Lymphatic Filariasis Detection in microscopic images

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Summary

In Africa, the propagation of parasites like the lymphatic filariasis is complicating seriously the efforts of health professionals to cure certain diseases. Although there are medicines capable to treat the lymphatic filariasis, the condition needs to be discovered first, which is not always an easy task having into account that in most countries affected by this disease it can only be detected at night (nocturne). The lymphatic filariasis is, a parasitical infection which can originate changes or ruptures in the lymphatic system as well as an abnormal growth of certain areas of the body causing pain, incapacity and social stigma.

Approximately 1.23 billion people in 58 countries from all over the world are threatened by this disease which requires a preventive treatment to stop its propagation which makes it even more important for the existence of a mechanism that is less costly and more agile in the analysis of a blood smear to verify the existence of microfilariae (little worms that are produced by other adult worms while housed in the lymphatic system).

The lymphatic filariasis is caused by an infection with nematodes (“roundworms”) of the Filariodidea family in which three types are inserted: Wuchereria bancrofti, responsible for 90% of all cases; Brugia malayi, responsible for almost every remaining; B. timori also causing the disease. All three have characteristics that can differentiate them and which allow them to be identified.

The current identification process of the disease consists on the analysis of microfilariae in a blood smear with a blood sample through a microscope and its identification by the observer.

Taking this into account, it is intended to develop image analysis and processing techniques for the recognition and counting of the two principal types of filarial worms from a thick blood smear. Also the use of a smartphone and a portable microscope makes the detection possible without the need of a health professional and consequently can automate the process. To make this possible an adapter smartphone-microscope can be used to obtain an image with the magnification of 1000x. The images can then be analyzed in a server or in the smartphone, if it has enough processing power for it. It is expected from this process that the need to resort to labs to process the blood smear gets unnecessary making the process more accessible and agile instead of costly and slow.

For the detection of the parasites from the acquired images it is intended to implement, experiment and choose the more adequate operations. These comprise pre-processing operations
with the goal to enhance the acquired images and eliminate possible artifacts coming from the acquisition system. However, the main operations should be those that allow the verification of existence or nonexistence, recognition and classification of the pretended parasites. Processing and analysis techniques that are common in these processes are based in the extraction of features (e.g. SIRF, SURF, and FLANN) template similarity, edge detection and description of contours and recognition of statistical patterns.

Once detected and recognized one or more parasites and its types, a rule should be defined and used to declare the presence of the disease and its type.
**Resumo**

Em África, a propagação de parasitas como a filariase linfática está a dificultar seriamente os esforços dos profissionais de saúde para curar determinadas doenças. Apesar de existirem medicamentos capazes de tratar a filariase linfática, esta precisa primeiro, de ser detetada, o que nem sempre é uma tarefa simples tendo em conta que na maioria dos países afetados por esta doença, esta apenas pode ser detetada de noite (noturna). A filariase linfática é, uma infeção parasitária que pode gerar alterações ou ruturas no sistema linfático assim como um crescimento anormal de certas regiões do corpo causando dor, incapacidade e estigma social.

Aproximadamente 1.23 biliões de pessoas em 58 países de todo o mundo são ameaçados por esta doença que requer um tratamento preventivo para parar a sua propagação, o que torna ainda mais importante, a existência de um mecanismo que seja mais acessível e mais ágil na análise de uma amostra de sangue para verificar a existência de microfilárias (pequenos vermes que são produzidos por outros vermes adultos enquanto alojados no sistema linfático).

A filariase linfática é causada por uma infeção por nemátodos (vermes redondos) da família Filariodidea, na qual, se inserem os três tipos: *Wuchereria bancrofti*, responsável por 90% de todos os casos; *Brugia malayi*, responsável por quase todos os restantes; *B.timori* também causadora da doença. Todas as três têm características que permite diferenciá-las e consequentemente identificá-las.

O método de detecção da doença atual, consiste na análise de microfilárias numa amostra de sangue através de um microscópio e a sua identificação por um observador com as qualificações necessárias.

Tendo isto em conta, é pretendido o desenvolvimento de técnicas de análise e processamento de imagem para o reconhecimento e contagem dos dois principais tipos de filárias a partir de uma lamela com uma amostra de sangue. O uso de um smartphone e de um microscópio portátil também tornam a deteção possível sem a necessidade de um profissional de saúde e, consequentemente, o processo pode ser automatizado.

Para tornar isto possível, um adaptador smartphone-microscópio pode ser usado para obter uma imagem com uma magnificação de 1000x. Posteriormente, as imagens poderão ser
analisadas num servidor ou no smartphone, se este tiver capacidade de processamento suficiente
para o propósito. Deste processo, é esperado que a necessidade de recorrer a laboratórios para
processar a amostra de sangue se torne desnecessária, tornado o processo mais acessível e ágil
em vez de custoso e lento.

Para a deteção de parasitas a partir das imagens adquiridas, é pretendida a implementação,
teste e escolha das operações mais adequadas. Estas compreendem operações de pre-
processamento com o objetivo de melhorar as imagens adquiridas e eliminar possíveis artefatos
provenientes do sistema de aquisição. Contudo, as operações principais devem ser aquelas que
permitem a verificação da existência ou não existência, reconhecimento e classificação dos
parasitas pretendidos. Técnicas de análise e processamento que são comuns nestes processos são
baseadas em extração de características (ex. SIRF, SURF, FLANN), semelhança de texturas,
deteção de arestas e descrição de contornos e reconhecimento de padrões estatísticos.

Uma vez detetados e reconhecidos um ou mais parasitas e os seus tipos, uma regra deverá
ser definida e usada para declarar a presença da doença e o seu tipo.
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Rui Pedro Menezes da Rosa Neves
Attitude is a little thing that makes a big difference

Winston Churchill
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<th>Definition</th>
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<tr>
<td>ARM</td>
<td>Advanced RISC Machine</td>
</tr>
<tr>
<td>B</td>
<td><em>Brugia</em></td>
</tr>
<tr>
<td>CN</td>
<td>Caudal nuclei</td>
</tr>
<tr>
<td>CPU</td>
<td>Central Processing Unit</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
</tr>
<tr>
<td>FCA</td>
<td>Filarial circulating antigen</td>
</tr>
<tr>
<td>JNI</td>
<td>Java Native Interface</td>
</tr>
<tr>
<td>kNN</td>
<td>K-Nearest-Neighbor</td>
</tr>
<tr>
<td>(l_{a\beta})</td>
<td>Luminance, a and b color channels</td>
</tr>
<tr>
<td>L1</td>
<td>First larval stage</td>
</tr>
<tr>
<td>L2</td>
<td>Second larval stage</td>
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<td>L3</td>
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<td>LF</td>
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<td>NDK</td>
<td>Native Development Kit</td>
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<tr>
<td>PCR</td>
<td>Polymeral chain reaction</td>
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<tr>
<td>SoC</td>
<td>System on Chip</td>
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<tr>
<td>SVM</td>
<td>Support Vector Machine</td>
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<tr>
<td>W</td>
<td><em>Wuchereria</em></td>
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Chapter 1

Introduction

1.1 Context

The lymphatic Filariasis in humans, commonly known as elephantiasis, it’s one of the 4 most important tropical diseases identified by the World’s Health Organization along with onchocerciasis, Chagas disease and leprosy [1]. It is an infectious disease which causes changes in the lymphatic system and deformation of body parts causing pain and incapacity [2] and that is acquired, normally in childhood, when filarial parasites are transmitted from person to person through mosquito bites.

It is estimated that approximately 120 million people are currently infected in 73 countries across the tropical and subtropical regions of Asia, Africa, Occidental Pacific and part of Caribbean and South America and 1.23 billion live in areas where the Filariasis is capable of transmission being therefore at risk [3].

The economic costs of this disease are huge, estimated over US$1 billion for year only for India [4]. Besides this economic costs there are also social costs in the measure that this disease besides causing pain and discomfort it also stigmatizes socially [5]. In fact, it is estimated that 40 million infected individuals are seriously incapacitated and disfigured by LF [6] and there are studies that suggest that depression is prevalent in individuals with LF.

It is for this set of factors that the LF is considered a public health problem to eliminate until 2020 [8]. Pursuing that goal, in 2000 there was launched the Global Program to Eliminate Lymphatic Filariasis (GPELF) which recommends a strategy of chemoprevention that goes through the massive administration of medicines (MMA) anthelmintic over the populations at risk which implies an investment of at least US$ 154 million per year during the following 5 years [9].

It is, therefore, in this context of massive utilization of medicines where there is no access to a fast and efficient diagnosis, due to the inexistence of health professionals qualified and
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other means that are sophisticated and costly, that comes up the necessity of developing a cheaper diagnosis system of easy execution and interpretation and practicable in areas of limited access to quality health care.

The use of a mobile-based framework seems to accomplish these requirements once that it comprises both a mobile-phone adaptable cheap microscope and an automatic analysis system. The latter consists on automatically assessing the cellular blood components of blood smears as well as the number of filarial parasites, which would replace the time-consuming expertise-dependent standard procedure.

1.2 Motivation and Objectives

The main objectives of this project are summed up below:

- Develop a robust automated image processing methodology capable of finding the parasites and estimate the number of parasites in low-quality smartphone-acquired thick blood smear images;

- Develop a robust image processing methodology to differentiate the two types of microfilariae (*Wuchereria bancrofti* and *Brugia malayi*)

- Integration of image processing methodologies into mobile devices

1.3 Dissertation structure

Besides the introduction this thesis contains more six chapters. In chapter 1 we have the identification and context of the problem, the motivation, and objectives for the solution (the dissertation). In chapter 2 we present the biology of the parasites and describe the actual techniques used to detect these parasites, followed by a review of image processing techniques and similar projects in chapter 3. In chapter 4 the solution developed is presented. Chapter 5 will address the integration of the image processing methodology developed in Android, presenting the mobile application. Following, in chapter 6, the results will be presented and consequently, the discussion associated to those results. Finally, in chapter 7, conclusions and future work are presented.
Chapter 2

Lymphatic Filariasis Characterization

Lymphatic Filariasis is a mosquito-borne disease [10] whose main symptoms are related with the damage of the lymphatic system both in lymphatic ganglia and lymphatic vessels. Approximately 90% of LF is caused by parasite worms of the type *Wuchereria bancrofti*, being the majority of the remaining 10% caused by *Brugia malayi* [11]. On a general way these worms are known as filariae.

2.1 Biology, life cycle and transmission

The filariae *Wuchereria bancrofti* (Cobbold, 1877) and *Brugia malayi* (Bucley & Edeson, 1956) are parasite nematodes belonging to the Filarial superfamily. They have as a preferential habitat the tissues and the circulatory and lymphatic systems of the definitive host where they can live for a period of 8 to 10 years [12]. They are threadlike, white milky colored, opaque, with a smooth cuticle and present sexual dimorphism being the females bigger than males, 80-100 x 0.25mm and 40 x 0.1 mm, respectively. The adults of both types are very identical in all their characteristics being the females undistinguishable. However, the males from *B. malayi* only have around half of the size of the males from *W. bancrofti* [13].

After mating, millions of eggs, existent in the uterus of female worms hatch and give origin to larval states, known as microfilariae which present a cuticle and measure in average 260-300 x 8 μm and have the somatic cores in a simple line where, in the case of *W. bancrofti* don’t reach the caudal end while for the *B. malayi* the two last cores are separated and the last one, very small, is isolated in the caudal apex [14] (Figure 2.1).
The life cycle of both parasites is comprised of two stages given that it needs an intermediate definitive host, the man, to complete its evolutionary cycle [15] (Figure 2.2). In a first stage, when mosquitos bite an infected person they ingest contaminated blood with microfilariae. These loose the cuticle in the stomach of the mosquito and migrate to its torax muscles [16] where they become smaller and thicker originating the first larval state (L1) which after 5 to 7 days becomes the second larval state (L2). Between the 10th and 11th day they suffer a new change and transform themselves in the infecting state (L3) (1500 x 20 μm) which is very active. When mature, these L3 larvae migrate to the mouth of the mosquito to be transmitted to the definitive host when the insect is feeding [17]. The L3 larvae can remain live and active for about 46-50 days or as much time as the live of the mosquito [18].

**Figure 2.1** Differentiation of species of microfilariae on the basis of presence or absence of caudal nuclei (CN)

**Figure 2.2** General Lifecycle of filariae
Lymphatic Filariasis Characterization

The second stage initiates when an infected mosquito bites a definitive host. Once introduced, the L3 larvae migrate to the lymphatic system where they install and develop themselves until adulthood. After about 8 months, in the *W. bancrofti* and 3 months in the *B. malayi*, the pregnant females produce millions of microfilariae that circulate through the blood stream until they get housed in the deep capillaries of several regions of the organism having a lifetime of approximately a year [19].

Similarly to other filariae, *W. bancrofti* and *B. malayi* exhibit periodicity, this is, the concentration of microfilariae in the peripheral blood of the host varies along the day and they can be influenced by cardiac rhythms of the intermediary host [20], namely for the time of biggest biting activity of the vector mosquito [21, 22].

In diverse areas, where the parasites are transmitted by mosquitos with nocturne habits the periodicity of *W. bancrofti* and *B. malayi* is nocturne achieving a peak of concentration around midnight and not being detectable or being in low concentrations around mid-day [23]. There are, however, nocturne sub periodical strains of *W. bancrofti* and *B. malayi* and diurnal sub periodical strains of *W. bancrofti* [15]. In these, the microfilariae are continuously present in the peripheral blood but the concentrations are superior during the night or day, respectively.

The transmission of Filariasis is done exclusively by the bite of mosquitos of types *Aedes, Anopheles, Culex* and *Mansonina* which are the intermediary hosts and vectors of the LF [24, 25, 26].

### 2.2 Filariasis Physiopathology

The ganglia and lymphatic vessels are the places to where the larvae and adults of *W. bancrofti, B. malayi* and *B. timori* go, making them expectable to be the places where the pathology and the pathological process associated are more intense.

In a general way, LF presents big diversity of clinical manifestations such as fever, lymphangitis, chyluria, hydrocele and elephantiasis. These clinical manifestations are the result of the action of both the adult worms, which cause the dilation of the lymphatic vessels and the thickness of its endothelium and the microfilariae which act over the lymphatic ganglia producing extra lymphatic manifestations [27, 28].

The clinic course of lymphatic Filariasis comprises the asymptomatic, acute and chronic stages by progression order.
Lymphatic Filariasis Characterization

2.2.1 Asymptomatic stage

The asymptomatic stage is characterized by the presence of microfilariae on the peripheral blood, although there is no clinical manifestation of the LF. At least half of the patients with LF are clinically asymptomatic [29].

The asymptomatic stage is directly connected to the quality of the immunity response of the patient [30, 31].

2.2.2 Acute stage

The acute clinical manifestations of LF, are characterized by episodic attacks of discomfort, fever and chills, for the appearance of swollen and painful lymphatic ganglia (lymphadenitis), and inflammation of the lymphatic channels (lymphangitis) derived of the block of lymphatic vessels which prevents the free circulation of lymph which has as a consequence the swelling of members or scrotum (lymphedema) [29].

In male individuals the genitalia is frequently affected. The acute attacks can be uni or bilateral and it is common that they occur in individuals with chronic manifestations [32].

2.2.3 Chronic Stage

The chronic stage of Filariasis develops usually between the 10th and 15th years after the occurrence of the first symptoms and it is characterized amongst other for a low density of the microfilariae in the blood which can remain undetected. As the inflammatory process keeps evolving the affected regions becomes harder in the subcutaneous layer due to the high concentration of protein present in the liquid accumulated. An invasion of the subcutaneous tissues is also verified with the consequent loss of elasticity of skin and development of elephantiasis [20] (Figure 2.3).

Figure 2.3 Chronicle clinical manifestations of bancroftian lymphatic filariasis. A – Inferior left member with lymphedema. B – Locker signal (arrows), normally observed after digital compression in the affected member with lymphedema. C – Elephantiasis
The chronic stage is related with the appearance of elephantiasis of members, breast and genitalia, hydrocele and chyluria [33].

4 2.3 Diagnosis

The diagnosis of Filariasis can be done by different parasitological, immunologic or molecular methods and also by imaging techniques.

8 2.3.1 Fresh exam technique

The standard method used in the diagnosis of Filariasis consists on the observation and counting directly on the optical microscope of a blood smear having in attention the periodicity of the parasite [23]. The blood smear sample should be thick and stained with Giemsa or with hematoxylin and eosin.

To increase the sensitivity of the method, concentration techniques can be used like centrifugation and filtration [34].

16 2.3.2 Serologic Techniques

The serologic techniques provide an alternative to the fresh exam technique. The infected individuals have high level of antibodies ant filarial Ig4 on the blood and can be detected using routine exams [35, 36, 37]. Using the same exams they can also search for the filarial circulating antigen (FCA) [38].

22 2.3.3 Polymerase chain reaction (PCR)

The PCR tests are highly specific and sensitive allowing the detection of the presence of parasite DNA in infected individuals as well as intermediary hosts in bancroftian Filariasis and by Brugia [39, 40].
2.3.4 Imaging Technique

Recently the ultrasound allowed locating and visualizing the movements of the adult living worms of *W. bancrofti* and lymphatic dilation in patients with bancroftian Filariasis. This technique allows the diagnosis in individuals classified as asymptomatic with microfilariae [41, 42].

Also the lymphoscintigraphy has shown in individuals classified as asymptomatic the presence of lymphatic abnormalities in affected members of host individuals of microfilaria [31].
Chapter 3

Literature Review

3.1 Introduction

In this chapter, diverse image processing techniques are presented and their contribution for parasite detection and blood cells segmentation. Initially there is a review of the technologies that will be used in the project followed by the review of image processing techniques and stages on similar projects like Detection of Malaria or Blood Cells Segmentation. Lastly different and diverse image processing techniques are presented and explained for the different stages of the processing: image enhancement, image segmentation, feature extraction and classification of features.

3.2 Technology Review

3.2.1 Android

Android is a mobile operating system based on the Linux core and it was developed by Google. It is currently the most used mobile operating system in the world and in 2013 it possessed the biggest percentage in world sales of mobile operating systems. In July 2013, the application store had more than a billion applications available, downloaded more than 50 billion times. A study has shown that 71% of the programmers develop applications for the Android system. The operating system code is made available by Google under an open source license. Android is really popular in technology companies that need a customizable and low cost software for high technology devices. As it is open source it has led to an encouragement of a programmers community on adding resources or bringing Android to devices that initially weren’t launched with the operating system.
3.2.2 Skylight

Skylight is a mechanical device in which its base clamps to a microscope eyepiece. It enables the phone camera to be positioned and held steady over the eyepiece, so that the image from the microscope focuses on the phone camera and consequently displayed on the phone screen. This way, we can obtain an image from a microscope onto our smartphone.

Skylight was designed to be used with desktop microscopes and only a few phones are not compatible with this device.

![Skylight adaptor on the left. Skylight on microscope with mobile device ready to take pictures.](image)

3.2.3 Fraunhofer Microscope Prototype

There is a microscope prototype being developed within the MalariaScope project by Fraunhofer to obtain images from microscope directly to the smartphone. By placing the smartphone on the area designed for it (on top of the microscope) and aligning the smartphone camera with the image sensor we can place a blood smear on the microscope and obtain a set of images.
Literature Review

images that we can use for the image processing stage. A fixed magnification in the image acquisition step is used.

An example of an obtained image will be similar to the one presented in Figure 3.3:

Figure 3.3 Example of a microscopic image containing the parasite

3.2.4 OpenCV

OpenCV is an open source computer vision library originally developed by Intel which is used for the development of applications in the area of Computer Vision. This library was developed in C/C++ programming languages but also gives support to Java, Python and Visual Basic users. There are versions of the library ready for Windows, Linux, Mac, Android and iOS.
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3.2.5 FastCV

The FastCV library offers a mobile optimized computer vision library which includes the most frequently used vision processing functions across a wide array of mobile devices. It contains two implementations, one designed to run efficiently on ARM processors and the second one runs only on Qualcomm SoC’s (mobile processors). The second one provides hardware accelerated implementations of the computer vision functions included in the library.

3.3 Related Work

3.3.1 Segmentation of Blood Cells Components

For image processing there are usually a set of stages that we should follow in order to obtain better and accurate results. Those stages are: image enhancement followed by image segmentation followed by feature extraction and finally, and this one is optional depending on the goal of the method we are constructing, classification of extracted features.

3.3.1.1 Pre-processing

It is imperative to distinguish the blood components because we will need to differentiate them from the parasites and this is not as evident as it seems. The main objective is to recognize, classify and count the diverse types of cells present in the image. For this, it is needed to do a pre-processing of the image in order to make the image more suitable for the following operations. The work described in [43] simply converted the image from RGB to 256 gray levels and used a Gaussian noise reduction filter with the size 3x3 and a standard deviation of 2. Filtering is required to reduce the presence of noise, specially the one produced by low performance cameras. In [44] pre-processing operations like edge detection, spatial smoothing filtering and adaptive histogram equalization to detect and extract the red blood cells from the images were used. The work presented in [45] uses edge detection and highlighting to highlight the edges of the image. It finds the maxima of the gradient of the image to locate edges. This gradient is calculated with the derivative of a Gaussian which will smooth the image and reduces noise and unwanted details and textures.

3.3.1.2 Image Segmentation

The image segmentation goal is to divide the image in interest regions with similar attributes. Again from reference [43] we see the use of Otsu’s method to find the best threshold to divide the image in two classes and the maximum correlation criterion which will evaluate the thresholds using a cost-function. In the end, they used a criterion that uses two thresholds to
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discriminate background, red blood cells and leucocytes. They also use the watershed algorithm to detect and separate two joined cells by computing and plotting as a height the shortest distance between each pixel and the background where the peaks represent the center of the cell. By applying a dilation process having both centers as seed points for the operation they could recover the joined cells. The work described in [45] uses methods based on the thresholding of luminance or color components of the image.

3.3.1.3 Feature Detection and Classification

In feature detection we need to extract the relevant features, in this case the blood cells that will be classified, and we need some processing to find the features that characterize the blood cells.

The authors on [44] used Hough Transform, which was firstly introduced as a method for detecting lines but has been extended to detect low-parametric objects as circles. Each point in the x-y plane yields a line in the a-b plane and if the lines of two points intersect then that is also true for all the points contained in the line. The same principle can be used for the detection of circles by using the circle function instead of a line function. It then counts the number of red blood cells by the following formula:

\[
\text{Actual rbc count per cumm} = \frac{\text{rbc counted by Hough Transform}}{\text{input image area}} \cdot \frac{\text{magnification}^2 \cdot \text{film thickness}}{\text{dilution factor}}
\]  

where film thickness is 0.1mm, the standard medical practice.

Reference [43] searches for a black range pixel, makes a dilation and marks the feature to compute its area. If the area is greater than a defined area threshold then it is a cell. It classifies red cells by auto clustering using the nearest centroid.

The work in [45] uses Artificial Neural Network to find and classify the features. It is structured by layers in which each layer makes independent computations on the data that receives, processes it based on a weighted sum of the inputs and passes to the next layer. A sub group of one or more processing elements with the use of a threshold function determines the output of the network.
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3.3.2 Parasite Detection

For parasite detection, the methods can be similar to those used in the early stages of blood cell segmentation but they differ to find distinctive characteristics of the parasites because these have unique features and need to be distinguished from the blood cells and sometimes they can be mistaken with a blood cell due to similar shapes or in the malaria detection case, we need to analyze the blood cell to check for the parasite inside of it.

3.3.2.1 Pre-processing

The authors of [46] and [47] converted the image to a gray scale space because the image showed some degree of color variability and used a 5x5 median filter to substitute each pixel in the image with the median value of the intensity level in the neighborhood which will result in reduced sharp transitions in the pixel intensities which is the indication of random noise. In reference [48] the authors firstly resize the image for magnification or to reduce image size to speedup processing, reduces noise with also a 5x5 median filter and then enhances image contrast by transforming the original images into the HSI because HSI color space highlights some parasite information.

3.3.2.2 Image Segmentation

In the image segmentation process, the work from [46] segmented the blood cells by converting the pre-processed image into a binary image with a threshold that maximally divides the background from the foreground objects. As a single global threshold was not sufficient they recurred to Otsu’s method to better divide the image into classes.

Reference [48] used histogram thresholding by getting the correspondent HSI image, obtain the green component from RGB image and Hue and Saturation from HSI image and determined the coordinates of bounding rectangles enclosing erythrocytes by using Otsu’s algorithm to segment in green, Hue and Saturation components. It also did the same with other algorithm similar to Otsu’s, named Zack’s algorithm, that locates global minimum points in an image histogram which will be the threshold used to separate objects from background. Thus, two techniques were tested here.

3.3.2.3 Feature Detection and Classification

To extract the features (infected and healthy erythrocytes), the work from [46] used the difference of intensity distribution between possibly infected erythrocytes and healthy ones and the area defined by the erythrocyte. This can be done because the gray-scale infected erythrocytes have intensity values close to 0 (parasite nucleus). With the fact that parasite’s cytoplasm appears lighter while parasite’s nuclei appears darker each possibly infected cell can
be divided into 3 regions using a multiple thresholding to segment them consequently confirming an infected cell or not. Using a Bayes classifier divided in two stages the cells were classified using probabilities. For the first stage infected red blood cells were separated from normal ones through intensity differences. In the second stage a ratio of white to black area was used to detect a false positive or a real infected red blood cell (through difference of intensity from parasite’s nuclei and cytoplasm).

In reference [47] after the gray-scale conversion and median filtering the parasite’s will have a darker color than the rest of the cells so an algorithm was developed to calculate the average of the maximum and minimum pixel intensities and each pixel that had an intensity higher than the average would be part of a possible parasite. Parameters like specificity and sensitivity were used to evaluate the detection.

The work in [48] trained a multilayer back propagation Artificial Neural Network to partition the image in two regions: erythrocytes and background and used two sets of features to train two Artificial Neural Networks (one with RGB pixel values and other with both RGB and HSI pixel values) and the features were divided into two classes: erythrocytes and background pixel values.

### 3.4 Image Processing Techniques

Image processing is the method to perform operations on a digital image with the goal of getting an enhanced image or extract the desired information from it. Image processing involves many processing steps like: image acquisition, image enhancement, image restoration, colour image processing, wavelets and multiresolution processing, compression, morphological processing, segmentation, representation and description, object recognition, tracking and knowledge base.

#### 3.4.1 Image Enhancement

The aim of Image Enhancement is to improve the information in images for human viewers better interpretability or perception and providing better input for other automated processing techniques. This is achieved by modifying attributes of an image to make it more suitable for a given task or specific observer [49]. Therefore, it is important to choose carefully the attributes to modify best suit the needs because the following operations results will depend on the enhancement that has been made.

The enhancements can be divided in two categories: spatial domain methods and frequency domain methods.
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Spatial domain techniques deal directly with the image pixels, applying to each pixel a transformation resulting in a new pixel with the desired enhancement.

\[ g(x, y) = T[f(x, y)] \]  

where \( T \) is the transformation and \( f(x, y) \) the pixel to modify.

For frequency domain techniques the image needs to be transferred first to the frequency domain through the Fourier Transform and all the enhancements are applied to the Fourier Transform of the image and then to get the resultant image it is needed to apply the Inverse Fourier Transform. These are used to enhancement operations such as modifying the image brightness, contrast or the distribution of gray levels, and so the image needs to be converted to gray scale.

The most basic transformation \( T \) is to compute the negative of an image where the pixel grays are inverted. Every pixel value from the original image is subtracted from 255. Negative images can enhance the white or grey detail in the dark regions of the image and can make some details clearer (e.g. in human body tissues) (Figure 3.4).

\[ s = 1.0 - r \]

**Figure 3.4** Negative of an image

Threshold transformations are useful for segmentation where we want to isolate an object from the background converting images into binary images by selecting a threshold that maximally divides the image into two classes of intensities: one for the intensities of background and the other for the intensities of the foreground (Figure 3.5).

**Figure 3.5** Thresholding method
Logarithmic transformations or fractional power-law transformations are used for expanding lower gray pixel values into a wider range, while compressing the high values. The opposite occurs when inverse logarithmic or power-law transformations, with and higher-than-1 exponent are used.

Piecewise linear transformations uses an arbitrary user-defined transform instead of a well-defined mathematical function (Figure 3.6).

![Figure 3.6 Piecewise linear transformation](image)

Another image enhancement technique is the histogram equalization used for adjusting image intensities to enhance contrast. Histogram equalization automatically determines a transformation function seeking to produce an output image with a uniform histogram. Another method is to generate an image having a specified histogram named histogram matching (Figure 3.7).

![Figure 3.7 Histogram Equalization and results](image)

Other technique used is smoothing where the main idea is to suppress either the high frequencies in the image or the low frequencies enhancing or detecting edges in the image. An image can be filtered either in the frequency or in the spatial domain.
The first one involves transforming the image into the frequency domain, multiplying it with the frequency filter function and transform the result into the spatial domain again. The corresponding process in the spatial domain is to convolve the input image with the filter function.

We have to approximate the filter function with a discrete and finite kernel and we shift this kernel over the image and multiply its value with the corresponding pixel values of the image. The size and form of the kernel determine the characteristics of the operation (Figure 3.8).

![Median vs. Gaussian filtering](image)

**Figure 3.8** Comparison between Gaussian and Median filters

Here we can see that Gaussian filter is not optimal to remove salt and pepper noise and the image gets blurred when we apply the filter with higher deviations. Median filter suppresses well the noise as we can see when using the 5x5 kernel.

### 3.4.2 Morphological Image Processing

Morphological image processing describes a range of image processing techniques that deal with the shape of features in an image. Operations of this type are used to simplify image data, preserve essential shape characteristics and eliminate noise [50].

We have the basic of set operations which consists on the following operations:

\[ C = A \cup B \]  
\[(3.3)\]
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2 Produces a set that contains the elements of both A and B.

4 Intersection = AND

6

\[ D = A \cap B = \{ p \mid p \in A \land p \in B \} \]  

(3.4)

8 Produces a set that contains the elements that are in A and B. If they have no common elements the sets are called disjoint.

12 Complement

\[ A_c = A \cup 0 \]  

(3.5)

14 Produces a set that contains the elements that are not contained in A

16

18

20 Reflection

\[ I \ast = \{-p \mid p \in I\} \]  

(3.6)

22 Where p is a point (position) belonging to the original image I.

This operation reflects a point in relation to the origin (0,0).

24 Dilation and Erosion are basic morphological processing operations. They are employed as the basic elements of many algorithms and are produced through the interaction of a structuring set and a set of pixels of interest in the image.

28 A structuring element is a binary image (or mask) that allows us to define arbitrary neighborhood structures and we can think of a binary image as the set of all pixel locations in the foreground or background:
A dilation of an image $A$ by the structuring element $B$ is given by the set operation:

$$A \oplus B = \{ s | (\hat{B})_{s} \cap A \}$$  \hspace{1cm} (3.8)

where $\hat{B}$ is the reflection of $B$ about its origin followed by a shift $s$.

A dilation is used for repairing breaks and intrusions and can also be computed as an erosion of the foreground and an erosion can be computed as a dilation of the background. Thus, we represent erosion with the set operation:

$$A \ominus B = \{ s | (\hat{B})_{s} \subseteq A \}$$  \hspace{1cm} (3.9)

Erosion can split apart joined objects and can strip away protrusions. It is used for shrinking or thinning operations whereas dilation grows and thickens the objects in a binary image. We can make joint operations with erosion and dilation that can be useful for smoothing the contour of an object, eliminate thin protrusions, eliminate small holes and fill gaps in the contour.

Opening operation is an erosion followed by a dilation:

$$A \circ B = (A \ominus B) \oplus B$$  \hspace{1cm} (3.10)

where stray foreground structures that are smaller than the $B$ structuring element will disappear and larger structures will remain in the image.

Closing operation is a dilation followed by an erosion:

$$A \cdot B = (A \oplus B) \ominus B$$  \hspace{1cm} (3.11)

and it consists of filling the holes in the foreground that are smaller than $B$.

If we want to get the outline of the object we can apply an outline operation that consists of a dilation followed by a subtraction and if we wish to skeletonize the object then we just need to repeatedly run erosion and stop when the thickness is at 1 pixel.
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We can also apply morphological operations in gray scale images. For this we need to define erosion and dilation as the maximum or minimum operation, respectively, in the neighborhood covered by the structuring element B (Figure 3.9 and 3.10):

\[
(A \oplus B)(u, v) = \max_{(i,j) \in H} \{A(u + i, v + j) + B(i, j)\} \tag{3.12}
\]

\[
(A \ominus B)(u, v) = \min_{(i,j) \in H} \{A(u + i, v + j) + B(i, j)\} \tag{3.13}
\]

![Figure 3.9 Closing operation](image1)

![Figure 3.10 Opening Operation](image2)

3.4.3 Image Segmentation

Good image segmentation is characterized by its efficiency, meaning the process speed, shape matching and better shape connectivity with segmentation result. The goal is to identify and isolate the regions corresponding to individual surfaces, objects or natural parts of objects. It can be used for image compression, object recognition, image editing and others. We have several segmentation algorithms and these can be divided in two basic properties: discontinuity and similarity. In the first one, partition is based on the changes in intensity levels or grey levels of an image and the focus is to identify points, lines and edges. In similarity pixels that are similar, for example, in brightness levels, are grouped and to do this we have approaches like thresholding, region growing and region splitting and merging.

Based on this we can classify segmentation techniques in four categories [51]:

- Segmentation by Edge Detection
The segmentation by Edge Detection divides the image into object and its background. It observes the changes of intensity in the pixels of the image where if the intensity values do not change much along a direction it is an edge. There are first derivative and second derivative edge detection operators but the second order ones give the most reliable results. An example of a second derivative edge detector is the Canny edge detector presented in Figure 3.1:

![Original Image](image1.png) ![Edge Image](image2.png)

**Figure 3.11 Edge Detection**

First the image is converted to gray scale and a Gaussian filter is applied for filtering out any noise that may exist on the image. After smoothing the image it finds the edge strength by taking the gradient of the image and estimating the gradient in the x-direction columns and y-direction rows to find the absolute gradient magnitude. When edge strength is found, it finds the direction through the gradient of x and y directions and nonmaximum suppression is applied which means that any pixel value along the direction found that is not considered to be an edge is set to 0 (black).

Next, the segmentation by thresholding is one of the simplest approaches to segment an image. This is done by looking at the intensity levels of the pixels. We can define global or local thresholding.

Global thresholding distinguishes the objects from the background by comparing with the threshold defined and uses binary partition to segment the image.

Local thresholding, as it says, is local, meaning that the threshold values vary over the image depending on the characteristics found at that region. A histogram is used to segment the image afterwards.

Otsu’s method [52] is an automatic thresholding method and it is automatic because it chooses the thresholding value based on the foreground and background variances (measure of
spread) of each threshold. It iterates through all possible thresholding values and by checking the pixels that fall in foreground or background it calculates a weighted variance based on that, measuring the separability of two classes. The goal is to find the threshold value where the sum of foreground and background spreads is at its minimum. To calculate the class-variance the following formula is used:

\[ \sigma_w^2 = w_b(t)\sigma_b^2(t) + w_f(t)\sigma_f^2(t) \] (3.14)

The segmentation by region groups the pixels that are related to some object. The area that is detected for segmentation should be closed. A single pixel is taken at first and the region grows by comparing all the unallocated neighboring pixels through the difference of the pixel’s intensity value and region’s mean intensity value. The one with the smallest difference gets allocated to the respective region.

Finally, the segmentation by clustering groups the pixels based on its attributes. A cluster contains a group of similar pixels that belong to a specific region and are different from the other regions. Images can also be grouped according to their content. In this case, grouping is made according to some inherited characteristics of the pixels like shape, texture, etc. A well-known clustering technique is the K-Means technique.

Choosing an initial number of centroids (K) the data points are assigned to the nearest centroid minimizing the square of the distance from the data points to the cluster.

\[ J = \sum_{j=1}^{k} \sum_{i=1}^{n} \left| x_i^{(j)} - c_j \right|^2 \] (3.15)

### 3.5 Machine Learning

Machine learning is a core subarea of artificial intelligence. The goal is to create learning algorithms that do the learning automatically without the need of human intervention or any type of assistance. The learning is always based on some sort of observations or data, direct experience or instruction. It learns from the past to behave better in the future.

#### 3.5.1 Feature Extraction

A feature is a part of an image that contains interesting details. It is a general term since to define a feature we need to know what is the goal of the image processing or which kind of
structures are we identifying hence which characteristics do they have (features). Some of the most commonly used features are: edges, corners, blobs, etc. Gabor filters, shape based, texture based, wavelet transforms, SIFT, SURF, Fourier descriptors, Haar transforms are amongst the several techniques for feature extraction.

3.5.1.1 SIFT (Scale-Invariant Feature Transform)

A method is needed to be able to locate objects in an image containing other objects. This method needs to be scale, rotation, and luminance invariant because we need to locate an object in an image and this image will not always have the same magnification, color intensities or rotation. SIFT provides a set of features of an object that are not affected by any of those complications. This method is also very resilient to the effects of “noise” in the image [53].

Stage 1: Scale Space Extrema Detection

To achieve scale invariance, SIFT looks for stable features across all possible scales using a continuous function of scaling, constructing the scale space. It uses the Difference of Gaussian filter for scale space because it is an efficient filter and has the same stability as the scale-normalized Laplacian of Gaussian (used in detection of edges, blobs).

The Difference of Gaussian algorithm is a gray scale image enhancement algorithm that subtracts a blurred version of the original gray scale image from another blurred version of the original image. These images are obtained by convolving the original gray scale image with Gaussian kernels with different deviations, producing different blurs. This blur will suppress only high frequency spatial information and by subtracting one image from the other we will preserve the spatial information that lies between the ranges of frequencies that are preserved in the two blurred images.

Stage 2: Keypoint Localization

This step goal is to find keypoints in the 3D scale space.

Some pixel is selected if it is larger or smaller than all its 26 neighbors, 8 on the same scale and 9 on the scale above and below.

Here, points with low contrast and poorly localized along edge data are discarded.

This is done by calculating the Laplacian value for each key point found in stage 1. The location of extremum, z is given by: 
If the function value at $z$ is below a threshold value the point is excluded (extrema with low contrast). To eliminate key points with poor localization it is noted that in these cases there is a large principle curvature across the edge but a small curvature in the perpendicular direction in the difference of Gaussian function. If this difference is below the ratio of largest to smallest eigenvector, from the 2x2 Hessian matrix at the location and scale of the keypoint, the keypoint is rejected.

**Stage 3: Orientation Assignment**

By assigning a consistent orientation, the keypoint descriptor can be orientation invariant. Assign dominant orientation as the orientation of the keypoint.

First, the Gaussian-smoothed image $L(x, y, \sigma)$ at the keypoint’s scale sigma is taken so that all computations are performed in a scale-invariant manner. For an image sample $L(x, y)$ at scale sigma, the gradient magnitude, $m(x, y)$, and orientation, $\theta(x, y)$, are precomputed using pixel differences:

- Form an orientation histogram from gradient orientations of sample points.
- Locate the highest peak in the histogram. Use this peak and any other local peak within 80% of the height of this peak to create a keypoint with that orientation. Some points might have more than one orientation.

**Stage 4: Keypoint Descriptor**

The local gradient data used in the stage above, is also used to create keypoint descriptors. The gradient information is rotated to line up with the orientation of the keypoint and then weighted by a Gaussian with variance of 1.5 times the keypoint scale. This data is then used to create a set of histograms over a window centered on the keypoint.

Keypoint descriptors typically uses a set of 16 histograms, aligned in a 4x4 grid (Figure 3.12), each with 8 orientation bins, one for each of the main compass directions and one for each of the mid-points of these directions. These results in a feature vector containing 128 elements.
These resulting vectors are known as SIFT keys and are used in a nearest-neighbors approach to identify possible objects in an image [52].

### 3.5.1.2 SURF (Speed UP Robust Features)

SURF is inspired by SIFT and the standard version of SURF is several times faster than SIFT. SURF uses an integer approximation of the determinant of the Hessian matrix for blob detection, which can be computed with three integer operations using a pre-computed integral image and uses Haar wavelet for feature descriptor that can also be computed using an integral image [54].

#### 3.5.1.2.1 Interest Point Detection

Integral images are used. Integral images allow for fast computation of box type convolution filters. The entry of an integral image $I_\Sigma(x)$ at a location $x = (x,y)^T$ represents the sum of all pixels in the input image $I$ within a rectangular region formed by the origin and $x$.

\[
I_{\Sigma}(x) = \sum_{i=0}^{i=a} \sum_{j=0}^{j=b} I(i,j)
\]  

The detector is based on the Hessian matrix and it detects blob-like structures at locations where the determinant is maximum. The determinant is also used for scale selection. Given a point $x = (x,y)$ in an image $I$, the Hessian Matrix $H(x, \sigma)$ in $x$ at scale sigma is defined as:

\[
H(x, \sigma) = \begin{bmatrix} L_{xx}(x,\sigma) & L_{xy}(x,\sigma) \\ L_{xy}(x,\sigma) & L_{yy}(x,\sigma) \end{bmatrix}
\]  

Where $L_{xx}(x,\sigma)$ is the convolution of the Gaussian second order derivative $\partial^2 / \partial x^2 * g(\sigma)$ with the image $I$ in point $x$. As we had seen in SIFT Gaussians are optimal for scale-space analysis.
The approximation of the Hessian Matrix to second order Gaussian derivatives goes further with box filters. 9x9 box filters represent the lowest scale (highest spatial resolution) for computing the blob response maps (Figure 3.13).

Interest points need to be found at different scales so these are usually implemented as an image pyramid. The images are repeatedly smoothed with a Gaussian and then sub-sampled in order to achieve a higher level of the pyramid. Scale space is analyzed by up-scaling the filter size rather than iteratively reducing the size of the image. The scale space is divided into octaves with one octave representing a series of filter response maps obtained by convolving the same input image with a filter of increasing size.

In order to localize interest points in the image and over scales, a non-maximum suppression in a 3x3x3 neighborhood is applied. The maxima of the determinant of the Hessian Matrix are then interpolated in scale and image space.

3.5.1.2.2 Interest Point Description and Matching

To be invariant to image rotation, a reproducible orientation for the interest points is found. First the Haar wavelet responses are calculated in x and y directions and weighted with a Gaussian. The dominant orientation is estimated by calculating the sum of all responses within a sliding orientation window of size $\pi/3$. The horizontal and vertical responses within the window are summed and these two summed responses yield a local orientation vector where the longest one will define the orientation of the interest point.

For the extraction of the descriptor a squared region centered around the interest point and oriented along the orientation selected is constructed. The region is split up regularly into smaller 4x4 square sub-regions which preserve important spatial information. For each sub-region Haar wavelet responses at 5x5 regularly spaced sample points are computed. The responses in horizontal and vertical directions are summed up over each sub-region and form a first set of entries in the feature vector. Each sub-region has a four-dimensional descriptor vector for its underlying intensity structure which is done by extracting the absolute values of the Haar wavelet responses. Concatenating the four-dimensional descriptor vector for each of all 4x4 sub-regions results in a descriptor vector of length 64. By turning the descriptor into a unit
vector, contrast invariance is achieved. Wavelet responses are invariant to an offset in illumination.

For fast index during the matching stage, sign of the Laplacian for the underlying interest point is included. It gives a positive response for dark blobs on bright backgrounds and a negative response for bright blobs on dark backgrounds. In the matching stage features are compared to check if they have the same type of contrast which allows for faster matching (Figure 3.14).

![Figure 3.14 Matching](image)

### 3.5.2 Classification Methods

Classification is the assignment of an object to a certain class based on its similarity to previous examples of other objects.

#### 3.5.2.1 Support Vector Machines (SVMs)

Support Vector Machines is a function-based learning algorithm which aims to compute the maximum margin hyperplane separating two classes. SVM may not be able to find any separating hyperplane if the data contains misclassified instances and this can be fixed by using a soft margin that accepts some misclassifications.

Most real-world problems involve non-separable data for which no hyperplane exists that separates the positive from negative instances in the training set. A solution to this problem is to map the data onto a higher dimensional space and define a separating hyperplane there. Kernel trick can be used, to allow inner products to be calculated directly in feature space without the need to map the data into high dimensional space.

#### 3.5.2.2 Bayesian Classifiers

Bayesian Classifiers minimize the probability of misclassification. The idea behind a Bayesian Classifier is that if an agent knows the class it can predict values of the other features. If not, Bayes’ rule can be used to predict the class given some feature values. The simplest case of a Bayesian Classifier is the naïve Bayesian classifier.
3.5.2.3 Decision Tree

Decision Trees are trees that classify instances by sorting them based on feature values. Each node in a decision tree represents a feature in an instance to be classified, and each branch represents a value that the node can assume. Instances are classified starting at the root node and sorted based on their feature’s values.

The feature that best divides the training data will be associated to the root node. The source set gets split into sub-sets based on attribute values and this is made recursively for each of the sub-sets until there is no more value on splitting again. The most well-known algorithm for building decision-trees is the C4.5 and the most commonly selected attribute value test is information gain which is based on the concept of entropy from information theory. The information gain is the entropy reduction of a particular features achieved by learning the classification.

3.5.2.4 k-Nearest Neighbor (kNN)

k-Nearest Neighbor is a lazy learning algorithm and its principle is that the instances within a dataset will generally exist in close proximity to other instances that have similar properties. If the instances are tagged with a classification label, then the value of the label of an unclassified instance can be determined by observing the classes of its nearest neighbors. It located the k nearest instances and determines its class by identifying the most common class on the k nearest neighbors. The relative distance between the instances is determined by using a distance metric (e.g. Euclidean distance, Manhattan distance).

3.6 Relevant Remarks

In relation to blood cells segmentation the work on [44] diluted samples with an anticoagulant liquid to separate the cells and prevent their overlapping. The counting method presented here had good results but blood cells that did not appear completely on the image weren’t counted, like it would be done in a manual blood test, so improvements need to be made in order to count those cells that appear cut. In [43] the authors they were able to discriminate between red and white blood cells with their proposed framework. The software developed could give the precise result of a blood test if appropriate calibration of the microscope was used. The classification method based on nearest centroid performed well having a total recognition rate of 98%. The work described in reference [45] used the layered feed-forward Artificial Neural Network and was able to detect between complete and healthy, incomplete non circular and sickled red blood cells. It had an accuracy of 81% counting red and white blood
cells and this accuracy can be improved by increasing the training of the Artificial Neural Network.

For parasite detection, the work in reference [46] proposed method presents reasonable results but in the results possible infected erythrocytes with low white/black ratio were identified as infected which should only happen with a higher ratio level. When the erythrocyte is in the very early trophozoite stage the cytoplasm is not very visible so the algorithm failed in identifying those erythrocytes for some cases. A future work is to classify the parasite based on shape and size. Reference [47] results had some false positives due to impurities or white blood cells having the same color as the malaria parasite but overall it had a good performance on identifying the infected cells and the malaria parasite although accuracy needs to be improved.

In reference [48] the accuracy of the method was 95% in detecting the presence of plasmodium parasites which proved that Artificial Neural Network classifiers are reliable and accurate. Artificial Neural Network based only on RGB colors performed better than the one with RGB and HSI features. A technique for segmenting plasmodium parasites based on Zack’s algorithm was developed which determines the best threshold value.

For image pre-processing, median filter could be a good option since almost all of the proposed methods used it and stated it performed better than the Gaussian filter. In every method images were converted to gray-scale except on the work from [48] which maintained color and converted only to HSI in some cases. For image segmentation most of the methods used thresholding (histogram, automatic, luminance), applying different algorithms to separate blood cells from parasites or to segment blood cells only.
Chapter 4

Lymphatic Filariasis Detection

As said before, lymphatic Filariasis is a parasitic infection caused by nematodes of the type *Wuchereria* and *Brugia*. *Wuchereria bancrofti* is responsible for 90% of the cases, *Brugia malayi* is responsible for almost all of the remaining cases and *B. timori* falls out of the scope of the project because it causes almost none of the cases. The unique features that characterize the two main species (section 2.1) provide the basis for the actual method of Lymphatic Filariasis detection which is the observation by microscope of a blood smear stained with Giemsa and the manual identification of the parasites with special attention to their periodicity. The periodicity depends on the type of parasite where there are parasites that have a maximum concentration during the day and others during the night. That is the case of the two microfilariae that we need to identify, which means that the blood collection needs to be done during the night. The method is not an agile method and it always requires an health professional to perform the analysis of the blood smear which difficulties the task for regions like Africa where it is impossible to analyze so many cases in so little time and where it is hard to have many health professionals and acquire medical material to perform the analysis.

Given this, there is a need for a reliable detection method that is more agile and accessible to detect the disease which is the goal of this project.

By acquiring microscopic images from a portable microscope into a smartphone, the smartphone will be able to process those images using image processing techniques and analyze the blood smears to verify whether there are parasites or not. As it is an autonomous system, it doesn’t need any human intervention besides the image acquisition which makes a possible solution for regions like Africa where a person that is not an health professional and doesn’t have that much knowledge about the disease can still conduct the test on the smartphone that will tell if the blood smear on analysis has infected parasites or not. As it will identify the two types of parasites on the blood smear, this solution makes possible for the identification of
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parasite as well as its recognition, differentiating the *Wuchereria bancrofti* of the *Brugia malayi*, allowing the health professionals to medicate based on it.

### 4.1 The Process Model

The goal of this project was to detect microfilariae in microscopic images of a Giemsa stained blood smear and to classify the detected microfilariae into one of the two most common types of microfilariae that cause Lymphatic Filariasis (*Brugia malayi* and *Wuchereria bancrofti*). The process for this type of problems is, generally, very similar and as it follows.

It starts with the image acquisition phase where images are acquired to be analyzed and is followed by a pre-processing phase, where certain operations are applied to the image in order to reduce noise and enhance certain aspects that are relevant for the goals of the project, in this case, enhancing the microfilariae and its characteristics. Is continued by the segmentation phase where the interest regions of the image are isolated (microfilariae), the feature extraction phase where we extract features from the segmentation developed, in order to be able to use machine learning to train and classify those features extracted, making this the last phase. All of these steps are outlined in figure 4.1.

![Figure 4.1 Different stages of the methodology](image)

### 4.2 Image Acquisition

The first phase of the process was to acquire images from the microscope(s) to build our database of images. The samples were obtained through the Institute of Tropical Diseases and were exclusively of thick blood. Two samples were given and used but only images that contained one parasite were included in the database construction. There are some requirements for the image to enter the database as we can see in the following section.
4.2.1 Image Dataset Requirements

- The thick blood smears must be collected by a specialized doctor;
- The thick blood smears should be obtained from two different patients;
- The images should be acquired with a 1000x magnification;
- The images should be acquired with a low-cost commercial microscope or with a microscope-smartphone adapter using a high resolution camera from a smartphone device;
- Every image must contain, at least, one parasite, except for the ones used for testing the segmentation which contain only cells;
- The database should have approximately the same number of images from each sample;
- The images should be annotated by a specialist;

4.2.2 Image Dataset Construction

The images were taken from a normal microscope using the Skylight adapter and from Fraunhofer’s prototype. Only images with one parasite were added to the dataset, except for some testing images that contained only cells, to test the segmentation results when there is no parasite present. Due to scarcity of parasites in the samples, the number of different parasites found on each sample is of 9 and 5 for *Brugia malayi* and *Wuchereria bancrofti* samples, respectively. To address this problem, several images of the same parasite were taken, in different positions on the optical circle, obtaining images with different light and background conditions and with the sample upside down to simulate new parasites. Together, the dataset is composed of a total of 109 images, 55 from the first sample and 54 from the second one.

4.3 Pre-Processing

In Pre-Processing the aim is to reduce the noise of the images and to enhance the details of the structures that are relevant for the goals, to make it suitable for the next phase, which is segmentation. The lesser noise in the image and the more visible and distinctive the relevant structures are, the clearer the segmentation will be. In this phase, there will be three main steps. The first one is to resize the image by cropping it, suppressing the size of it to the size of the diameter of the optical circle found in the image. The second one is to reduce noise, performing a color normalization, channel subtraction and applying a median filter. This color normalization will make it possible to have the images from *Brugia malayi* and *Wuchereria*...
bancrofti samples in the same range of colors, as they have different colorations. By subtracting the blue channel from the green channel, from the RGB color space, the result will be a grayscale image. Applying a median filter will reduce some of the background noise of the stains. The final step will be to enhance the structure, in this case, the filaria, by applying a morphological operation to enhance the contrast between the filaria and the cells present in the image. This morphological operation is based on the calculation of the image histogram in grayscale.

4.3.1 Image Cropping/Resize

The resizing of the image is performed to remove the extra black pixels we get from the photography taken because these will cause extra processing and may alter the results. By applying this cropping to the image, we will get an image with the size of the diameter of the optical circle that is present in the picture. To do so, it is needed to threshold the image to separate the background from the ROI (region of interest). The ROI contains the sample itself and all the elements that compose it. By applying successive median filters, which will blur the image making the elements blend, it becomes easier to apply a threshold based on the Otsu’s method to separate the image into two regions, the background and the foreground. The size of the median filter is estimated as shown below:

\[
window = \begin{cases} 
\frac{imgWidth}{windowFactor} & \text{if } imgWidth > imgHeight \\
\frac{imgHeight}{windowFactor} & \text{if } imgHeight > imgWidth 
\end{cases} \tag{4.1}
\]

where \( windowFactor \) is a constant with a value of 40.

Figure 4.2 Original image on the left. Segmented optical circle in the middle. After cropping with diameter size on the right. Parasite of type *Wuchereria bancrofti*
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The result is the optical circle pixels in white color and the rest of the pixels in black. Afterwards, the diameter is calculated using the first white pixel found on the top of the image and the last white pixel found on the bottom of the image. With the diameter, the image is resized both in height and width to the value of the diameter found.

4.3.2 Color Normalization

The color normalization process, as it was mentioned above, will make it possible to have images from both samples in the same range of colors, dealing with inconsistencies [55], making it possible to have a single processing algorithm for both the samples. To perform this color normalization, Reinhard’s method was used. This method matches the color distribution of the input image with the color distribution from the target image by use of a linear transform in a perceptual color space [56]. The color space chosen is the \( l\beta \) color space which has recently been applied in digital histology images [56]. To perform this normalization both the input and target images are converted to the color space, the mean and standard deviations of both images are calculated and then the means and standard deviations of each color channel in the two images in the color space are matched according to the equations below:

\[
l_{\text{normalized}} = \frac{l_{\text{original}} - \bar{l}_{\text{original}}}{\tilde{l}_{\text{original}}} l_{\text{target}} + \bar{l}_{\text{target}} \tag{4.2}
\]

\[
\alpha_{\text{normalized}} = \frac{\alpha_{\text{original}} - \bar{\alpha}_{\text{original}}}{\tilde{\alpha}_{\text{original}}} \alpha_{\text{target}} + \bar{\alpha}_{\text{target}} \tag{4.3}
\]

\[
\beta_{\text{normalized}} = \frac{\beta_{\text{original}} - \bar{\beta}_{\text{original}}}{\tilde{\beta}_{\text{original}}} \beta_{\text{target}} + \bar{\beta}_{\text{target}} \tag{4.4}
\]
In the end, the image is converted back to the *RGB* color space, obtaining the final result.

The final result, will have the filaria in a lighter color, lighter than most of the cells, and of a very distinctive color from the color of the background. This will allow that the following operation has a more effective result. The target image chosen for the process was a well illuminated Giemsa stained image of a microfilaria with a 1000x magnification and the input image is the negative of the original image since it differentiates the background color from the color of the cells and filaria which will have a similar color.

![Figure 4.4](image1.png)

*Figure 4.4* Original image on the left. Negative of the original image in the middle. Result of color normalization applied to the negative image and after cropping on the right. Parasite of type *Wuchereria bancrofti*

![Figure 4.5](image2.png)

*Figure 4.5* Original image on the left. Negative of the original image in the middle. Result of color normalization applied to the negative image and after cropping on the right. Parasite of type *Brugia malayi*

### 4.3.3 Channel Subtraction and Median Filter

After the color normalization, the *Blue* channel is subtracted from the *Red* channel and a median filter is applied based on the diameter of the image. The *Blue* and *Red* channels were the ones that produced better results and no other color space had better results than the *RGB* color space after color normalization. Either one of the samples would have great results and the
other very bad ones, or both samples had fair results but some images would fail because it had a more loaded background.

In relation to the median filter, it is applied to larger images with a size of 9, for medium images a size 5 will suffice, and for the smaller images the size has to be 3. If a median filter is applied with a bigger size than the one that is supposed to be applied, the thinner parts of the filaria will become disconnected from the rest of the filaria. With the proposed sizes, the noise reduction is optimal being able to detach some of the background from the filaria.

![Figure 4.6 Images from both samples after channel subtraction and median filter. Parasite of type Wuchereria bancrofti on the left and Brugia malayi on the right](image)

**4.3.4 Morphological Operation to reduce background**

The morphological operation used after performing the median filter is the Top Hat. The Top Hat operation is the difference between the original image and its opening (morphological operation applied on the original image). With the adequate size, we can eliminate most of the background that usually is of a similar color as the color of the filaria. To find the samples that have a more loaded background, the histogram is calculated for each image as presented in equation below. The histogram is, for each color present in the image, the number of pixels found in the image.

The probability assigned to each gray level can be given by the relation:
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\[ p_r(r_k) = \frac{n_k}{N} \quad 0 \leq r_k \leq 1, \quad k = 0,1,2,L - 1 \] (4.5)

Where \( r_k \) is the normalized intensity value, \( L \) is the number of gray levels in the image, \( n_k \) is the number of total pixels with the gray level \( r_k \) and \( N \) is the total number of pixels.

After testing several images, a pattern was found. Images with a loaded background had high numbers of pixels between the intensity values 32 and 45, which are the intensity values where most of the background is located. The threshold found for deciding which size should be applied for the structuring element was of 300000 pixels with the intensity values between 32 and 45. Images with a less loaded background had a lower number of pixels between this two values while images with a more loaded background had more than 300000 pixels, and usually, in the order of millions. Images with lower values had less background which means that the structuring element can be bigger because it does not need to eliminate much of the background while images with high levels of pixels between 32 and 45 have a more loaded background which means that it needs to be reduced and removed. To find the size of the structuring element several tests were conducted on different sized images and the relation found through the diameter was the following:

\[
\text{elementSize} = \begin{cases} 
2 \times (0.0367 \times \text{diameter} - 4.5) + 1 & \text{if } \sum_{i=32}^{i=45} \text{hist}[i] < 300000 \\
2 \times (0.0256 \times \text{diameter} - 1.22) + 1 & \text{if } \sum_{i=45}^{i=32} \text{hist}[i] > 300000 
\end{cases} \] (4.6)

The structuring element is the pattern that will be searched for in the image and where the operation is applied. For the structuring element, a rectangular shape was used, meaning that, the element to look for, are rectangles with the intended size. The implementation library also supports ellipse shaped elements that produced more accurate results, with the boundaries of the filariae smoother, but as this is a very costly operation, and the results are not of a significant difference in relation to rectangular shapes, those were the ones chosen, which makes the processing much faster and still produces very good results.
After applying this morphological operation, pixels with more intensity are the ones relevant and the morphological operation will allow that the background pixels intensity from both samples is at about 32 or lower for what the cells and the filaria can be found between 32 and 255 values of intensity.
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Most of the filaria is located at higher intensity pixels, but there are certain parts, especially thinner parts of the filaria that we don't want to lose, that have a lower intensity. With this in mind, a simple threshold is applied to cut the pixels that are below 32 of intensity, which will clear some parts of the image that may contain some background and isolating more the filaria to produce better results in the segmentation phase.

![Image after morphological operation on the left. Results of pixel threshold by 32 on the right. Type: *Wuchereria bancrofti*](image)

**Figure 4.9** Image after morphological operation on the left. Results of pixel threshold by 32 on the right. Type: *Wuchereria bancrofti*

![Image after morphological operation on the left. Results of pixel threshold by 32 on the right. Type: *Brugia malayi*](image)

**Figure 4.10** Image after morphological operation on the left. Results of pixel threshold by 32 on the right. Type: *Brugia malayi*


4.4 Segmentation

The segmentation phase is where the regions of interest are isolated to be able to extract features after. For this work, the regions of interest are the filariae, meaning that in the segmentation phase, the aim is to remove everything that is not a filaria, or of importance to maintain the structure of the filaria. Here, most of the cells that are within the sample will be removed as well as the remaining background from the Pre-Processing. The segmentation process has three phases. The first is to threshold the grayscale resulting image from pre-processing, separating it in two different intensity values, 0 and 1, where 0 is black and 1 is white, making it binary. The pixels with the highest intensity value will be the pixels that belong to the interest regions, filaria, cells and background. Afterwards, another threshold is made, this time an area thresholds to remove all the objects from the image that are smaller than a certain size. This will remove the cells that are not attached to the filaria and the background. The latest step, will be to remove the cells that are attached to the filaria because as they have a very similar color to the filaria, they will be thresholded together with the filaria, making them one single component.

4.4.1 Adaptive Threshold

To make the image binary, adaptive threshold was used. Adaptive threshold calculates a threshold for small regions (mean of the block size of the neighborhood) instead of defining a single threshold for the whole image. To calculate this threshold the adaptive Gaussian mean function is being used.

\[
dst(x,y) = \begin{cases} 
    \text{maxValue} & \text{if } src(x,y) > T(x,y) \\
    0 & \text{otherwise} 
\end{cases} \quad (4.7)
\]

\[
T(x,y) = \text{mean}_{x,y}(\text{blockSize} * \text{blockSize}) - C \quad (4.8)
\]

As there are varying lighting conditions on the samples, adaptive threshold produces a much better result, whereas in the threshold performed through the Otsu’s Method, a single threshold was found and the remaining of the background was segmented along with the filaria, causing it to have background attached which would be difficult to remove afterwards.
To find the best values for the adaptive threshold function, tests with all the images were conducted at different sizes. A formula was found to relate the window size through the diameter of the image:

\[
\text{windowSize} = \text{roundUP}(0.1364 \times \text{diameter} + 15)
\]

### 4.4.2 Area threshold

To remove the cells and the remaining background, an area threshold is performed. To do so, all the contours on the image are found. As the filariae and cells will have the highest intensity values by the previous threshold applied, only the contours with white pixels surrounded by black pixels will be taken into account. This will also allow to ignore the parts of the filaria that make a loop. To find cells and pieces of background, the minimum enclosing circle is found for each contour and then, the area of the minimum enclosing circle is compared against the contour area, as a rough way to find circularity. As the filariae have the shape of a worm, the minimum enclosing circle area will never be similar to the contour area, the last one will always be much smaller. To find the range of areas for the cells and background, several images were tested in different resolutions and with different area thresholds to find out which was the maximum difference between the minimum enclosing circle area and contour area that could cover all the cells and pieces of background. The equation found has, as variables, the
magnification used for the image in the microscope and the image diameter calculated in the pre-processing phase. For loaded background images, the constant used is slightly bigger than for the other images to be sure that background pieces get segmented.

\[
\text{minCircleArea} - \text{ContourArea} \leq 50 \times \text{diameter} \times \left( \frac{\text{diameter}}{\text{magnification}} \right) \tag{4.10}
\]

\[
\text{minCircleArea} - \text{ContourArea} \leq 30 \times \text{diameter} \times \left( \frac{\text{diameter}}{\text{magnification}} \right) \tag{4.11}
\]

With the above, the cells and pieces of background are removed, isolating the filaria.

![Figure 4.12 Result of Area Threshold on both samples. Cells are removed.](image)

Type: *Wuchereria bancrofti* on the left and *Brugia malayi* on the right

The next step was to remove the cells attached to the filariae that cannot be segmented in this step.

### 4.4.3 Attached Cells Removal

To remove the cells attached to the filaria the method is similar to pre-processing. There is a color normalization with the same target image, but this time, the input image is cropped before, which will make that the lightest regions become black pixels. Afterwards, a morphological operation is applied to restrict the size of the areas that become black followed by a threshold, and finally the image is segmented to isolate only the cells.
4.4.3.1 Pre-Processing

As the cells and some parts of the filaria have stronger colors, these are the lightest areas after color normalization and they will become black once it is applied. Following, the normalized image is converted to grayscale and a morphological operation is applied, but this time a Black Hat, which is the difference between the input image and it’s closing (closing morphological operation applied on the input image). The size of the structuring element in relation to the diameter is given by:

\[
elementSize = 2 \times \text{roundUP}(0.010765 \times \text{diameter} + 2.1566) + 1
\] (4.12)

Adding the result of the operation to the grayscale image, the result is the image with all the black regions but only those who are smaller or equal to the size of the structuring element.

![Original cropped image on the left. Result of color normalization in the cropped image in the middle. Grayscale conversion of the normalized image and addition of the result of the morphological operation. Type: *Wuchereria bancrofti*](image1)

![Original cropped image on the left. Result of color normalization in the cropped image in the middle. Grayscale conversion of the normalized image and addition of the result of the morphological operation. Type: *Brugia malayi*](image2)
4.4.3.2 Segmentation

To segment the image first we need to apply a threshold. Adaptive threshold is used again to do so, exactly for the same reasons pointed out on the pre-processing step to find the filaria. The same algorithm is used to find the threshold, the adaptive Gaussian mean function, and the window size is, again, defined with an expression having the diameter as variable:

\[
\text{windowSize} = 3.81554 \times 10^{-6} \text{diameter}^2 - 0.0291736 \times \text{diameter} - 16.9013 \quad (4.13)
\]

Both the cells and some parts of the filaria will be black regions at these points, for what there is the need to segment and get only the cells, which are the main focus. Consequently, the “circularity” is verified exactly as described in the previous segmentation phase, Area Threshold. Finding all the contours and, for each, subtracting the minimum enclosing area to the real area of the contours and finding all the contours that obey to the following area restriction:

\[
\text{minCircleArea} - \text{Contour Area} \leq 2 \times \text{diameter} \times \left( \frac{\text{diameter}}{\text{magnification}} \right) \quad (4.14)
\]

\[
\text{area} > \text{diameter} \times 1.7 \quad (4.15)
\]

Figure 4.15 Adaptive Threshold of the added image on the pre-processing. Type: \textit{Wuchereria bancrofti} on the right and \textit{Brugia malayi} on the left

There is also an extra restriction where the area needs to be bigger than a threshold, to prevent smaller contours that are not cells and are, possibly, parts of the filaria to be segmented and consequently cut parts of the filaria.
After the cells are subtracted from the image that resulted from the area threshold, a morphological opening is performed in order to detach smaller pieces that remain after subtraction. The size of the structuring element is given by:

\[
elementSize = 2 \times \text{round}(0.00358 \times \text{diameter} - 0.7188) + 1
\]  

(4.16)

4.5 Filaria Reconstruction

The result of the segmentation of the cells will be subtracted from the segmented image with the filaria. As it was mentioned before, the method to remove the attached cells may cause unexpected cuts on the filaria due to cells that are in front of the filaria or simply because some regions of the filaria are lighter than others, causing the algorithm to mistaken them as cells. To address this problem, all the contours are calculated, as the separation of the filaria will form two different bodies. Then, if the contours are within a certain distance, they are connected through the drawing of lines between those points. The area of each of the bodies has to be superior to a certain value in order to be a valid contour, guaranteeing this way that no other form is connected to the filaria causing errors on isolating the filaria. The area and the distance are calculated through the following formulas:
where distanceFactor is a constant with the value 4.72089.

\[
area > 6.7384 \times diameter - 2627.71 
\]

(4.17)

\[
distance \leq 0.03189 \times diameter + distanceFactor 
\]

(4.18)

In the image of figure 4.17, the filaria is connected allowing to get a better segmentation result which will be essential for feature extraction and classification phases, as it will appear to it as a whole body and not as different bodies to train and classify. This will avoid errors in training and classification such as wrong length, area, perimeter calculation and more.

### 4.6 Feature Extraction

After the segmentation phase, the features from the isolated areas need to be obtained. These features will allow the machine learning to determine whether the image contains a parasite or not, and which type of parasite it is in case of positivity. To distinguish the filaria from the cells there is an obvious difference in sizes, shape, and circularity, as well as there are differences in texture and color. To distinguish one type of parasite from the other, geometry might not be the way to go as the geometry is really similar between them, and to find the differences on the computer would be a very difficult job. Only color and texture features were
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capable of that distinction and consequently used here. As one type has disperse nuclei in the body and the other has a body loaded with nuclei the differences will be clear.

So, three main groups of features were extracted: color features, texture features and geometry features.

6  4.6.1 Color & Texture Features

For color and texture features there are two types of statistics that can be obtained from a texture analysis. The first and second order statistics [57].

First-order statistics can be taken from a calculated histogram of an image to quantitatively describe the first-order statistical properties of the image [57]:

Mean. Average level of intensity

Variance. Variation of intensity around the mean

Skewness. Zero if histogram symmetrical around the mean, positive or negative depending if it has been skewed above or lower the mean.

Kurtosis. Measure for flatness of the histogram.

Energy.

\[ Energy = \sum_{i,j} p(i,j) \]  \hspace{1cm} (4.19)

Entropy. Measure of histogram uniformity.

\[ Entropy = \sum_{i,j} p(i,j) \log(p(i,j)) \]  \hspace{1cm} (4.20)

For color features these calculations are made for \( l,a,b \) color space.
Second-order statistics are defined as the co-ocurrence matrix which is a square matrix of dimension equal to the number of intensity levels in the image, for each distance and orientation. As it has too many elements for texture analysis, some features can be calculated using the co-ocurrence matrix to discriminate the texture and avoid costly computations:

**Contrast.**

\[
Contrast = \sum_{i,j} (i-j)^2 p(i,j) \tag{4.21}
\]

**Correlation.**

\[
Correlation = \sum_{i,j} \frac{p(i,j)(i-u_i)(j-u_j)}{\sigma_1 \sigma_2} \tag{4.22}
\]

**Homogeneity.**

\[
Homogeneity = \sum_{i,j} \frac{p(i,j)(i-u_i)(j-u_j)}{\sigma_1 \sigma_2} \tag{4.23}
\]

**Inverse Difference.**

\[
InvDiff = \sum_{i=0}^{G-1} \sum_{j=0}^{G-1} \frac{p(i,j)}{1 + (i-j)^2} \tag{5.4}
\]

**Entropy.**

\[
Entropy = \sum_{i=0}^{G-1} \sum_{j=0}^{G-1} p(i,j) \log(p(i,j)) \tag{4.24}
\]
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Maximum Probability.

\[ \text{max} P = \text{max}_{i,j} p(i,j) \quad (4.25) \]

Absolute Value.

\[ \text{AbsValue} = \sum_{i=0}^{n} \sum_{j=0}^{n} |i - j| p(i,j) \quad (4.26) \]

6 4.6.2 Geometry Features

Geometry features can be obtained through boundary points, contour-based methods. Such features are perimeter, perimeter-resultant measures such as area, circularity, solidity and so on. Below, there are some shape-based features presented and their calculation method [58]:

**Area.** Area of the shape that can be obtained through contour.

**Perimeter.** Perimeter of the shape that can be obtained through contours.

**Convex Hull.** Smallest convex that contains a shape.

**Convexity.** Ratio between the perimeter of the convex hull and the original shape.

\[ \text{Convexity} = \frac{P_{\text{Convex Hull}}}{P_{\text{Original Shape}}} \quad (4.27) \]

**Solidity.** Whether a shape is convex or concave. Convex shapes have value 1 of Solidity.

\[ \text{Solidity} = \frac{A_{\text{shape}}}{A_{\text{Convex Hull}}} \quad (4.28) \]
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Circularity. How close is a shape from a circular form.

\[ \text{Circularity} = \frac{P_{\text{shape}}^2}{4\pi A_{\text{shape}}} \] (4.29)

Rectangularity. How close is a shape from a rectangular form.

\[ \text{Rectangularity} = \frac{A_{\text{shape}}}{A_{\text{MinBoundingBox}}} \] (4.30)

Thinness Ratio.

\[ \text{Thinness Ratio} = \frac{4\pi A_{\text{shape}}}{P_{\text{shape}}^2} \] (4.31)

Euler Number. Difference between the contiguous parts and the number of holes in an image.

\[ \text{Euler Number} = S - N \] (4.32)

Holes Area Ratio. Ratio between the sum of the holes areas and the area of the shape

\[ H \ A \ R = \frac{\sum A_{\text{holes}}}{A_{\text{shape}}} \] (4.33)
**Eccentricity.** It is the measure of aspect ratio and can be calculated through the ratio of the length of major axis to the length of the minor axis.

\[
Eccentricity = \frac{\lambda_{\text{MinorAxis}}}{\lambda_{\text{MajorAxis}}} \tag{4.34}
\]

**Minimum Bounding Box Area.** Minimum bounding box is the smallest rectangle that contains every point of the shape. In the minimum bounding box the major axis is the length and the minor axis the width. Therefore, the area of a rectangle is \( length \times width \) so:

\[
\text{areaBoundingBox} = \lambda_{\text{MinorAxis}} \times \lambda_{\text{MajorAxis}} \tag{4.35}
\]

**Elongation.** Another concept based on eccentricity. Shapes with large ratio will have an elongation of 1 while shapes that are more symmetric will have an elongation of 0.

\[
f_{\text{elong}} = \frac{i_{2}}{\sqrt{i_{1}}} \tag{4.36}
\]

**Minimum Enclosing Circle.** Smallest circle that contains every point of the shape. Here, the area, perimeter, radius of the circle can be obtained.

\[
\sigma_{i} = \sum_{i,j} p(i,j)(i - w_{i})^{2} \tag{4.37}
\]
4.7 Classification

4.7.1 Classification Methods

There are two classification processes in this project. The first one is to classify features between cells and parasite and the second one, with the features of objects classified as parasite from the first classification, to classify between Brugia malayi and Wuchereria bancrofti. Both classifications were tested with two different machine learning algorithms, k-Nearest-Neighbor (kNN) and Support Vector Machines (SVM) using radial basis function (RBF). For each it was performed a parameter optimization as well as a verification of the algorithm results using a k-Fold-Cross-Validation. Also the accuracy, specificity and sensibility were calculated and based on those results the best parameters were chosen. For kNN the parameter to optimize is the default K which determines the class of the feature based on the class that has the most neighbors assigned, from the K neighbors. On the SVMs, the parameters to optimize are C and γ according to:

\[ K(x_i, x_j) = e^{-\gamma \|x_i - x_j\|^2}, \quad \gamma > 0 \]  

(4.38)

To perform all the tests a different project was set up to get the final results and the final training files to be used on the classification process. The diagram in figure 4.18 shows the flow of the supportive program:

![Diagram](image)

Figure 4.18 Support program flow to perform train and classification
Lymphatic Filariasis Detection

The training and classification images were obtained by using the Monte Carlo’s algorithm with image balancing where 67% of the dataset is for training and the remaining 33% for classification dividing the dataset into 73 images for training and 36 images for classification.

4.7.2 Classification Results

Both the kNN and SVM got impressive levels of accuracy when optimized. This also depended on the features used in the training. To differentiate between cells and parasite, geometry is a very important feature because the size of the filaria by itself will be a determining parameter, but to differentiate between both types of parasite the geometry features would confuse the algorithm and get lower results so the best way to differentiate both types is to use only color and texture features as the texture from both filariae has significant differences.

With the results presented in tables 4.1 and 4.2 the chosen algorithm was the SVM using a radial basis function for both the first and second SVMs. The optimization process was conducted by performing training and classification with different parameters and adjusting them according to the previous results. The best results were with SVMs that had $\gamma$ lower than 1 and for both, the SVM using $\gamma$ with the value of 0.1 and $C$ with the value of 1 got optimal results. For the first SVM it was needed to set class weights due to the disparity between the positive and negative masks. The first SVM is the one to classify between parasite and cells and the number of parasite features was of 76 while the number of negative features was of 16448 features in training. By setting class weights, the SVM knows of this disparity and classifies based on it. Without the class weights the SVM failed in every classification, classifying everything as cells but when the class weights were added the SVM produced optimal results. For the second SVM this was not needed as the images used for training were evenly distributed (55 belonging to the first sample and 54 belonging to the second).
Table 4.1 Results of machine learning algorithms in classification between parasite and cells, using color, texture and geometry features

<table>
<thead>
<tr>
<th>Classifier Algorithms</th>
<th>Accuracy</th>
<th>Sensibility</th>
<th>Sensitivity</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>kNN</td>
<td>99.96%</td>
<td>92.30%</td>
<td>100.00%</td>
<td>k = 3</td>
</tr>
<tr>
<td>kNN</td>
<td>99.96%</td>
<td>92.30%</td>
<td>100.00%</td>
<td>k = 5</td>
</tr>
<tr>
<td>kNN</td>
<td>99.96%</td>
<td>92.30%</td>
<td>100.00%</td>
<td>k = 7</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.95%</td>
<td>89.74%</td>
<td>100.00%</td>
<td>C = 1; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.96%</td>
<td>92.30%</td>
<td>100.00%</td>
<td>C = 1; y = 0.3</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.92%</td>
<td>84.61%</td>
<td>100.00%</td>
<td>C = 1; y = 0.5</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.91%</td>
<td>82.05%</td>
<td>100.00%</td>
<td>C = 1; y = 0.7</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.95%</td>
<td>89.74%</td>
<td>100.00%</td>
<td>C = 0.8; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.95%</td>
<td>89.74%</td>
<td>100.00%</td>
<td>C = 0.6; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.95%</td>
<td>89.74%</td>
<td>100.00%</td>
<td>C = 0.4; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>99.95%</td>
<td>89.74%</td>
<td>100.00%</td>
<td>C = 0.2; y = 0.1</td>
</tr>
</tbody>
</table>

Table 4.2 Results of machine learning algorithms in classification between both types of filariae, using color and texture features

<table>
<thead>
<tr>
<th>Classifier Algorithms</th>
<th>Accuracy</th>
<th>Sensibility</th>
<th>Sensitivity</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 3</td>
</tr>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 5</td>
</tr>
<tr>
<td>kNN</td>
<td>92.30%</td>
<td>94.44%</td>
<td>90.47%</td>
<td>k = 7</td>
</tr>
<tr>
<td>kNN</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>k = 9</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>90.47%</td>
<td>100.00%</td>
<td>C = 1; y = 0.3</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>87.17%</td>
<td>79.16%</td>
<td>100.00%</td>
<td>C = 1; y = 0.5</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>79.48%</td>
<td>70.37%</td>
<td>100.00%</td>
<td>C = 1; y = 0.7</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.8; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.6; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.4; y = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.2; y = 0.1</td>
</tr>
</tbody>
</table>
4.7.2.1 k-Fold Cross Validation

To validate the results of the second SVM, as it was the most critical one due to the similarity in geometry and basing only the training in color and texture, k-Fold Cross Validation was used. In k-Fold Cross Validation the dataset is divided into n groups of equal size. As the dataset consists of 109 images, 55 from a type and 54 from another, the resulting division was of 10 groups of 10 images each and 1 group with 9 images. The images in each group are also balanced to have 5 images from the first sample and 5 from the second except for the group with 9 images that has 5 images from the first sample and 4 from the second one. No images could be repeated. After the division, the aim of the k-Fold Cross is to train n-1 groups and classify the remaining one and to repeat this process n times, each time picking a different group to classify and consequently, having different trainings.

Table 4.3 Visualization of k-Fold Crossing. Each column represents a group, each row an iteration.

<table>
<thead>
<tr>
<th>Train</th>
<th>Train</th>
<th>Train</th>
<th>Train</th>
<th>Train</th>
<th>Train</th>
<th>Train</th>
<th>Train</th>
<th>Train</th>
<th>Train</th>
</tr>
</thead>
<tbody>
<tr>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
</tr>
<tr>
<td>Train</td>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
</tr>
<tr>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
</tr>
<tr>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
</tr>
<tr>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
</tr>
<tr>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
</tr>
<tr>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
<td>Train</td>
</tr>
<tr>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Train</td>
<td>Classify</td>
<td>Train</td>
</tr>
</tbody>
</table>

The results of the classification were grouped and the final result was a recognition of 58 features from parasites from type *Brugia malayi* out of 58 and 54 features from parasites from type *Wuchereria bancrofti* out of 57 with a final accuracy of 97.39%.
Lymphatic Filariasis Detection

Table 4.4 Results of the classification with the optimized SVM

<table>
<thead>
<tr>
<th></th>
<th>Brugia malayi</th>
<th>Wuchereria bancrofti</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brugia malayi</td>
<td>58</td>
<td>0</td>
</tr>
<tr>
<td>Wuchereria bancrofti</td>
<td>3</td>
<td>54</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Classifier Algorithm</th>
<th>Parameters</th>
<th>Accuracy 97.39%</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVM: RBF</td>
<td>C = 1; γ = 0.1</td>
<td></td>
</tr>
</tbody>
</table>
Chapter 5

Image Processing in Android

The ultimate goal of this thesis work was to develop an automated image processing methodology capable of detecting and classifying between the two types of filariae. This automated methodology is to be integrated in the MalariaScope project, especially in its mobile-application. In this chapter the MalariaScope project is presented as well as the application. Later, the integration of the image processing methodology developed with the MalariaScope application will be addressed.

5.1 Integration with MalariaScope Project

The MalariaScope project is a mobile-based solution to provide automatic detection of Malaria in blood samples. It is used as a cheap alternative to microscopes, constituting a prototype developed by Fraunhofer to obtain images from a blood sample which will be analyzed with an automated image processing tool to detect the malaria parasite. The analysis to perform has two main stages: the determination of the parasite density in a blood sample and the identification of the species. As Lymphatic Filariasis is a severe problem in regions like Africa where it is difficult the access to health specialists, medical equipment and staff to process all the cases that exist, an automated tool would be a life changer and that is where this project comes in. The idea is to use the same prototype and application for different automated image processing methodologies in order to automate the detection of certain parasites which allows anyone to use the application without the need of a specialist or an expensive microscope.
5.2 Application Overview

The MalariaScope application has a simple built-in database. It is represented by one-to-many relations between patients and sessions and, sessions and samples which means that a patient can have one or more than one session and one session can have one or more samples. Given this, the main screen of the application, after the logo intro, is the patient database where it is possible to create a new patient or use/delete a patient that already exists. From here, after accessing a patient, the patient screen is shown, where there are the patient samples and the user can create or load a sample.

![Patient database, main screen after logo intro.](image-a)
![Samples screen of “Patient 1”](image-b)
![Screen to add sample](image-c)

**Figure 5.1** a) Patient database, main screen after logo intro. b) Samples screen of “Patient 1”. c) Screen to add sample

Each sample has views, and these views can be of three types: Camera Manual, Camera Automatic or Gallery. Camera Manual allows the user to explore the blood smear through the smartphone that is controlling the microscope, moving along the blood smear. Camera Automatic takes a certain number of pictures defined by the user, automatically, by moving along the blood smear with an auto-focus algorithm. Finally, images can simply be imported from the smartphone Gallery to be analyzed. Images will appear in the views with a blue color if they haven’t been analyzed yet and green when the analysis is finished.
The image can then be processed running in background, meaning that there can be more than one view being processed and there is no need to be on the application waiting for the analysis to be finished. After the analysis is finished, a notification appears on the smartphone informing that the analysis has been finished. While it isn’t finished, there is also a status bar showing how many images have been analyzed.

**Figure 5.2** a) Screen when accessing a sample. b) Adding a view to a sample. c) Image loaded to the view and ready to be analyzed (in blue)

**Figure 5.3** a) Hitting the button to process an image, notification saying it will run in background. b) Status screen of analysis running in background. c) Notification of when the analysis is finished
The analysis is performed at the smartphone level, meaning that there is no need to have an external connection or processing to a server. In the end, the analysis is displayed in a new activity with the form of a report showing the image with the parasite, if found, marked in green.

![Image](image.png)

**Figure 5.4** Final result with the found parasite marked in green and respective classification below

This application was adapted to this project in order to maintain the application consistency between the MalariaScope applications.

### 5.3 Image Processing Integration

To perform the analysis in the application, a sample must be added with images from the gallery or microscope/prototype. After images have been added, clicking the arrow icon causes the program to perform the analysis of every image that is listed in the sample view. After completion, the image thumbnail will become of a different color (green) and the image can be seen already with the analysis performed and the parasite marked. Since the analysis task is a CPU (Central-Processing-Unit) intensive task and it may take some time to perform the analysis to all the images, it is needed to be performed in background. That is done by moving the task
Image Processing in Android

from the User-Interface thread to another thread created by the class IntentService which handles each request in turn using a worker thread. Therefore, the user can be viewing the results or performing other analysis while the application is performing the image processing operations.

The image processing methodology was integrated in the application using JNI (Java Native Interface), which defines a way for managed code written in Java to interact with native code written in C++. This way, the Java code can invoke the local functions written in C++. But this is not enough as it is needed a cross-development tool to compile the C++ code. The tool, supplied by Google, is the Android NDK (Native Development Kit). The NDK will allow the application to run on Android devices. With the NDK there is direct access to the CPU and hardware, and OpenCV functions in Java can be optimized invoking implementations in C++. NDK is therefore, the right option to choose to integrate the methodology in native code, even though it will need to be compiled for every architecture in order to work on the majority of Android devices. OpenCV also has the libraries prepared for Android which need to be integrated in the application in order to recognize the OpenCV functions. This is as simple as add a module to the project and import the OpenCV Android module that can be downloaded from their main page.
Chapter 6

Results

In this chapter the results from segmentation and classification will be presented. Several classifications with different features and different algorithms were performed in order to choose the best classification algorithm with optimized parameters. The results for these classifications will be presented in this chapter as well as a discussion about the different classifications and consequently, the conclusions to be drawn. Afterwards, a discussion about the deficiencies and strong points of the methodology will be addressed.

6.1 Results & Discussion

The segmentation phase was able to detect and segment all the filariae in the dataset. With the dataset used, it is guaranteed that the filariae will be present in the final image. However, some of the images may have some background or remaining cells from the attached cells removal method in section 4.4.3 attached to the filaria. Therefore, to evaluate the segmentation phase results, resulting images were assigned to 4 groups. Images of the filaria after segmentation that have no background attached to its boundaries and no remaining cells were considered well isolated, figure 6.1. Images with a low amount of background or one cell attached to the filaria were assigned to the + group, figure 6.1. Images with a significant amount of background attached to the boundaries or more than one cell were assigned to the ++ group figure 6.2 For the contrary, images where the filaria is not complete (overly segmented) these images were assigned to the – group, figure 6.2.
Results

Based on these assumptions, the results of the segmentation phase can be observed in table 6.1.

**Figure 6.1** Well isolated filaria on the left. Image assigned to the + group on the left due to some background attached to the filaria. Both filariae from type: *Wuchereria bancrofti*

**Figure 6.2** Image that will be assigned to the ++ group on the left due to having more than one cell attached to the filaria. Image that will be assigned to the group − on the right due to a missing piece. Types: *Wuchereria bancrofti* on the left and *Brugia malayi* on the right.
Results

Table 6.1 Results of segmentation phase

<table>
<thead>
<tr>
<th>Well Isolated</th>
<th>+</th>
<th>++</th>
<th>-</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Brugia malayi</strong></td>
<td>45</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td><strong>Wuchereria bancrofti</strong></td>
<td>22</td>
<td>23</td>
<td>9</td>
</tr>
</tbody>
</table>

The images that had more than the filaria were mostly from *Wuchereria bancrofti*’s sample which is the one that has a more intense blue color due to being left to stain for more time. This results in a more loaded background and the cells having similar colors to the filaria which might be the cause to poor isolation only on one sample. As it can be observed in figure 6.1 and 6.2, the images from the groups + and – still have a good segmentation even though there are minor artifacts or a missing piece. We believe that these minor artifacts or missing pieces do not have a significant impact on the classification phase. For the ++ group images, these might cause some impact on the train and classification as the filariae becomes really different but as they are only 9 images in a dataset of 109 and the other groups of images (well isolated, + and -) got well classified this is not a decisive factor.

In relation to the classification phase, the same machine learning algorithm was chosen for both the classifications, between cell and parasite, and between the two species of parasites but, with different parameters. The final results can be observed in the table below:

Table 6.2 Results of final classification between parasite and cell

<table>
<thead>
<tr>
<th>Parasite</th>
<th>Cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parasite</td>
<td>36</td>
</tr>
<tr>
<td>Cells</td>
<td>0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Classifier Algorithm</th>
<th>Parameters</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVM: RBF</td>
<td>C = 1; γ = 0.3</td>
<td>99.9%</td>
</tr>
</tbody>
</table>
Table 6.3 Results of final classification between the two types of parasites

<table>
<thead>
<tr>
<th></th>
<th>Brugia malayi</th>
<th>Wuchereria bancrofti</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brugia malayi</td>
<td>19</td>
<td>1</td>
</tr>
<tr>
<td>Wuchereria bancrofti</td>
<td>0</td>
<td>19</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Classifier Algorithm</th>
<th>Parameters</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVM: RBF</td>
<td>C = 1; γ = 0.1</td>
<td>97.43%</td>
</tr>
</tbody>
</table>

where the levels of accuracy, sensibility and specificity are very high. This could be due to the difference in color on both samples, which depends on the time that it is left for stain. One sample may have a different color from another if it is left for stain for more or less time. This might have helped the results but tests were conducted in order to verify if the results were highly dependable of certain features. To do so, classifications using only texture, color or geometry features were performed, as well as a classification with the usage of the three feature groups (texture, color and geometry).

The results were as expected and as can be observed in the original images from both types (Brugia malayi and Wuchereria bancrofti). Color is definitely a decisive feature, the colors are very different between both the types. Texture is also very distinctive as the images from Wuchereria bancrofti’s sample have a texture with small nuclei separate from each other while images from Brugia malayi’s sample have a more unified texture. Now for geometry, is very difficult to find a distinctive measure to differentiate both types from geometry features and, although it didn’t performed that poorly, the results were not the best, hence, making geometry an indecisive feature. These results can be observed in tables 6.1, 6.2 and 6.3.

As geometry is not the most distinctive feature, and the results are pretty similar between the classifications with all features (table 6.7) and with texture and color features only (section 4.7.2, table 4.2), geometry features were not included in the final classification algorithm to speed up the process.
## Results

**Table 6.4** Results of classification with different machine learning algorithms using only geometry features

<table>
<thead>
<tr>
<th>Classifier Algorithms</th>
<th>Accuracy</th>
<th>Sensibility</th>
<th>Sensitivity</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>kNN</td>
<td>76.92%</td>
<td>91.66%</td>
<td>70.37%</td>
<td>k = 3</td>
</tr>
<tr>
<td>kNN</td>
<td>74.35%</td>
<td>84.61%</td>
<td>69.23%</td>
<td>k = 5</td>
</tr>
<tr>
<td>kNN</td>
<td>69.23%</td>
<td>81.81%</td>
<td>64.42%</td>
<td>k = 7</td>
</tr>
<tr>
<td>kNN</td>
<td>66.66%</td>
<td>75.00%</td>
<td>62.96%</td>
<td>k = 9</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>82.00%</td>
<td>87.50%</td>
<td>78.26%</td>
<td>C = 1; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>82.00%</td>
<td>87.50%</td>
<td>78.26%</td>
<td>C = 1; γ = 0.3</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>76.92%</td>
<td>81.25%</td>
<td>73.91%</td>
<td>C = 1; γ = 0.5</td>
</tr>
<tr>
<td>SVM: RBF</td>
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<td>83.33%</td>
<td>80.95%</td>
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<tr>
<td>SVM: RBF</td>
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<td>54.54%</td>
<td>58.82%</td>
<td>C = 0.8; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>53.84%</td>
<td>52.17%</td>
<td>56.25%</td>
<td>C = 0.6; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>53.84%</td>
<td>52.17%</td>
<td>56.25%</td>
<td>C = 0.4; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>51.28%</td>
<td>50.00%</td>
<td>53.33%</td>
<td>C = 0.2; γ = 0.1</td>
</tr>
</tbody>
</table>

**Table 6.5** Results of classification with different machine learning algorithms using only texture features

<table>
<thead>
<tr>
<th>Classifier Algorithms</th>
<th>Accuracy</th>
<th>Sensibility</th>
<th>Sensitivity</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>kNN</td>
<td>92.30%</td>
<td>94.44%</td>
<td>90.47%</td>
<td>k = 3</td>
</tr>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 5</td>
</tr>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 7</td>
</tr>
<tr>
<td>kNN</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>k = 9</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.3</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.5</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.7</td>
</tr>
<tr>
<td>SVM: RBF</td>
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<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.8; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
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<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.6; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>C = 0.4; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>92.30%</td>
<td>94.44%</td>
<td>90.47%</td>
<td>C = 0.2; γ = 0.1</td>
</tr>
</tbody>
</table>
Results

Table 6.6 Results of classification with different machine learning algorithms using only color features

<table>
<thead>
<tr>
<th>Classifier Algorithms</th>
<th>Accuracy</th>
<th>Sensibility</th>
<th>Sensitivity</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>kNN</td>
<td>76.92%</td>
<td>91.66%</td>
<td>66.66%</td>
<td>k = 3</td>
</tr>
<tr>
<td>kNN</td>
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<td>92.30%</td>
<td>73.07%</td>
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</tr>
<tr>
<td>kNN</td>
<td>79.48%</td>
<td>92.30%</td>
<td>73.07%</td>
<td>k = 7</td>
</tr>
<tr>
<td>SVM: RBF</td>
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<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.3</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>C = 1; γ = 0.5</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>92.30%</td>
<td>94.44%</td>
<td>90.47%</td>
<td>C = 1; γ = 0.7</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.8; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
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<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.6; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
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<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.4; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>92.30%</td>
<td>94.44%</td>
<td>90.47%</td>
<td>C = 0.2; γ = 0.7</td>
</tr>
</tbody>
</table>

Table 6.7 Results of classification of different machine learning algorithms using all features (texture, color, geometry)

<table>
<thead>
<tr>
<th>Classifier Algorithms</th>
<th>Accuracy</th>
<th>Sensibility</th>
<th>Sensitivity</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 3</td>
</tr>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 5</td>
</tr>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 7</td>
</tr>
<tr>
<td>kNN</td>
<td>94.87%</td>
<td>94.73%</td>
<td>95.00%</td>
<td>k = 9</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>94.87%</td>
<td>90.47%</td>
<td>100.00%</td>
<td>C = 1; γ = 0.3</td>
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<tr>
<td>SVM: RBF</td>
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<td>79.16%</td>
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<td>C = 1; γ = 0.5</td>
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<td>100.00%</td>
<td>C = 1; γ = 0.7</td>
</tr>
<tr>
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<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.8; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.6; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.4; γ = 0.1</td>
</tr>
<tr>
<td>SVM: RBF</td>
<td>97.43%</td>
<td>95.00%</td>
<td>100.00%</td>
<td>C = 0.2; γ = 0.1</td>
</tr>
</tbody>
</table>

Although the attached cells removal algorithm produces good results, it is not optimal as some cells are not detected due to differences in color which is a deficiency of the algorithm. Another downside is in the segmentation, where some remaining background may be attached to the filaria and although they are only small parts, they don’t produce a perfect segmentation. This is mainly due to the extra illumination on the edges of the optical circle or some extra noise from the camera or lens dust that will alter the colors of the image and introduce new colors.
Results

Although the classification got very good results, these results will be hard to maintain on bigger datasets where the number and variability of the images is bigger. The dataset is small, as the number of different filarias is reduced. So, the images to the filariae had to be taken in different positions, shifting the sample and rotating it upside down to produce the effect of new filariae. As the images will be different once the sample is shifted or rotated (different cells, background, illumination) it covers well the fact that the number of filariae is reduced but this might have helped in the classification phase.

Nonetheless, the algorithm does segment the parasite and classifies it accordingly whether it has the extra bodies or not and the worst results are the ones with a little of background attached to the boundaries of the filaria or some cells that have subsisted to the attached cells removal. The filaria will be intact in the end of the segmentation and, if the image is on focus and with no extra objects produced by dust on the camera or some extra light on the edges of the optical circle, the algorithm produces very good results in segmentation having no background attached to the filaria, hence, producing a smooth result.
Chapter 7

Conclusions and Future Work

The aim of the project was to develop a robust image processing methodology for low quality images acquired using the Fraunhofer mobile acquisition prototype or any other low cost device like Skylight to detect and classify parasites that cause Lymphatic Filariasis into the two most common types, *Brugia malayi* and *Wuchereria bancrofti*. The proposed algorithm consisted on the segmentation of the parasite and classification in one of the two types. The algorithms employed in the pre-processing and segmentation allowed the parasite to be segmented correctly in low quality images with variance in lighting and color. The classification phase results show a very good detection and discrimination achieving an accuracy of 99% to classify between cell and filariae and between the two types of filariae. As discussed that can be due to the small set of specimens provided by the Hygiene and Tropical Medical Institute.

For future work, the algorithm should be tested in high quality image datasets and other low quality datasets without noticeable differences in coloration to test the accuracy and reliability of the methodology. The Android application should be improved as it is a primary version only to test the results on mobile. Although it is adapted to the MalariaScope application this should be fully integrated in the application where it is possible to choose which type of analysis the user wants to perform on the blood sample. The speed of the algorithm should also be addressed as the high size used in the morphological operations and median filters is very costly. Possibly, with the insertion of threads to perform the attached cells removal in parallel with the segmentation the operations could become faster. The application can be extended to other parasites that are very similar with the parasites presented in this work.

In conclusion, this work provides a robust methodology to find Lymphatic Filariasis parasites and their types, which was never tried before, and a stable background work to expand the detection to other parasites of the same type (*Loa Loa*) and extend the range of detection of parasites in low quality images.
References


References


References


[34] http://www.cdc.gov/parasites/lymphaticFilariasis/diagnosis.html


References


References


References