147]</td>
<td>T2, ADC, T1-PC</td>
<td>WMHS</td>
<td>WP</td>
<td>SVM</td>
<td>15</td>
<td>Se=76%, Sp=86%</td>
</tr>
<tr>
<td>Shah et al.(2012)[17]</td>
<td>T2WI, ADC, DCE</td>
<td>WMHS</td>
<td>PZ</td>
<td>SVM</td>
<td>31</td>
<td>f-measure=89%</td>
</tr>
<tr>
<td>Matulewicz et al.(2013)[57]</td>
<td>MRS</td>
<td>WMHS</td>
<td>WP</td>
<td>ANN</td>
<td>18</td>
<td>AUC=0.968</td>
</tr>
<tr>
<td>Hambrock et al.(2013)[13]</td>
<td>T2WI, DWL, DCE</td>
<td>prostatectomy</td>
<td>PZ &amp; TZ</td>
<td>in-house-developed CAD system</td>
<td>34</td>
<td>experienced observers, AUC=0.91</td>
</tr>
<tr>
<td>Tiwari et al.(2013)[148]</td>
<td>T2WI, MRS</td>
<td>WMHS</td>
<td>WP</td>
<td>SeSMK-GE</td>
<td>29</td>
<td>AUC = 0.89</td>
</tr>
<tr>
<td>Peng et al. (2013)[27]</td>
<td>T2WI, ADC, DCE</td>
<td>prostatectomy</td>
<td>WP</td>
<td>LDA</td>
<td>48</td>
<td>AUC = 0.95</td>
</tr>
<tr>
<td>Ginsburg et al.(2014)[149]</td>
<td>T2WI, DWL, DCE</td>
<td>WMHS</td>
<td>PZ and CG</td>
<td>PCA-VIP</td>
<td>108</td>
<td>CG, AUC=0.85; PZ, AUC=0.79</td>
</tr>
<tr>
<td>Stember et al.(2014)[51]</td>
<td>T2WI, ADC</td>
<td>Biopsy</td>
<td>TZ</td>
<td>Naive Bayes classifier</td>
<td>18</td>
<td>predicted TZ tumor in all test patients</td>
</tr>
<tr>
<td>Niaf et al.(2014)[150]</td>
<td>T2WI, DWL, DCE</td>
<td>prostatectomy</td>
<td>WP</td>
<td>P-SVM</td>
<td>48</td>
<td>AUC = 0.889</td>
</tr>
<tr>
<td>Garcia Molina et al.(2014)[16]</td>
<td>T2WI, ADC, DCE</td>
<td>prostatectomy</td>
<td>PZ</td>
<td>incremental learning ensemble SVM</td>
<td>12</td>
<td>Se=84.4%, Sp=78.0%</td>
</tr>
<tr>
<td>Litjens et al.(2014)[62]</td>
<td>T2WI, DWL, DCE, PDWI</td>
<td>Biopsy</td>
<td>WP</td>
<td>Random Forest</td>
<td>347</td>
<td>AUC = 0.889</td>
</tr>
<tr>
<td>Kwak et al.(2015)[52]</td>
<td>T2WI, DWI</td>
<td>Biopsy</td>
<td>PZ and TZ</td>
<td>SVM</td>
<td>244</td>
<td>AUC of 0.89</td>
</tr>
<tr>
<td>Zhao et al.(2015)[151]</td>
<td>T2WI biopsy/follow-up</td>
<td>PZ and CG</td>
<td>ANN</td>
<td></td>
<td>71</td>
<td>CG, AUC=0.821; PZ, AUC=0.849</td>
</tr>
</tbody>
</table>
## Table 2

### Validation of CADx systems

<table>
<thead>
<tr>
<th>Reference</th>
<th>Ground truth on the histology</th>
<th>Candidate on MR image</th>
<th>Image Registration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chan et al. (2003)[122]</td>
<td>Biopsy</td>
<td>MO</td>
<td>NA</td>
</tr>
<tr>
<td>Puech et al. (2007)[39]</td>
<td>Needle biopsy or prostatectomy</td>
<td>MO</td>
<td>NA</td>
</tr>
<tr>
<td>Tiwari et al. (2007) [54]</td>
<td>Biopsy</td>
<td>Sextant location determined by radiologist</td>
<td>NA</td>
</tr>
<tr>
<td>Vos et al. (2008)[36]</td>
<td>WMHS+ MO</td>
<td>MO</td>
<td>3D rendering mode</td>
</tr>
<tr>
<td>Viswanath et al.(2008)[140]</td>
<td>WMHS+ MO</td>
<td>MANTRA</td>
<td>Multimodal image registration</td>
</tr>
<tr>
<td>Viswanath et al.(2009)[129]</td>
<td>WMHS</td>
<td>MANTRA</td>
<td>Multimodal image registration</td>
</tr>
<tr>
<td>Vos et al.(2009)[37]</td>
<td>WMHS</td>
<td>Not specified</td>
<td>Not specified</td>
</tr>
<tr>
<td>Tiwari et al.(2009)[55]</td>
<td>WMHS+ sextant boundaries</td>
<td>A joint review session of trial imagers and pathologists</td>
<td>NA</td>
</tr>
<tr>
<td>Artan et al.(2010)[142]</td>
<td>WMHS+ MO</td>
<td>Tumor location is transferred to the in vivo MRI from histologic images +ex vivo MRI</td>
<td>NA</td>
</tr>
<tr>
<td>Vos et al.(2010)[143]</td>
<td>WMHS+ MO</td>
<td>MO</td>
<td>Mutual information registration</td>
</tr>
<tr>
<td>Viswanath et al.(2011)[144]</td>
<td>WMHS+ MO</td>
<td>Registration from histologic images</td>
<td>MACMI</td>
</tr>
<tr>
<td>Lopes et al.(2011)[145]</td>
<td>WMHS+ drawn by urologists</td>
<td>Drawn by urologists</td>
<td>manual correspondence</td>
</tr>
<tr>
<td>Sung et al.(2011)[146]</td>
<td>Radical prostatectomy + MO</td>
<td>The radiologist matched the pathologic slices with corresponding MRI</td>
<td>NA</td>
</tr>
<tr>
<td>Tiwari et al.(2012)[56]</td>
<td>WMHS</td>
<td>MO +ex vivo MRI</td>
<td>Manual registration</td>
</tr>
<tr>
<td>Viswanath et al.(2012)[23]</td>
<td>WMHS+MO</td>
<td>Registration from histologic images</td>
<td>Multimodal Elastic Registration</td>
</tr>
<tr>
<td>Vos et al.(2012)[14]</td>
<td>Needle biopsy</td>
<td>Combining the findings with, histopathology of MR-guided samples by radiologist.</td>
<td>NA</td>
</tr>
<tr>
<td>Artan et al.(2012)[147]</td>
<td>WMHS+MO</td>
<td>MO +ex vivo MRI</td>
<td>Manual registration</td>
</tr>
<tr>
<td>Shah et al.(2012)[17]</td>
<td>WMHS+MO</td>
<td>Not specified</td>
<td>PSM</td>
</tr>
<tr>
<td>Matulewicz et al.(2013)[57]</td>
<td>WMHS+MO</td>
<td>MO</td>
<td>Manual registration</td>
</tr>
</tbody>
</table>

*Acad Radiol. Author manuscript; available in PMC 2017 August 01.*
<table>
<thead>
<tr>
<th>Reference</th>
<th>Ground truth on the histology</th>
<th>Candidate on MR image</th>
<th>Image Registration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tiwari et al. (2013)[148]</td>
<td>WMHS+MO</td>
<td>MO</td>
<td>Manual registration</td>
</tr>
<tr>
<td>Peng et al. (2013)[27]</td>
<td>WMHS</td>
<td>MO</td>
<td>Manual registration</td>
</tr>
<tr>
<td>Ginsburg et al. (2014)[149]</td>
<td>WMHS+MO</td>
<td>Registration from histologic images</td>
<td>Nonlinear registration</td>
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<tr>
<td>Stember et al. (2014)[51]</td>
<td>Needle biopsy</td>
<td>Not specified</td>
<td>NA</td>
</tr>
<tr>
<td>Niaf et al. (2014)[150]</td>
<td>Prostatectomy+MO</td>
<td>MO</td>
<td>Manual registration</td>
</tr>
<tr>
<td>Garcia Molina et al. (2014)[16]</td>
<td>Prostatectomy+MO</td>
<td>MO</td>
<td>Manual registration</td>
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<tr>
<td>Litjen et al. (2014)[62]</td>
<td>Needle biopsy</td>
<td>Not specified</td>
<td>NA</td>
</tr>
<tr>
<td>Kwak et al. (2015)[52]</td>
<td>Needle biopsy</td>
<td>Determined by radiologists</td>
<td>NA</td>
</tr>
<tr>
<td>Zhao et al. (2015)[151]</td>
<td>Biopsy</td>
<td>MO</td>
<td>NA</td>
</tr>
</tbody>
</table>

MO: manual outlined regions of lesions

MANTRA: multi-Attribute, non-initializing, texture reconstruction based ASM

MACMI: multi-attribute, higher order mutual information based elastic registration scheme

PSM: patient-specific molds

WMHS: whole mount histological sections

NA: no registration was used