Comprehensive Framework for Computer-Aided Prostate Cancer Detection in Multi-Parametric MRI

by

Audrey Chung

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in
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This thesis consists of material all of which I authored or co-authored: see Statement of Contributions included in the thesis. This is a true copy of the thesis, including any required final revisions, as accepted by my examiners.

I understand that my thesis may be made electronically available to the public.
Statement of Contributions

The following two papers are used in this thesis. They are described below:


This paper is incorporated in Chapter 3 of this thesis.

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Abstract

Prostate cancer is the most diagnosed form of cancer and one of the leading causes of cancer death in men, but survival rates are relatively high with sufficiently early diagnosis. The current clinical model for initial prostate cancer screening is invasive and subject to overdiagnosis. As such, the use of magnetic resonance imaging (MRI) has recently grown in popularity as a non-invasive imaging-based prostate cancer screening method. In particular, the use of high volume quantitative radiomic features extracted from multi-parametric MRI is gaining attraction for the auto-detection of prostate tumours since it provides a plethora of mineable data which can be used for both detection and prognosis of prostate cancer.

Current image-based cancer detection methods, however, face notable challenges that include noise in MR images, variability between different MRI modalities, weak contrast, and non-homogeneous texture patterns, making it difficult for diagnosticians to identify tumour candidates. In this thesis, a comprehensive framework for computer-aided prostate cancer detection using multi-parametric MRI was introduced. The framework consists of two parts: i) a saliency-based method for identifying suspicious regions in multi-parametric MR prostate images based on statistical texture distinctiveness, and ii) automatic prostate tumour candidate detection using a radiomics-driven conditional random field (RD-CRF).

The framework was evaluated using real clinical prostate multi-parametric MRI data from 20 patients, and both parts were compared against state-of-the-art approaches. The suspicious region detection method achieved a 1.5% increase in sensitivity, and a 10% increase in specificity and accuracy over the state-of-the-art method, indicating its potential for more visually meaningful identification of suspicious tumour regions. The RD-CRF method was shown to improve the detection of tumour candidates by mitigating sparsely distributed tumour candidates and improving the detected tumour candidates via spatial consistency and radiomic feature relationships. Thus, the developed framework shows potential for aiding medical professionals with performing more efficient and accurate computer-aided prostate cancer detection.
Acknowledgements

First and foremost, I would like to thank my supervisor Prof. Alexander Wong (fondly referred to as the “research dad”), whose unwavering support and eagerness to pursue all things computer vision related has long been a source of inspiration. Thank you for being a wonderful mentor, and for guiding me as I grow as a researcher. Your constant enthusiasm for new (and particularly ridiculous) ideas has shown me some of the most enjoyable aspects of research.

I would like to thank Prof. Paul Fieguth and Prof. Bryan Tripp for reviewing my thesis. Carving out the time from your exceedingly busy schedules to read and revise my thesis is greatly appreciated.

I would also like to thank the colourful and wildly eclectic mosaic of people that make up the Vision and Image Processing Research Group. Your collective love of collaborations has regularly demonstrated that great research happens at the crossroads of innovation and teamwork. There is no doubt that my research experience would have been noticeably less vibrant without you all.

In addition, I would like to thank my family and friends for their constant support and blind faith in my abilities. Thanks for listening to me spew information about topics you were unfamiliar with (but hopefully are now).

Lastly, I would like to thank the Natural Sciences and Engineering Research Council (NSERC) of Canada for funding my research in this thesis.
Dedication

This thesis is dedicated to my uncle, Peter Chung, who was recently diagnosed with prostate cancer.
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Nomenclature

$\alpha_i$  Saliency determined from statistical distinctiveness graphical model

$\alpha_{\text{max}}$  Maximum calculated saliency determined from statistical distinctiveness graphical model

$\beta_{i,j}$  Distinctiveness metric using Kullback-Leibler divergence

$\hat{A}$  Estimation of apparent diffusion coefficient map

$\Lambda_i$  Set of texture representations that corresponds to saliency $\alpha_i$

$\mathcal{E}$  Set of edges in graph

$\mathcal{V}$  Set of vertices (nodes) representing optimal label field $Y^*$

$\Phi_i$  $i^{th}$ principal component of $h(x)$

$\Psi(Y, X)$  Energy function factorized from different feature functions

$\Psi_c$  Set of potential functions for a clique $c$

$\psi_p$  Inter-voxel spatial and radiomic features-based pairwise function

$\psi_u$  Data-driven unary function

$\mu_i, \mu_j$  Mean of $t^r_i, t^r_j$

$\Sigma_i, \Sigma_j$  Covariance of $t^r_i, t^r_j$

$x_i$  Observations associated with voxel

$A$  Apparent diffusion coefficient map
a  Single value in apparent diffusion coefficient map
b  Gradient strength and pulse duration (b-value) in diffusion-weighted imaging
b_γ Given b-value associated with reference signal S_γ
b_i Given b-value associated with signal S_i
C  Set of cliques in conditional random field
c  Clique in conditional random field
C_p(i) Pairwise clique structure for a given neighbourhood N(i)
D  Strength of the diffusion in diffusion-weighted imaging
e_{ij} Edge, connectivity between a pair of nodes representing the relationship between two voxels
G(\mathcal{V}, \mathcal{E}) Graph realization of the radiomics-driven conditional random field
h(x) Textural representation comprised of sets of texture features
m  Number of representative texture atoms
M_{xy}(0) Initial magnetic resonance signal value
M_{xy}(t) Decayed magnetic resonance signal at time t
n  Number of neighbours in a clique
N(i) Set of n neighbours (neighbourhood)
P  Conditional joint probability density function
S  Diffusion-weighted signal in diffusion-weighted imaging
S_0 Signal intensity without diffusion weighting in diffusion-weighted imaging
S_γ Reference signal associated with b-value b_γ
t(x) Compact textural representation of h(x) produced using PCA
\{t(x)\}_z Subset of compact texture features used for testing
Sparse texture model

$\theta_i, \theta_j$ Representative texture atom corresponding to healthy or suspicious tissue

$V(x)$ Local sub-volume centred at $x$ in correlated diffusion imaging

$v_c$ Random variable within clique $c$

$w$ Number of principal component analysis (PCA) components selected

$X$ Set of observations (voxels) in MR image

$x$ Single voxel in MR image

$Y$ Label field associated with observation set $X$

$y$ Single label associated with voxel $x$

$Y^*$ Optimal solution for label field $Y$

$y^*$ Optimal solution for single label $y$

$y_\phi$ Subset of labels (clique) in the surrounding neighbourhood

$Z(X)$ Normalizing function given observation set $X$
Chapter 1

Introduction

Prostate cancer refers to the presence of uncontrolled cellular growth (i.e., a malignant tumour) in the prostate, and can negatively affect urinary and erectile function in men. The prostate is a walnut-sized gland in the male reproductive system responsible for making seminal fluid, which mixes with sperm from the testicles to make semen. The prostate is located below the bladder and in front of the rectum, surrounding part of the urethra (the tube that carries urine and semen through the penis). While changes to the growth and behaviours of the cells in the prostate may lead to non-cancerous (benign) conditions such as prostatitis and benign prostatic hyperplasia (BPH), they can also lead to precancerous conditions such as prostatic intraepithelial neoplasia (PIN), proliferative inflammatory atrophy (PIA), and atypical small acinar proliferation (ASAP) [3].

Prostate cancer is the most diagnosed form of cancer (excluding non-melanoma skin cancers) in Canadian and American men. According to the Canadian Cancer Society [4], there was an estimated 24,000 new cases and 4,100 deaths from it in 2015, making it the third most deadly cancer and accounting for approximately 10% of cancer deaths in Canadian men. Similarly in the United States, there was an estimated 220,800 new cases and 27,540 deaths from prostate cancer in 2015, making it the second most deadly cancer and accounting for approximately 9% of cancer deaths in American men [5]. The median patient survival time for metastatic prostate cancer is between 12.2 to 21.7 months [6]. However, prognosis is relatively good if the prostate cancer is detected early. As such, fast and reliable prostate cancer screening methods are crucial and can greatly impact patient survival rate, as the five-year survival rate in Canada is 96% for patients diagnosed with prostate cancer before the metastatic stage [7].
1.1 Current Cancer Screening Approaches

There are a variety of tests that are presently being used to screen for prostate cancer. Typically, the current clinical model for initial prostate cancer screening employs a digital rectal exam (DRE) and/or a prostate-specific antigen (PSA) test. The DRE is generally done as part of a routine physical exam for adult males over the age of 50 (or in the 40s in the case of men at higher risk of prostate cancer). A healthcare professional performs a DRE by inserting a gloved finger into the rectum and feeling the prostate gland through the rectum wall to check for enlargement or abnormalities. Generally done in tandem with the DRE, the PSA test measures the amount of the PSA protein in the blood. A protein made by the cells in the prostate gland, small amounts of PSA can normally be found in the blood. While many physicians refer to a normal PSA level as 4.0 ng/mL of blood or less, the level of PSA varies and tends to increase with age. A PSA level of 2.5 ng/mL is typical for men younger than 50, with the upper limit of the normal PSA range increasing by 1.0 ng/mL per decade of age [3].

Given a positive DRE or an elevated PSA, a patient undergoes a follow-up transrectal ultrasound (TRUS) guided multicore biopsy for risk stratification. TRUS imaging uses ultrasound to image the prostate via a small probe inserted into the rectum. The probe projects sound waves through the rectum and records the corresponding echoes. The biopsy involves the systematic regional sampling of the prostate where typically eight or more tissue samples are collected [8]. However, these samples are subject to sampling error as prostate tumours imaged by TRUS tend to appear isoechoic and are difficult to differentiate from surrounding tissue [9].

The PSA in particular has recently come under scrutiny, as recent studies [10, 11] have demonstrated that the PSA test has a significant risk of overdiagnosis. Only an estimated 50% of men who were screened as having elevated levels of PSA are actually diagnosed with prostate cancer. This oversensitivity leads to expensive, painful, and often unnecessary needle biopsies, and subsequent overtreatment [10–12]. These TRUS guided prostate biopsies cause discomfort, possible sexual dysfunction, and increased hospital admission rates due to infectious complications [13, 14]. In addition, the biopsies have a chance of missing the cancerous tissue altogether. The challenge diagnosticians currently face is how to improve prostate cancer screening by reducing the overdiagnosis due to conventional screening methods while still maintaining a high sensitivity.
1.2 Computer-aided Detection via MRI

The use of magnetic resonance imaging (MRI) has recently grown in popularity as a non-invasive imaging-based prostate cancer screening method. MRI is a medical imaging technique used in radiology to tomographically image anatomy and physiological processes. Using a magnetic field and radio frequency pulses, MRI repeatedly acquires detailed two-dimensional images (or “slices”) over a tissue volume. MRI has been shown to be a viable alternative to the clinical prostate cancer screening methods currently in use.

A diagnosis through MRI, however, requires an experienced medical professional to extensively review the data due to the subtle differences between cancerous and healthy tissue. Manual labelling of image data is time-consuming, and can lead to diagnostic inconsistencies due to variability between radiologists (inter-observer variability) and the variability of a radiologist over multiple sittings (intra-observer variability) [15–17]. To help raise the consistency of radiologists, the European Society of Urogenital Radiology (ESUR) introduced the Prostate Imaging – Reporting And Diagnosis System (PI-RADS) as a common set of criteria [18]. PI-RADS is a set of guidelines for interpreting multiple MRI images, and aims to raise the consistency between diagnosticians through a common set of criteria.

1.3 Multi-Parametric MRI

Within the realm of MRI, there are various types or modalities of MRI that allow for different tissue details to be emphasized. One basic type of MRI is T2-weighted MRI which focuses on measuring the fluid content of tissue, allowing for improved soft tissue contrast relative to other medical imaging techniques [19, 20]. Another commonly used modality is diffusion-weighted imaging (DWI); sensitive to the diffusion process of water molecules in biological tissues, DWI has also been shown to provide promising delineation between cancerous and healthy prostate tissue [21–23]. While T2-weighted and DWI have both been shown to provide diagnostically relevant information for prostate cancer localization, the use of multiple MRI modalities (multi-parametric MRI) has been shown to improve prostate cancer screening [22, 24]. This is done via the extraction and combination of unique information and features from each modality.

Despite PI-RADS and further development to standardize diagnostic practices across multi-parametric MRI [25], there is still a level of subjectiveness across different medical professionals when assessing MR images that can lead to inter-observer and intra-observer
variability. In addition, notable challenges include noise in MR images, weak contrast, non-homogeneous texture patterns, variability between different MRI modalities, and lack of registration and/or misalignment between different MRI modalities. These challenges make it difficult for diagnosticians to accurately and efficiently identify tumour candidates. As such, computer-aided cancer detection methods are being developed to help the physicians with the process.

1.4 Proposed Framework and Thesis Contributions

Figure 1.1: The proposed comprehensive framework for computer-aided prostate cancer detection via suspicious region detection and tumour candidate classification using multi-parametric MRI. In this thesis, the following multi-parametric MRI modalities were used: T2-weighted imaging (T2w), apparent diffusion coefficient maps (ADC), computed high-b diffusion-weighted imaging (CHB-DWI), and correlated diffusion imaging (CDI).

The purpose of this thesis is to propose a comprehensive framework for computer-aided prostate cancer detection via suspicious region detection and tumour candidate classification using multi-parametric MRI. There are two main contributions:

1. A saliency-based method for identifying suspicious regions in multi-parametric MR prostate images based on statistical texture distinctiveness is presented in Chapter 3.
In this approach, a sparse texture model is learned via expectation maximization from features derived from multi-parametric MR prostate images, and the statistical texture distinctiveness-based saliency constructed using the model is used to identify suspicious regions.

2. An approach for automatic prostate cancer detection using a radiomics-driven conditional random field (RD-CRF) framework is detailed in Chapter 4. In addition to the high-throughput extraction and utilization of a comprehensive set of voxel-level quantitative radiomic features, the proposed RD-CRF framework leverages inter-voxel spatial and radiomic feature relationships to ensure that the auto-detected tumour candidates exhibit interconnected tissue characteristics reflective of malignant prostate tumours.

The two main contributions of this thesis focus on creating a comprehensive framework to provide decision support and aid medical professionals with consistently and accurately localizing prostate cancer (shown in Figure 1.1). In this thesis, a review of relevant background concepts and state-of-the-art methods are presented in Chapter 2. Problem formulation, methods, and comparison test results against other state-of-the-art methods using patient data are detailed for the two main contributions in Chapters 3 and 4, respectively. Lastly, conclusions, recommendations, and future works are discussed in Chapter 5.
Chapter 2

Background

In this chapter, background information for the thesis and related works for each proposed contribution are presented. The background theory for the used MRI modalities is presented in Section 2.1, and the mathematical formulation for conditional random fields (CRF) is described in Section 2.2. An introduction to radiomics and related research is outlined in Section 2.3. A brief description of the existing types of approaches and their limitations are discussed to lead into the thesis contributions. The relevant work for suspicious region identification is discussed in Section 2.3.1, and the applicable literature for prostate tumour detection is presented in Section 2.3.2.

2.1 Imaging Modalities

In this thesis, the following multi-parametric MRI modalities were used: T2-weighted imaging [26], apparent diffusion coefficient maps [27], computed high-b diffusion-weighted imaging [28], and correlated diffusion imaging [29]. The modalities are summarized in the following subsections.

2.1.1 T2-weighted

One of the basic MRI pulse sequences, T2-weighted (T2w) [26] imaging is a MR imaging modality that characterizes the sensitivity of tissue using the differences in transverse (spin-spin) relaxation time of the applied magnetic field. $T_2$ is a time constant that characterizes the time required for the magnetic resonance signal to irreversibly decay to 37% (or $1/e$) of its initial value. The transverse relaxation decay is defined as:
Figure 2.1: Sample T2-weighting images from eight different patients. Note that cancerous regions are extremely difficult to identify visually in this modality.

\[ M_{xy}(t) = M_{xy}(0)e^{-t/T_2} \]  

where \( M_{xy}(t) \) is the decayed signal at time \( t \), and \( M_{xy}(0) \) is the initial signal value.

T2w imaging has been shown to provide some useful information for classification due to a small reduction in signal for cancerous tissue in the prostate gland [22]. Figure 2.1 shows some sample T2w prostate MR images.

### 2.1.2 Apparent Diffusion Coefficient

Diffusion-weighted imaging (DWI) is an imaging modality in which the sensitivity of the tissue to the Brownian motion water molecules is measured through the application of lobe gradients (pairs of opposing magnetic field gradient pulses) [23]. The diffusion-weighted signal \( S \) is formulated as:

\[ S = S_0e^{-bD} \]  

where \( S_0 \) is the signal intensity without diffusion weighting, \( b \) is the degree of applied diffusion weighting and indicated the gradient strength and pulse duration, and \( D \) is the strength of the diffusion. The diffusion-weighted images are typically generated using
different $b$ values, and can be used to estimate apparent diffusion coefficient (ADC) maps via least-squares or maximum likelihood methods \[23\] by rearranging 2.2 for the strength of diffusion $D$:

$$A = \frac{1}{b} \ln \frac{S}{S_0}$$

(2.3)

Cancerous tissue in ADC maps (shown in Figure 2.2) is usually presented with a darker intensity relative to surrounding healthy tissue \[27\].

### 2.1.3 Computed High-b Diffusion Weighted Imaging

Previous research has shown that high b-values in DWI data (e.g., b-values greater than 1,000s/mm\(^2\)) allow for increased delineation between healthy and cancerous tissue \[28, 30\]. Due to hardware limitations, acquiring high b-value images for prostate imaging is infeasible. CHB-DWI is a computational model for reconstructing high b-value DWI data using low b-value acquisitions \[28, 31\].

Using the general equation for a diffusion-weighted signal $S_i$:

$$S_i = S_0 e^{-(b_i-b_s)A}$$

(2.4)

Figure 2.2: Sample apparent diffusion coefficient maps computed from diffusion-weighted images from eight different patients. While lower resolution than T2w images, there is increased contrast between cancerous and healthy tissue.
where $S_\gamma$ is the reference signal corresponding to $b_\gamma$ and $A$ is the ADC map, an estimate $\hat{A}$ for the ADC map can be formulated as a Bayesian estimation problem [28]:

$$\hat{A} = \arg \max_A P(S|A)$$

(2.5)

$$\hat{A} = \arg \max_A \prod_i P(S_i|A)$$

(2.6)

where $P(S|A)$ is the conditional probability of the signal given $A$, and $S_i$ is a single DWI measurement corresponding to b-value $b_i$.

Using the estimate $\hat{A}$, computed diffusion-weighted images $\hat{S}_i$ can be obtained for any desired b-value $b_i$ as follows:

$$\hat{S}_i = S_\gamma e^{-(b_i-b_\gamma)\hat{A}}$$

(2.7)

Our patient data includes CHB-DWI images (Figure 2.3) constructed at a b-value of 2,000$s/mm^2$ using a Bayesian model with the least-squares estimation used to estimate our ADC maps.

### 2.1.4 Correlated Diffusion Imaging

Correlated Diffusion Imaging (CDI) [29] is a new diffusion MRI modality that leverages the joint correlation in signal attenuation across multiple gradient pulse strengths and timings.
to improve delineation between cancerous and healthy tissue. The overall characterization of the water diffusion is better represented via the correlation of signal attenuation across all b-values within a local sub-volume, which is obtained via signal mixing [29]:

$$CDI(x) = \int \cdots \int_{b_0}^{b_n} S_0(x) \cdots S_n(x) P(S_0(x), \ldots S_n(x)) \prod_{v(x)} V(x) \times dS_0(x) \ldots dS_n(x)$$  \hspace{1cm} (2.8)

where $b_i$ represents the utilized b-values, $x$ is the spatial location, $S$ is the acquired signals, $P$ represents the conditional joint probability density function, and $V(x)$ is the local sub-volume centred at $x$. Figure 2.4 shows some sample CDI prostate images.

2.2 Conditional Random Fields

First proposed by Lafferty et al. [32], conditional random fields (CRFs) are a framework for building probabilistic models to label data. There are several advantages of CRFs over hidden Markov models [33] and stochastic grammars, such as the ability to relax the strong independence assumptions made in those models. In addition, CRFs also bypass the label bias problem of maximum entropy Markov models [34] and other discriminative Markov models. The label bias problem [32] refers to the susceptibility of these models to favour states with few successor states, i.e., the transitions leaving a given state compete only
against each other, rather than against all other transitions in the model. CRFs solve this bias problem by having a single exponential model for the joint probability of the entire set of labels given the observations.

CRFs can be formulated as a graph $G(V, E)$, where $V$ is the set of vertices (nodes) representing the states $Y$ and $E$ is the set of edges of the graph. Given $G$, $(X, Y)$ is a conditional random field where the random variables $y \in Y$ obey the Markov property with respect to the graph when conditioned on $X$. In this thesis, we use an undirected CRF, the probability distribution of which can be represented using a product of non-negative functions of the maximal cliques of graph $G$ [35]:

$$p(v) = \frac{1}{Z} \prod_{c \in C} \Psi_c(v_c)$$

(2.9)

where factors $\Psi_c$ are potential functions of $v_c$ within clique $c \in C$; conditionally independent nodes belong to different cliques and do not appear within the same factor.

The general model for a CRF is formulated as

$$P(Y|X) = \frac{1}{Z(X)} \prod_{c \in C} \Psi_c(X_c, Y_c)$$

(2.10)

where $P(Y|X)$ is the conditional probability of labels $Y$ given observations $X$, and $Z(X)$ is the normalization function given the observations $X$.

### 2.3 Radiomics

More recently, a particularly promising and powerful approach to cancer detection that could have significant potential for prostate cancer detection is radiomics [2,36–39], which involves the high-throughput extraction and utilization of a large amount of quantitative features for characterizing tumour phenotype. Radiomics facilitates for a high-dimensional mineable feature space that can be utilized for both detection and prognosis [36]. Studies on lung and head-and-neck cancer patients have confirmed the prognostic power of radiomics features when it comes to patient outcome prediction for personalized medicine [36–38]. However, the prognostic capability of radiomics features has only very recently been investigated for prostate cancer detection (general framework shown in Figure 2.5) and the quantitative characterization of prostate tumour phenotype.
2.3.1 Suspicious Region Identification

One specific area of research is the identification of suspicious regions to aid physicians with performing a more efficient and accurate diagnosis. The current method for identifying suspicious regions is to threshold apparent diffusion coefficient (ADC) maps, as low ADC values are associated with tumorous tissue [22]. Cameron et al. [1] proposed a threshold-based approach where tissue associated with ADC values within a threshold range are automatically identified as suspicious.

\[ y = \begin{cases} 
1 & a \in A, a < 1000 \times 10^{-6} mm^2/s \\
0 & \text{otherwise} 
\end{cases} \]  (2.11)

where \( a \) is a single value in the ADC map \( A \). However, this method depends on fixed thresholds, making it susceptible to noisy MR images and ADC variations across different sets of multi-parametric MRI data.

2.3.2 Computer-Aided Prostate Tumour Detection

Current methods for radiomics-driven automatic computer-aided prostate tumour detection typically use a supervised method trained on a set of low-level features calculated from multispectral MR images. Lemaître et al. [40] recently published a comprehensive review of
state-of-art methods for prostate cancer detection and diagnosis. These radiomics-driven methods use hand-crafted feature sets and some combination of the following classifiers: Bayesian, random forest, logistic regression, neural network, linear discriminant, GentleBoost, support vector machines, relevance vector machines, fuzzy Markov random fields, and conditional random fields.

Madabhushi et al. [41] extracted three-dimensional texture features from MRIs that had been corrected for background inhomogeneity and nonstandardness. For each image voxel, a trained Bayesian classifier assigns a malignancy “likelihood” to each feature independently, and the “likelihood” images are then combined using an optimally weighted feature combination scheme. Using similar features as [41], Tiwari et al. proposed a method that combines structural and metabolic imaging data for prostate cancer detection and detected cancerous regions within prostate tissue using a random forest classifier.

Duda et al. [42] published a semi-automatic multi-image texture analysis method that simultaneously analysed several images (each acquired under different conditions) of the same organ. In addition to the features used by [41], Duda et al. also extracted fractal-based and run length features and assessed the potential of multi-image texture analysis using different classifiers including logistic regression, a neural network, and support vector machines.

Litjens et al. [43] used features that represent pharmacokinetic behaviour, symmetry and appearance, and other anatomical aspects. In a two-stage computer-aided prostate cancer detection system, Litjens et al. detected initial candidates via multi-atlas-based prostate segmentation using a selective and iterative method for performance level estimation (SIMPLE) [44]. The authors then experimented with three different classifiers for voxel classification and candidate classification: linear discriminant, GentleBoost, and random forest.

Similar to Litjens et al., Vos et al. [45] also developed a two-stage classification approach for computer-aided prostate cancer detection. Vos et al. analysed lesion candidates and discriminated prostate cancer from benign abnormalities using a linear discriminant classifier. Ozer et al. [46] used parametric images derived from DCE MRI, and proposed the use of relevance vector machines (RVM) with a Bayesian framework. Ozer et al. then evaluated the method against support vector machines (SVM) with the same framework.

Khalvati et al. [39] proposed a multi-parametric MRI texture feature model for radiomics-driven prostate cancer analysis. The texture feature model, based on the one proposed by Peng et al. [47], comprises of 19 low-level texture features extracted from each MRI modality, including features extracted from the gray-level co-occurrence matrix (GLCM). Khalvati et al. [2] more recently published radiomics-driven models as an extension of the
previous texture feature model. An attempt at designing comprehensive quantitative feature sequences, the radiomics-driven models include additional MRI modalities, additional low-level features, and feature selection.

Recently, the use of random fields for prostate cancer detection has grown in popularity. The use of fuzzy Markov random fields (MRFs) in particular has been investigated [48–50], as fuzzy MRFs are an unsupervised method for cancerous tissue classification. Liu et al. [48] proposed a new method for estimating the parameters of the Markovian distribution of the measured data, and applied it to feature vectors extracted from multispectral prostate MRI datasets for prostate cancer detection. An extension on previous work, Ozer et al. [49] also propose the use of fuzzy MRFs as an unsupervised alternative to the previously proposed SVM and RVM, and evaluate the classifiers using feature vectors formed from the peripheral zone of multispectral prostate MRI datasets.

Lastly, Artan et al. [50] presented a cost-sensitive SVM cancer localization method as an extension to the conventional SVM for prostate cancer detection. Trained via a full grid search over the $\nu$ and $\gamma$ values (SVM kernel parameters) to determine the optimal parameters, the cost-sensitive SVM shows improved results relative to the conventional SVM. Artan et al. also proposed a new segmentation method by combining conditional random fields (CRF) with a cost-sensitive framework. Using node potential parameters $w$ obtained from SVM and subsequently estimated edge potential parameters $\kappa$, the cost-sensitive CRF demonstrated improved cost-sensitive SVM results by incorporating spatial information.
Chapter 3

Suspicious Region Identification in Multi-Parametric Prostate MRI

In this chapter, the proposed method for identifying suspicious regions using prostate multi-parametric MRI is described. The components of the method are region-based textural representations, sparse texture model, statistical textural distinctiveness, and suspicious region identification via saliency, and are presented in Section 3.2, Section 3.3, Section 3.4, and Section 3.5, respectively. The patient data used in this thesis is described in detail in Section 3.6. Experimental setup and applicable quantitative metrics are outlined in Section 3.7, and both quantitative and visual results are shown in Section 3.8. At the end of this chapter, a brief summary of the method and implications of the results are discussed.

3.1 Problem Formulation

One specific area of research is the identification of suspicious regions to aid physicians with performing a more efficient and accurate diagnosis. The current method for identifying suspicious regions is to threshold apparent diffusion coefficient (ADC) maps, as low ADC values are associated with tumorous tissue [22]. Cameron et al. [1] proposed a threshold-based approach where tissue associated with ADC values within a threshold range are automatically identified as suspicious. However, this method depends on fixed thresholds, making it susceptible to noisy MR images and ADC variations across different sets of multi-parametric MRI data.
Figure 3.1: Proposed framework for identifying suspicious regions using prostate multi-parametric MRI. Unique texture features extracted from different MRI modalities are used to learn a sparse texture model, and suspicious regions are identified via a statistical textural distinctiveness-based saliency map.

To facilitate a more reliable diagnosis, a novel method for identifying suspicious regions indicative of potential prostate cancer using texture-based saliency in multi-parametric MR images is proposed. The proposed method uses unique texture information from each MRI modality to learn a sparse texture model, and better characterize suspicious tissue within a patient’s MRI data.

A novel method is proposed for identifying suspicious regions to better aid physicians with performing more efficient and accurate diagnoses. The proposed method uses multi-parametric MR images and incorporates cross-modality texture features to better identify suspicious regions via statistical textural distinctiveness. Figure 3.1 shows the general algorithmic framework developed.
3.2 Region-based Textural Representations

Region-based (or image patch-based) textural representations are used to allow for the characterization of texture features indicative of suspicious regions in prostate MR images. For region-based textural representations, we incorporate the feature set proposed by Khalvati et al. [39], which consists of sets of 19 low-level texture features extracted each from T2-weighted (T2w) images, apparent diffusion coefficient (ADC) maps, computed high-b diffusion-weighted imaging (CHB-DWI) data, and correlated diffusion imaging (CDI) data, to better capture healthy and cancerous tissue characteristics. These MRI modalities were selected based on their potential to separate cancerous from healthy prostate tissue.

The sets of texture features are combined into a single textural representation \( h(x) \), and a compact version of the textural representation is produced using principal component analysis (PCA). PCA [51] is used to increase the variance between the elements of the texture descriptor and to improve the efficiency of the subsequent sparse texture model and statistical textural distinctive model stages. A compact textural representation \( t(x) \) is produced using the \( w \) principal components of \( h(x) \) with the highest variance:

\[
\begin{align*}
  t(x) &= \{\Phi_i(h(x))\}_{1 \leq i \leq w} \\
  \text{where } \Phi_i \text{ is the } i^{\text{th}} \text{ principal component of } h(x). \\
\end{align*}
\]  

where \( \Phi_i \) is the \( i^{\text{th}} \) principal component of \( h(x) \). While \( w \) can be selected based on variance compactness, \( w \) components of \( h(x) \) were selected to represent 90% of the variance of all the textural representations as determined through extensive empirical testing via parameter grid search.

3.3 Sparse Texture Model

To characterize healthy and suspicious tissue for a patient, a sparse texture model is learned using the extracted multi-parametric MRI texture features [39]. The sparse texture model incorporates unique texture features from each MRI modality to learn tissue characteristics via cross-modality texture information. Thus, the sparse texture model can better identify healthy and suspicious tissue.

Using a subset \( t(x)_T \) of the compact textural representations \( t(x) \) as training data, a global texture model is defined to represent the heterogeneous characteristics of healthy and suspicious prostate tissue. As global texture modelling is computationally expensive, we generalize an MRI slice as being composed of a set of regions where a particular texture
pattern is repeated over a given area. In addition, the number of areas with unique texture patterns is assumed to be much fewer than the number of individual voxels in the training data, allowing for a sparse texture model to fully characterize the MRI data.

Using this generalization, we can establish a textural sparsity assumption, and the global textural characteristics of prostate tissue can be well-represented using a small set of distinctive local textural representations. This allows for the use of a sparse texture model, defined as a set of \( m \) representative texture atoms:

\[
T^r = \{ t^r_i | 1 \leq i \leq m \} \tag{3.2}
\]

The sparse texture model used in the proposed method is a set of representative texture atoms corresponding to healthy or suspicious tissue, where each texture atom represents the mean and covariance (i.e., \( t^r_i \sim \mathcal{N}(\mu_i, \Sigma_i) \)) of a particular texture pattern characteristic of healthy or suspicious tissue. The representative atoms in the sparse texture model are learned via expectation maximization [52] in a Gaussian mixture model [53].

### 3.4 Statistical Textural Distinctiveness

Suspicious regions in prostate MRI data can be characterized as areas that are highly unique and texturally distinct. Using the concept of statistical textural distinctiveness [54], we quantify the distinctiveness of texture patterns and uncover the underlying saliency by using the statistical relationship between texture patterns across different MRI modalities.

To define statistical textural distinctiveness between two representative texture atoms (denoted as \( t^r_i \) and \( t^r_j \)) in the sparse texture model, we use Kullback-Leibler (KL) divergence [55] to measure the statistical difference between the representative texture atoms in the sparse texture model:

\[
\beta_{i,j} = \log \frac{\Sigma_j}{\Sigma_i} - w + \text{trace}(\Sigma_i^{-1} \Sigma_j^{-1}) + \frac{(\mu_j - \mu_i)^T \Sigma_i^{-1} (\mu_j - \mu_i)}{2} \tag{3.3}
\]

where \( u \) is the number of PCA components selected, \( \mu_i \) and \( \mu_j \) represent the mean of \( t^r_i \) and \( t^r_j \), respectively, and \( \Sigma_i \) and \( \Sigma_j \) represent the covariance of \( t^r_i \) and \( t^r_j \), respectively. Thus, the distinctiveness metric \( \beta_{i,j} \) increases as the texture patterns become more distinct from one another.
3.5 Suspicious Region Detection via Saliency Map Computation

As the majority of prostate tissue is considered to be healthy, salient regions can be interpreted as suspicious due to the uniqueness and statistical occurrence of the corresponding cross-modality texture characteristics. Given a subset of compact texture features used for testing (denoted as \( t(x)_Z \)), the saliency map for a given MRI image can be computed using the previously determined statistical textural distinctiveness graphical model. The saliency \( \alpha_i \) is defined as:

\[
\alpha_i = \sum_{j=1}^{m} \beta_{i,j} P(t^*_i | t(x)_Z) \tag{3.4}
\]

where \( P(t^*_i | Z) \) is the occurrence probability of \( t^*_i \) in \( t(x)_Z \).

For \( \Lambda_i \) being the set of texture representations that corresponds to saliency \( \alpha_i \), voxels belonging to salient representative texture atoms \( \Lambda_i \) (i.e., \( \alpha_i > \alpha_{\text{max}}^2 \)) are classified as regions of suspicious tissue, with all other voxels classified as healthy tissue. That is, each voxel \( x \) in a given MRI image is assigned a label \( y \):

\[
y = \begin{cases} 
1 & x \in \Lambda_i, \alpha_i > \frac{\alpha_{\text{max}}}{2} \\
0 & \text{otherwise}
\end{cases} \tag{3.5}
\]

3.6 Patient Data

The performance of the proposed framework was evaluated using the MP-MRI data of 20 patients (17 with cancer and 3 without cancer) acquired using a Philips Achieva 3.0T machine at Sunnybrook Health Sciences Centre, Toronto, Ontario, Canada. Institutional research ethics board approval for this study was obtained at Sunnybrook Health Sciences Centre. For each patient, the following MP-MRI modalities were obtained (Table 3.1): i) T2w, ii) DWI, and iii) CDI. The patients’ age ranged from 53 to 83. Table 3.1 summarizes the information about the 20 patients’ datasets used in this research. MP-MRI imaging data were processed in the ProCanVAS (Prostate Cancer Visual Analysis System) platform developed jointly by Sunnybrook Research Institute, Toronto, ON, Canada and the
University of Waterloo, Waterloo, ON, Canada. The data set includes segmentation information to isolate the prostate, and ground truth data for tumour size and location. All imaging data were reviewed and marked as healthy and cancerous tissue by a radiologist with 18 and 13 years of experience interpreting body and prostate MRI, respectively, and the radiologists’ markings were confirmed by pathology data to be Gleason scores of 7 or above.

### Table 3.1: Description of the prostate T2w, DWI, and CDI imaging data

<table>
<thead>
<tr>
<th>Modality</th>
<th>DFOV ($cm^2$)</th>
<th>Resolution ($mm^3$)</th>
<th>TE (ms)</th>
<th>TR (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>T2w</td>
<td>22 × 22</td>
<td>0.49 × 0.49 × 3</td>
<td>110</td>
<td>4,687</td>
</tr>
<tr>
<td>DWI</td>
<td>20 × 20</td>
<td>1.56 × 1.56 × 3</td>
<td>61</td>
<td>6,178</td>
</tr>
<tr>
<td>CDI</td>
<td>20 × 20</td>
<td>1.56 × 1.56 × 3</td>
<td>61</td>
<td>6,178</td>
</tr>
</tbody>
</table>

### 3.7 Experimental Setup

Each patient dataset had corresponding T2w images, ADC maps, CHB-DWI data, and CDI data. Using the radiologist contour of the prostate, a rectangle cropped around the prostate gland was selected as the region of interest (ROI) for each MRI slice. The performance of each method was evaluated using leave-one-patient-out cross-validation. A subset of the training texture features were randomly selected and used to train the classifier, and the voxels in a single MRI slice were classified as either healthy or cancerous tissue and assigned the saliency value of the nearest texture atom.

In addition, the number of texture atoms used to compute the spare texture model (as described in Subsection 3.3) was varied to determine the optimal number of representative texture atoms for identifying suspicious regions in prostate MR images. The ADC-based method was compared against the proposed texture distinctiveness method (TD) via sensitivity, specificity, and accuracy metrics.

\[
\text{Sensitivity} = \frac{TP}{P} \quad \text{Specificity} = \frac{TN}{N}
\]

\[
\text{Accuracy} = \frac{TN + TP}{N + P}
\]
where the performance of each method was quantified by the metrics' closeness to one. TP is the number of voxels in the intersection of the identified cancerous tissue and the radiologist’s tissue segmentation, TN is the number of voxels not in the identified tissue that are also not in the radiologist’s segmentation, N is the number of voxels not in the radiologist segmented tissue and P is the number of voxels in the radiologist segmented tissue.

### 3.8 Experimental Results

The proposed textural distinctiveness method (TD) was evaluated using both four-atom and six-atom sparse texture models. Table 3.2 shows the performance metrics for the ADC-based method [1] and the proposed method. The testing data contained 52 tumours (as identified by an experienced radiologist) across the slices from 13 different patients.

<table>
<thead>
<tr>
<th></th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADC-based method [1]</td>
<td>0.7911</td>
<td>0.7107</td>
<td>0.7115</td>
</tr>
<tr>
<td>TD (4 texture atoms)</td>
<td>0.8088</td>
<td>0.8285</td>
<td>0.8283</td>
</tr>
<tr>
<td>TD (6 texture atoms)</td>
<td>0.8103</td>
<td>0.8303</td>
<td>0.8301</td>
</tr>
</tbody>
</table>

As seen in Table 3.2, the proposed TD method outperformed the ADC-based method [1] in terms of sensitivity, specificity, and accuracy. While there was only a relatively small increase in sensitivity (approximately 1.5%), TD demonstrated an increase of at least 10% in specificity and accuracy relative to the ADC-based method. This is especially beneficial, as a low specificity negatively impacts a diagnostician’s ability to perform quick and accurate assessments of MRI data. By increasing specificity, TD minimizes the number of wrongly detected regions that contain no tumor candidates. This is important for procedures such as radical prostatectomy where an extremely high specificity rate is required.

Figure 3.2 shows a visualization of the suspicious regions detected using the ADC-based method [1] and the proposed TD method using four and six representative texture atoms. While all methods identify the cancerous regions as suspicious, the ADC-based method in particular has a tendency to be over-sensitive and often identifies a large portion of the prostate tissue as suspicious. A visual inspection of the identified suspicious regions
shows that TD consistently produces spatially compact and useful regions regardless of the number of texture atoms.

### 3.9 Summary

A novel method was proposed to aid physicians in efficiently and accurately diagnosing patients via the identification of suspicious regions in prostate MR images. We extracted unique textural information from different MRI modalities, and used a sparse texture model to learn tissue texture characteristics. As the majority of prostate tissue is considered to be healthy, texturally distinct regions can be interpreted as suspicious due to the uniqueness and statistical occurrence of the corresponding cross-modality texture characteristics.

![Figure 3.2: Visual comparison of identified suspicious regions (shown in red) between (a) ADC-based method [1], (b) TD using four texture atoms, (c) TD using six texture atoms, and (d) radiologist segmented regions.](image)
The proposed statistical textural distinctiveness approach (using four-atom and six-atom sparse texture models) was evaluated against the ADC-based method [1]. In both cases, statistical textural distinctiveness produced higher sensitivity, specificity, and accuracy values than the state-of-art ADC-based method. In additional, statistical textural distinctiveness also identified suspicious regions on a per patient basis, rather than relying on a fixed ADC value characteristic of typical cancerous tissue (as is the case with the ADC-based threshold method). Thus, statistical textural distinctiveness shows potential for more flexible and visually meaningful identification of suspicious tumour regions.

Achieving a 1.5% sensitivity increase and a 10% specificity and accuracy increase over current the state-of-the-art method, the proposed statistical textural distinctiveness method shows potential for more visually meaningful identification of suspicious tumour regions and allows for more accurate tumour classification. Particularly noteworthy is the 10% specificity increase of the proposed statistical textural distinctiveness approach, as a high specificity is crucial for imaging-guided procedures such as biopsy or radical prostatectomy.

Future work includes the further investigation of additional MRI modalities, and the use of spatial consistency to enforce more compact identified suspicious areas. Applications include identifying suspicious regions for clinicians to better stream-line a patient’s diagnosis, and automatically identifying regions of interest for computer-aided tumour detection methods. In the next chapter, a novel algorithm for prostate cancer tumour detection is presented.
In this chapter, the method for prostate tumour detection is presented. The quantitative radiomics-driven feature model is introduced in Section 4.2, and the radiomics-driven conditional random field (RD-CRF) is formulated in Section 4.3. Experimental setup is outlined in Section 4.4, and results were obtained using the data previously described in Chapter 3. Experimental results are shown in Section 4.5, and detailed discussion of the results is presented in Section 4.6. A brief summary of the method and related future works are discussed at the end of this chapter.

4.1 Problem Formulation

Inspired by genetic profiling for cancer risk stratification (genomics), Khalvati et al. [2] introduced comprehensive and quantitative radiomics feature models consisting of hundreds of radiomics features derived from MP-MRI data via feature selection and classification for the purpose of voxel-resolution prostate tumour detection. As the first voxel-level radiomics feature models based on MP-MRI data designed for prostate tumour characterization to be introduced, the results of the studies demonstrated the prognostic power of radiomics for the purpose of automatic voxel-resolution prostate cancer detection. Nevertheless, one limitation of such voxel-resolution radiomics-driven prostate tumour detection approaches is that they utilize the extracted quantitative radiomics features associated with individual voxels on an independent basis, and do not account for the interconnected tissue characteristics reflective of cancerous tumours. As such, the incorporation of additional information
regarding the spatial and radiomics feature relationships between voxels has significant potential for achieving a more reliable voxel-resolution radiomics-driven prostate cancer detection performance.

We propose a novel approach for automatic prostate cancer detection based on a radiomics-driven conditional random field (RD-CRF) framework. The proposed RD-CRF framework utilizes the quantitative MP-MRI radiomics feature model proposed in [2] and incorporates not only the spatial relationships between voxels, but the quantitative radiomics feature relationships between voxels into a conditional random field model. This facilitates for the enforcement of interconnected tissue characteristics reflective of cancerous tumours, thus better representing the actual cancerous tissue phenotype.

Here, we present the methodology and underlying principles of a novel method for automatic pixel-resolution prostate cancer detection via a radiomics-driven conditional random field (RD-CRF) framework. While existing prostate cancer detection methods can generally identify tumours, these methods tend to identify regions that are sparsely distributed (as shown in Section 4.2) that are unrepresentative of typical cancerous tissue. As such, the proposed RD-CRF framework leverages the full set of pixel-level quantitative radiomics features and incorporates inter-pixel spatial and feature relationships via a conditional random field to detect tumours more reliably by taking into account the inter-connected tissue characteristics reflective of cancerous tumours.

An overview of the proposed framework for pixel-resolution prostate cancer detection is shown in Figure 4.1. First, a large amount of quantitative radiomics features are extracted from MP-MRI data using a quantitative radiomics feature model. Second, a classifier trained using such quantitative radiomics features from training data is used to perform initial pixel-resolution cancer detection. A radiomics-driven conditional random field (RD-CRF) framework is then used to perform final pixel-resolution cancer detection using the initial results produced by the classifier and the full set of extracted quantitative radiomics features of the current patient case. The detailed methodology behind each step of the proposed framework is described below.

4.2 Quantitative Radiomics-driven Feature Model

First, the proposed framework incorporates the quantitative radiomics feature model proposed in [2], which was specifically designed for extracting a large number of quantitative radiomics features from MP-MRI data for the purpose of prostate cancer detection. Sets of 96 low-level radiomics features (4 from first-order and 72 from second-order statistical features, 8 from Kirsch edge detection, and 12 from Gabor filters [2]) were extracted
Figure 4.1: The proposed framework for automatic voxel-resolution prostate cancer detection. A large amount of quantitative radiomics features are extracted from MP-MRI data using a quantitative radiomics feature model. A classifier trained using quantitative radiomics features from training data is used to perform initial voxel-resolution cancer detection. A radiomics-driven conditional random field (RD-CRF) framework is then used to perform final voxel-resolution cancer detection using the initial results produced by the classifier and the full set of extracted quantitative radiomics features of the current patient.

from each of following 8 different forms of MP-MRI data: i) T2-weighted (T2w) imaging data, ii) apparent diffusion coefficient (ADC) data, iii) computed high-b diffusion-weighted imaging (CHB-DWI) data, iv) correlated diffusion imaging (CDI) data, and v-viii) diffusion-weighted imaging (DWI) data acquired at 4 individual b-values (0, 100, 400, and 1000s/mm$^2$). This results in a total of 768 features used within the comprehensive quantitative radiomics feature model that can better capture both healthy and cancerous tissue characteristics. A SVM classifier was then trained using quantitative radiomics features extracted from training data using this radiomics feature model, which can then be used to perform voxel-resolution classification on a new unseen patient case.

The MP-MRI modalities used in this radiomics feature model were selected based on their potential to separate cancerous from healthy prostate tissue. T2w imaging data
has been shown to provide some localization information for tumour candidates due to a small reduction in signal in the cancerous tissue [22]. Similarly, cancerous tissue in ADC data is typically represented by a darker intensity relative to surrounding tissue [27]. Previous research has shown that high-b values in diffusion weighted imaging data allows for increased delineation between cancerous and healthy tissue [28,30]. Lastly, CDI data takes advantage of joint correlation in signal attenuation across multiple gradient pulse strengths and timings to improve delineation between cancerous and healthy tissue [29,56].

One challenge with classifying tissues into cancerous and healthy as performed in existing state-of-the-art voxel-resolution radiomics-driven cancer detection methods (e.g., [2]) is that the detected tumour candidates tend to be sparsely distributed across the prostate region (see Figure 4.2), leading to high false positive and negative rates. This sparsity is a result of existing approaches not taking into account inter-voxel spatial relationships and radiomics feature relationships between voxels, and results in situations where there are missing voxels within a tumour candidate, as well as scattered voxels that may be considered as cancerous voxels. As cancerous tissue tends to be localized to one or two regions of a prostate, and exhibit interconnected tissue characteristics, this motivates the use of a probabilistic radiomics-driven framework that better accounts for such characteristics to improve prostate cancer detection performance, which will be described in the next section. Therefore, the proposed framework incorporates cross-modality spatial priors to enforce spatial relationships between voxels while maintaining relationships between cross-modality radiomic features.

Figure 4.2: Examples of tumour candidates detected by [2]. The identified regions (shown in red) are sparsely distributed across the prostate tissue, contrary to the compact regions characteristic of cancerous tissue.
4.3 Radiomics-driven Conditional Random Field

The theory behind the radiomics-driven conditional random field (RD-CRF) framework used to perform final voxel-resolution cancer detection based on the initial results produced by the classifier and the full set of extracted quantitative radiomics features of the current patient case can be described as follows. Conditional random fields \[32, 35\] were first proposed by Lafferty \[32\] et al. and have previously been used for image labelling \[57\]. The proposed RD-CRF model extends the conditional random field (CRF) model to leverage the full set of voxel-level quantitative radiomics features derived from MP-MRI data while taking into account the spatial relationships and quantitative radiomics feature relationships between voxels to better enforce interconnected tissue characteristics.

Given the prostate MP-MRI data pertaining to the patient, the tumour detection problem is formulated as the conditional probability of a set of binary labels (i.e., suspected prostate tumour or healthy tissue). The RD-CRF framework models the conditional probability of the binary label field \(Y\) and corresponding \(X\) observations as follows:

\[
P(Y|X) = \frac{1}{Z(X)} \exp(-E(Y, X)) (4.1)
\]

where \(Z(X)\) is the normalizing function and \(E(Y, X)\) represents the energy function factorized from different feature functions. The labelling problem of the prostate as healthy or cancerous is optimized using a Maximum A Posteriori (MAP) approach where minimizing the energy function \(E(Y, X)\) produces the best classification of healthy tissue and tumour candidates:

\[
Y^* = \arg \min_Y E(Y, X) (4.2)
\]

where \(Y^*\) is the optimal solution given the patient’s MP-MRI. As \(Y^*\) maximizes the probability of healthy and cancerous tissue classification via the minimization of \(E(Y, X)\), the formulation of the energy function is crucial in obtaining an accurate binary label field.

\(E(Y, X)\) can be formulated as some combination of unary and pairwise potential functions, and is generally defined as:

\[
E(Y, X) = \sum_{i=1}^{n} \psi_u(y_i, X) + \sum_{\varphi \in C} \psi_p(y_\varphi, X) (4.3)
\]
\( E(Y, X) \) incorporates the data-driven unary function \( \psi_u(y_i, X) \) and the inter-voxel spatial and radiomic features-based pairwise function \( \psi_p(y_\phi, X) \) into the model. In Equation 4.3, \( y_i \in Y \) is a single state in the set \( Y = \{ y_i \}_{i=1}^n \) and represents the label of a single voxel in the binary label field. \( X = \{ x_i \}_{i=1}^n \) is the corresponding set of observations from the prostate MR-MRI data as represented by the quantitative radiomic features, and \( y_\phi \in Y \) is the subset of labels (clique) in the surrounding neighbourhood. For each node \( i \), there is a set of neighbours \( N(i) \):

\[
N(i) = \{ j | j = 1 : n, j \neq i \}
\]

(4.4)

where \(|N(i)|\) was empirically chosen to be 24 (i.e., a 5 \( \times \) 5 neighbourhood centred at node \( i \)). The proposed RD-CRF framework uses a pairwise clique structure

\[
C = \{ C_p(i) \}_{i=1}^n
\]

(4.5)

\[
C_p(i) = \{ (i, j) | j \in N(i) \}
\]

(4.6)

to enforce the consistency of spatial and radiomics feature relationships in the identification of tumour candidates. Each node \( i \) represents a single voxel in a patient’s MP-MRI data, and \( N(i) \) represents the corresponding neighbouring axial voxels.

For the proposed RD-CRF framework, the results of the trained classifier are used as the data-driven unary term \( \psi_u(.) \), while the pairwise term \( \psi_p(.) \) is factorized based on the set of clique structures \( C \) for a pairwise clique \( c \in C \).

By taking full advantage of the comprehensive set of 768 quantitative radiomics features extracted from the different MP-MRI modalities along with spatial and radiomics feature relationships within the RD-CRF framework, one can better characterize healthy and cancerous tissue phenotype and thus achieve improved final voxel-resolution cancer detection.

To obtain the final voxel-resolution tumour detection results, the energy function \( E(Y, X) \) is minimized using gradient descent, and the binary label field is assigned to each voxel as \( y^* \in \) optimal solution \( Y^* \) (as defined in Equation 4.2):
\[ y^* = \begin{cases} 
1 & \text{cancerous tissue} \\
0 & \text{healthy tissue} 
\end{cases} \]  

Graph \( G(\mathcal{V}, \mathcal{E}) \) is the realization of the proposed RD-CRF for an axial MRI slice, where \( \mathcal{V} \) is the set of nodes representing the states \( Y^* = \{0, 1\} \) and \( \mathcal{E} \) is the set of edges of the graph.

Figure 4.3: Realization of proposed radiomics-driven conditional random field framework. For each node \( y_i \), there is a set of corresponding measurements as radiomic features \( x_i \). The connectivity between each pair of nodes \( y_i \) and \( y_j \) is represented by the solid red line \( e_{ij} \).

As shown in Figure 4.3, each node \( y \) represents a single voxel and has a set of associated observations \( x_i \). Each \( x_i \) represents the set of quantitative radiomics features \([2]\) calculated across all MP-MRI modalities for a given voxel. The edge \( e_{ij} \) is the connectivity between a pair of nodes, and represents the relationship between two voxels.

### 4.4 Experimental Setup

The experimental setup used to assess the efficacy of the proposed RD-CRF framework for voxel-resolution prostate cancer detection is described as follows. Using the data previously
described in Chapter 3, each patient dataset had corresponding T2w imaging data, ADC
data, CHB-DWI data, and CDI data. For each modality, the testing data contained a total
of 40,975 samples (40,369 healthy and 606 cancerous samples confirmed by the radiologist)
across the slices from 20 different patients\(^1\). Using the radiologist contour of the prostate,
a rectangle cropped around the prostate gland was selected as the region of interest (ROI)
for each MP-MRI slice.

The voxel-resolution cancer detection results produced by the proposed RD-CRF frame-
work was compared with that produced using the state-of-the-art method proposed in [2]
via sensitivity, specificity, and accuracy metrics (as shown in Chapter 3), where the perfor-
ance of each method was quantified by the metrics’ closeness to one. TP is the number
of voxels in the intersection of the method’s identified cancerous tissue and the voxels con-
tained in the radiologist’s tissue markings, TN is the number of voxels not in the method’s
identified tissue that are also not in the radiologist’s markings, N is the number of voxels
not in the radiologist’s marked tissue, and P is the number of voxels in the radiologist’s
marked tissue.

In addition, the Jaccard index [58] and Sørensen–Dice coefficient [59] were used to
calculate the overlap between the radiologist’s marked tissue and the voxel-resolution cancer
detection results produced by the proposed RD-CRF framework and the state-of-the-art
method proposed in [2].

\[
\text{Jaccard} = \frac{|A \cap B|}{|A \cup B|} \quad \text{Sørensen–Dice} = \frac{2|A \cap B|}{|A| + |B|}
\]

where \(A\) is the set of voxels contained in the method’s identified cancerous tissue and
\(B\) is the set of voxels contained in the radiologist’s tissue markings. The performance
of each method was evaluated using leave-one-patient-out cross-validation, with training
radiomics features were selected from all but one patient and used to train the classifier,
and the voxels in the remaining patient’s MRI dataset were classified as either healthy and
cancerous tissue.

\(^1\)To balance the training data in terms of number of healthy and cancerous samples, when training
the SVM classifier, smaller subsets of healthy samples are randomly selected to be the same size as the
cancerous samples and this is repeated 50 times and the classification results are averaged. This way, the
training data used for training SVM classifier becomes 50% – 50% leveled with respect to cancerous and
healthy samples [2].
4.5 Experimental Results

Table 4.1: RD-CRF performance and Khalvati et al. [2] performance metrics for different numbers of cases. Statistical significance is shown across all performance metrics (p-values below 0.05).

<table>
<thead>
<tr>
<th>Cases/Results</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Accuracy</th>
<th>Jaccard</th>
<th>Sørensen–Dice</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 cases</td>
<td>70.21%</td>
<td>91.38%</td>
<td>90.59%</td>
<td>26.64%</td>
<td>37.12%</td>
</tr>
<tr>
<td>RD-CRF</td>
<td>71.47%</td>
<td>91.93%</td>
<td>91.17%</td>
<td>28.56%</td>
<td>39.13%</td>
</tr>
<tr>
<td>p-value</td>
<td>0.0413</td>
<td>0.000293</td>
<td>0.000163</td>
<td>0.0217</td>
<td>0.0312</td>
</tr>
</tbody>
</table>

Table 4.1 shows the average sensitivity, specificity, accuracy, Jaccard index, and Sørensen–Dice coefficient for the proposed RD-CRF and Khalvati et al. [2] across all patients. The proposed RD-CRF method shows significantly improved (p-values < 0.05) for all metrics.

Table 4.2 shows the improvement of the proposed RD-CRF framework with respect to [2] for different cases starting with 5 worst cases for sensitivity obtained by [2]. As it can be seen from Table 4.2, for the first 5 worst cases, the proposed RD-CRF framework improved sensitivity by 3.2%. When all 20 cases were taken into account, the proposed RD-CRF framework improved sensitivity, specificity, and accuracy by 1.10%, 0.55%, and 0.59%, respectively. Given that [2] already achieved high specificity rates (e.g., 92%), the proposed RD-CRF framework was only able to improve it slightly (0.59%).

Figure 4.4 shows the improvement in sensitivity and specificity obtained by the proposed RD-CRF framework when compared to [2]. Figures 4.5 and 4.6 show the actual sensitivity and specificity numbers for [2] and the proposed RD-CRF framework.

The fact that the proposed RD-CRF framework increased the sensitivity, specificity, and accuracy of [2] indicates that taking into account inter-voxel spatial and radiomics feature

Table 4.2: RD-CRF performance improvement with respect to [2] for different numbers of cases.

<table>
<thead>
<tr>
<th>Cases/Results</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 cases</td>
<td>3.20%</td>
<td>0.52%</td>
<td>0.61%</td>
</tr>
<tr>
<td>10 cases</td>
<td>1.41%</td>
<td>0.55%</td>
<td>0.60%</td>
</tr>
<tr>
<td>15 cases</td>
<td>1.50%</td>
<td>0.48%</td>
<td>0.53%</td>
</tr>
<tr>
<td>20 cases</td>
<td>1.10%</td>
<td>0.55%</td>
<td>0.59%</td>
</tr>
</tbody>
</table>
relationships helps noticeably in mitigating the identification of sparsely distributed tumour candidates and thus improving prostate cancer detection performance. Figure 4.8 shows examples of how the proposed RD-CRF framework mitigates sparsely distributed tumour candidates while promoting consistency in spatial and radiomics feature relationships to ensure the identified tumour candidates better reflect cancerous tumours.

Figure 4.8 shows the prostate tumour candidates detected using [2] and that detected using the proposed RD-CRF framework in comparison to the radiologist’s markings. Figure 4.7 shows sample results for [2], RD-CRF, and the radiologist’s markings along with the corresponding pathology data. The pathology data indicates additional smaller tumours present in the prostate tissue. While [2] and the proposed RD-CRF framework failed to detect these small areas of cancerous tissue, the radiologist’s markings also did not identify these small tumours, which illustrates that such small areas of cancerous tissue may not exhibit significantly distinctive image-based features in the MP-MRI data.

Figure 4.4: RD-CRF performance improvement in sensitivity (shown in red) and specificity (shown in blue) compared to [2]; the graph shows the percent improvements in both sensitivity and specificity for different numbers of cases.
Figure 4.5: RD-CRF sensitivity (shown in blue) for different number of cases compared to [2] (shown in red).

Figure 4.6: RD-CRF specificity (shown in blue) for different number of cases compared to [2] (shown in red).
4.6 Discussion

Computer-aided detection (CAD) algorithms for prostate cancer are mainly based on exploiting the imaging characteristics to train a classifier such as SVM, enabling it to classify new cases. Radiomics attempt to augment this approach by utilizing a large amount of quantitative imaging-based features to improve the classification results. The common theme in these radiomics-driven approaches is the fact that single voxels are studied individually and inter-voxel spatial and feature relationships are usually overlooked. In contrast, we have proposed a radiomics-driven conditional random field (RD-CRF) framework to incorporate the inter-voxel relationships by jointly enforcing multi-parametric radiomics features consistency and spatial consistency with the pairwise term (Equation 4.3). This enforces the compactness and radiomics-based connectedness of the classification results, leading to the better representation of prostate tumours and a better separation of cancerous and healthy tissues.

The proposed RD-CRF framework can incorporate any given quantitative radiomics feature model that classifies image voxels for a given case such as prostate cancer. In prostate cancer, one of the challenges that the CAD algorithms face is the fact that the cancerous voxels are a small fraction of the prostate gland (i.e., less than 1%). This leads to poor sensitivity results in a significant number of cases. To demonstrate the effectiveness of the proposed RD-CRF framework, we calculated the results starting with the worst sensitivity cases obtained by [2]. As it can be seen, the proposed RD-CRF framework was able to improve sensitivity by 3.2% for first 5 cases and 1.10% for all 20 cases while in all...
Figure 4.8: Visual comparison of identified prostate tumour candidates produced by [2] and the proposed RD-CRF framework. RD-CRF produced results that show a noticeable reduction in sparsely distributed tumour candidates and increased spatial consistency in detected tumour candidates. Left to right: radiologist’s markings, results produced by [2], results produced by RD-CRF, and the results overlaid with the radiologist’s markings.
cases specificity was improved by 0.60%. The low gain in specificity is mainly due to high specificity rates obtained by [2] (e.g., 92%) making it difficult for the RD-CRF framework to further improve performance.

The proposed RD-CRF framework can improve CAD algorithms with respect to sensitivity enabling them to become reliable for cancer screening programs where high sensitivity rates are required. Furthermore, [2] also tends to identify more sparsely distributed tumour candidates leading to false positive cases (Figure 4.8). The proposed RD-CRF framework can also help improve specificity of CAD algorithms to be used in procedures such as radical prostactomy where a very high specificity is required to prevent unnecessary surgeries. As seen in Figure 4.8, the proposed RD-CRF framework identified a single compact region (e.g., fifth row) and also extended the identified tumour candidate to an added region (e.g., second row).

While Table 4.2 indicates that the RD-CRF framework generally improved the classification results, RD-CRF is limited by the initial classification results from the trained classifier. Figure 4.9 shows examples of cases where the initial trained classifier failed to identify tumour candidates as cancerous, resulting in incorrect tumour candidate classifications when using the RD-CRF framework.

The example shown in the first and second row of Figure 4.9 are cases where the initial trained classifier failed to identify tumour candidates entirely while the third row shows a case where the initial trained classifier partially identifies the tumour. A visual inspection shows that the texture in the radiologist-marked regions appears similar to other healthy prostate tissue. This suggests that the MP-MRI modalities and corresponding extracted quantitative radiomics features do not capture all underlying texture characteristics of cancerous tissue, thus resulting in the radiomics feature model to not properly identify the suspicious regions. This suggests that more information (via new imaging modalities or the discovery of new quantitative radiomics features) is required to capture such cases.

4.7 Summary

A novel framework was proposed for automatic prostate cancer detection using multiparametric magnetic resonance imaging via a radiomics-driven conditional random field (RD-CRF) framework. Experimental results showed that the proposed RD-CRF framework improved the sensitivity, specificity, and accuracy compared to a state-of-the-art voxel-resolution radiomics-driven prostate cancer detection method. While the overall effectiveness of the RD-CRF framework is limited by the initial classification results, the
Figure 4.9: Examples where tumour candidates are not properly identified. Left to right: radiologist’s markings, initial classification results, and RD-CRF results.

RD-CRF framework was shown to improve the detection of tumour candidates by mitigating sparsely distributed tumour candidates and improving the detected tumour candidates via spatial consistency and radiomics feature relationships. Thus, the RD-CRF framework shows potential for more efficient and accurate computer-aided prostate cancer diagnosis for clinicians.
Similarly to Chapter 3, future work includes incorporating additional MRI modalities, and extending the pairwise potential used to characterize spatial and radiomics feature relationships to better enforce spatial and feature consistency and improve detection performance. The radiomics feature model used here incorporated CDI as one of the imaging modalities which, as shown in [2], boosts the initial results significantly. We have developed an enhanced version of CDI, called dual-stage correlated diffusion imaging (D-CDI), which has shown promise in enhancing separability of cancerous and healthy tissue in prostate MRI compared to CDI [56]. As future work, we will incorporate D-CDI into the radiomics feature model used by the RD-CRF framework to investigate its efficacy in improving prostate cancer detection performance. The introduction of a hierarchical RD-CRF framework will also be investigated for the incorporation of morphological features and asymmetric characteristics of cancerous tissue in the prostate gland. We will investigate using a hybrid morphological-textural feature model as an initial classifier for the RD-CRF framework where in addition to texture analysis, the morphological characteristics (e.g., shape) of candidate regions are taken into account to detect cancer [1].

Furthermore, the DWI data captured at different b-values are usually distorted due to patient movement during the image acquisition, and thus may reduce cancer separability and affect the cancer detection performance of the RD-CRF framework. We have presented preliminary results for co-registering the b-value images to compensate for patient movement [60] which we will incorporate into our proposed RD-CRF framework to investigate the effect on the accuracy of cancer detection. Finally, this method was developed with the intention of using the tumour region candidates defined by statistical textural distinctiveness in Chapter 3 as the initial regions used for classification and enforcing spatial consistency and radiomics feature relationships by RD-CRF.
Chapter 5

Conclusion

In this chapter, a brief summary of the thesis and the key contributions are described in Section 5.1. Recommendations of the methods are discussed in Section 5.2, and potential future work for this research is outlined in Section 5.3.

5.1 Summary of Thesis and Contributions

In this thesis, a comprehensive framework for computer-aided prostate cancer detection using multi-parametric MRI has been developed. The framework consists of two parts: i) a saliency-based method for identifying suspicious regions in multi-parametric MR prostate images based on statistical texture distinctiveness, and ii) automatic prostate tumour candidate detection using a radiomics-driven conditional random field (RD-CRF).

In Chapter 3, a suspicious region identification method via statistical textural distinctiveness-based saliency is described. Using unique texture features extracted from multi-parametric MRI modalities to learn a sparse texture model, the distinctiveness of texture patterns are quantified and used to uncover the underlying saliency via the statistical relationship between texture patterns across different MRI modalities. The quantitative improvement over the state-of-the-art method is reported in Table 3.2, where the statistical textural distinctiveness method achieves an approximately 1.5% increase in sensitivity, and at least a 10% increase in specificity and accuracy. Figure 3.2 shows the qualitative improvement of textural distinctiveness method; the state-of-the-art method has a tendency to be over-sensitive, while the textural distinctiveness method consistently produces spatially compact and useful regions.
In Chapter 4, automatic prostate cancer detection method based on a radiomics-driven conditional random field (RD-CRF) framework is presented. Using the quantitative MP-MRI radiomics feature model proposed in [2], the method incorporates both the spatial relationships between voxels and the quantitative radiomic feature relationships between voxels into a conditional random field model. This facilitates the enforcement of interconnected tissue characteristics reflective of cancerous tumours, thus better representing the actual cancerous tissue phenotype. Table 4.1 shows a comparison of the RD-CRF method with the state-of-the-art method [2] and statistically significant improvement is obtained across all performance metrics (p-values below 0.05), and Table 4.2 shows the performance improvement relative to [2]. Figure 4.8 shows a visual comparison of detected prostate tumour candidates produced by [2] and the RD-CRF method. The proposed RD-CRF method produced results that show a noticeable reduction in sparsely distributed tumour candidates and increased spatial consistency in detected tumour candidates.

5.2 Recommendations

The suspicious region identification method via statistical textural distinctiveness produced higher sensitivity, specificity, and accuracy values than the state-of-art ADC-based method. In addition, statistical textural distinctiveness also identifies suspicious regions on a per patient basis, rather than relying on a fixed ADC value characteristic of typical cancerous tissue (as is the case with the ADC-based threshold method). Thus, statistical textural distinctiveness shows potential for more flexible and visually meaningful identification of suspicious tumour regions, and should be used regularly by medical professionals to aid in suspicious region identification in place of the ADC-based method.

While the RD-CRF method generally improved the initial results, RD-CRF is limited by the initial classification results from the trained classifier. Sample cases where the initial trained classifier failed to identify tumour candidates as cancerous resulted in incorrect tumour candidate detection when using the RD-CRF. Though the overall effectiveness of the RD-CRF framework is limited by the initial classification results, the RD-CRF framework was shown to improve the detection of tumour candidates by mitigating sparsely distributed tumour candidates and improving the detected tumour candidates via spatial consistency and radiomics feature relationships. The RD-CRF framework shows potential for more accurate computer-aided prostate cancer detection for clinicians, and should be paired with a classifier with reasonably accurate initial classifications to further refine and fine-tune the results.
5.3 Future Work

5.3.1 Prostate Cancer Grading

The use of radiomic features is largely prevalent in current computer-aided prostate cancer detection and/or classification research. However, the application of radiomic features can also be extended to cancer grading and staging. Future works will include the use of radiomic features for computer-aided cancer grading and correlation to pathology, so as to better bridge the gap between radiology and pathology. The development of radiomics-based prostate cancer grading methods would enable a non-invasive method for assessing the severity and behaviour of prostate cancer.

5.3.2 Prognosis and Risk Stratification

The potential uses of radiomic features within the field of cancer detection and prognosis are extensive, and can lead to better patient care and survival rates through more reliable risk stratification. Risk stratification refers to a medical professional’s formal estimation of the probability of a person succumbing to a disease or benefiting from a treatment. Future works will also include the application of radiomic features to risk stratification, allowing for the development of a computer-aided decision support framework that helps medical professionals assess patients, and more efficiently and accurately provide prostate cancer risk stratification.

5.3.3 Discovery Radiomics

Current radiomic-driven methods for prostate cancer classification typically employ a set of pre-defined, hand-crafted quantitative features extracted from multi-parametric MR images. While hand-crafted imaging-based feature models for radiomics-driven methods for prostate cancer classification have been shown to be highly effective, the generic nature of such feature models for generating radiomic sequences can limit their ability to fully characterize unique prostate cancer tumour phenotype. Motivated by this, the concept of discovery radiomics [61, 62] was introduced, where we forgo the notion of predefined feature models by discovering customized, tailored radiomic feature models directly from the wealth of medical imaging data already available (as shown in Figure 5.1).

Discovery radiomics has the potential to find previously unexplored, abstract imaging-based features that capture highly unique tumour traits and characteristics beyond what
Figure 5.1: Overview of the proposed discovery radiomics framework where multi-parametric MRI data from past patients is used to discover a customized radiomic sequencer tailored for prostate cancer.

can be captured using predefined feature models, thus allowing for improved personalized medicine and risk stratification through radiomic sequencing.
References


