Non-invasive Glaucoma Screening Using Ocular Thermal Image Classification

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Ocular thermography is an important, emerging modality in the diagnosis and management of diseases related to eye. It is a non-invasive procedure to evaluate the presence of eye diseases and monitor the response to treatments. In this paper, we propose and evaluate a system designed using infrared thermal image processing that detects glaucoma. Euclidean distance based segmentation technique is used to threshold the IR image to obtain the region of interest, where the manifestation of glaucoma is predominant. Features are extracted using statistical moments from the temperature mapped IR image and Gray Level Co-Occurrence Matrix of the IR image. Two significant attributes, namely the homogeneity and area of region of interest are the inputs to a Support Vector Machine classifier to classify a given input ocular thermal image as a normal or diseased image. In our simulation study, one hundred ocular thermal images with even number of normal and diseased subjects were analysed. The classifier has achieved a maximum accuracy of 96% when homogeneity and area of region of interest are used as attributes, indicating the potential use of proposed method for screening patients even at early stages of glaucoma.

1. Introduction

Ocular thermography, which is a relatively new field in medical imaging, has so far been used to detect dry eye syndrome and certain inflammatory and non-inflammatory eye diseases [1]. Glaucoma is a disease that is known as a silent stealer of vision. It involves increase in eye pressure and results in damage of the optic nerve, thereby affecting vision. Increased corneal temperature is also an indication of glaucoma. The average value of corneal temperature ranges between 32°C to 35°C for a healthy individual. Glaucoma is usually detected using various tests as listed briefly in Table 1. However, these methods generally require examination by a proper ophthalmologist. In this paper, we propose infrared thermography as one of the potential alternate methods for glaucoma detection. Thermal imaging records the temperature distribution of the body and thus can provide insight in dysfunction associated with inflammation, abnormality or with pain. In addition, thermal imaging involves no ionizing radiation; the test is comfortable and analogous to the simple taking of a photograph [2]. Other modality of imaging does not reveal such abnormalities in a non-invasive manner that is shown by thermal imaging.

Infrared thermography (IRT) is an imaging science which has a vast potential in medical applications. The foundation of IRT is based on Planck's Law of blackbody radiation, according to which all normal matter with a temperature above absolute zero (0 K) emits electro-
magnetic radiation. Infrared thermography or thermal imaging involves detection of infrared energy emitted from an object and conversion of this energy into a temperature distribution. It can be viewed as a visualization technique to observe the invisible effects of change in temperature in a body and its surroundings. A special kind of camera known as IR camera (thermal camera) is used in IR thermography to quantitatively measure the surface temperatures of objects. This camera detects radiation in the long-infrared range which accounts to the wavelengths of 9 – 14 μm.

In the 1950s, thermography was an imaging technique used in defence research to observe any movement by the enemy at night. This approach had been incorporated into medical imaging in 1957 by Dr. R. Lawson who discovered that patients suffering from breast cancer had a higher skin temperature over the cancerous tissue than a normal tissue. Since then, thermography has been used as a tool of prognostic significance to assess the pathology of the breast and from the 1970s, thermography has been extended to medical applications in the detection of conditions like tumors, breast cancer, fractures, etc. Amir et al. [5] presented a fully automatic high accuracy technique for classification of suspicious areas in thermogram images with the aim of assisting physicians in early detection of breast cancer. The proposed algorithm consists of four main steps: pre-processing and segmentation, feature extraction, feature selection and classification. Statistical, morphological, frequency domain, histogram and Gray Level Co-occurrence Matrix (GLCM) based features are extracted and the methods proved to be efficient.

GLCM texture features emphasise spatial relationship between pixels, rather than one single pixel intensities [6]. GLCM texture analyses of medical images are relatively computationally efficient and the GLCM features provide fairly straightforward interpretations [7]. Maria et al. [8], performed novel thermal imaging analysis technique for detecting inflammation in thyroid eye disease. Maria and Robert [9] proposed automatic eye corner detection and tracking algorithm in sequence of thermal medical images. Wieck et al. [1], have reported the use of thermal imaging for medical applications while introducing pre-processing concepts such as contrast enhancement using histogram modification, median filtering and threshold segmentation. They explored image processing tools for conversion of thermal and visual images by evaluating area and distance based on statistical differencing. Duarte et al. [10], have outlined three techniques for segmentation in thermal image, namely thresholding, edge detection segmentation and region based segmentation in order to identify the region of interest. Rajendra Acharya et al. [11], has computed the average corneal temperature and temperature deviation along cornea among different age groups of humans by analysing the thermal images of individual eye. This helps in making a temperature mapping based on the color intensities of the pixels of the thermal image.

Morgan et al. [12], has discussed potential applications of ocular thermography. The dry eye syndrome and its diagnosis using thermal imaging are detailed. Also, the relationship between eye temperatures and ocular parameters has been examined. It has been demonstrated that inflammatory or dry eye disease causes a statistically significant eye temperature of the affected eye. N. Selvarasu et al. [13], compute Euclidean Distances between pixels to segment a particular region in thermal image and detect abnormalities such as arthritis, fractures, etc. The Euclidean distances between a pixel and the average pixel of the “disease hotspot” is calculated and a suitable threshold is used to segment the image for detecting fractures. The same principle has been applied to segment the ocular thermal images and extract the region of interest. Ravudu et al. [14], have extensively listed detection algorithm for the ocular diseases such as glaucoma, diabetic retinopathy and age related macular degeneration. In particular, glaucoma detection techniques involved manual determination of the optic disc size and the usage of Hough transform. Abràmoff et al. [15], presented a brief overview of the most prevalent causes of blindness in the industrialized world that includes age-related macular degeneration, diabetic retinopathy, and glaucoma. The review is devoted to retinal imaging modalities and image analysis methods and their clinical implications.

In conclusion, there has been extensive research done on thermal imaging in medical applications using various segmentation operations such as thresholding, morphological operations, etc., to extract features and thereby detect diseases like the other conditions manifested through change in relative surface temperature.

The objective of this paper is to develop an infrared thermal image processing system to detect glaucoma, thereby initiating another way for non-invasive glaucoma detection. Initial work on glaucoma can be referred in [16], [17]. The remainder of the paper has been organised as follows: Section 2 explains the proposed methodology. Section 3 illustrates the classification results based on our simulation. The conclusion is provided in Section 4.

## 2. Proposed Methodology

The overall idea is to process the thermal images, extract features from the images and use these features to build a classifier that classifies an image into “Normal” or “Diseased”. On visual inspection, the thermal images of diseased eyes seemed to possess higher temperature in the eyelid region than that of normal eyes. This led to the conclusion that glaucoma manifests itself in the eyelids in thermal images and hence the area of the “Region of Interest (ROI)”, in terms of number of pixels which is the portion of the eyelid with relatively high temperature, is taken to be the first attribute while constructing a classifier. Two methodologies have been proposed for obtaining the second feature for the classifier; the first based on GLCM feature extraction and the second based on temperature mapping. The hallmark of glaucomatous progression is the cupping of the optic nerve head, the optic nerve is located in the back of the eye, which transmits images to the brain. The damage will in turn increase the pressure of the eye, which is called the intra ocular pressure. Hence in the initial stage of using thermal imaging one can find the pressure difference which is manifested by large increase in temperature, which in turn can be extracted using GLCM features. GLCM features capture such variations in the eye. The temperature mapping catches the changes in the eye in terms of temperature difference. The classifiers that have been built

Table 1. Factor examined for clinical analysis of glaucoma.

<table>
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<td>Tonometry</td>
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<td>Expensive equipment; Mastery of technique requires skill and practice</td>
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2.1. Euclidean Distance Based Segmentation of ROI

As the image data with thermal images has details of the whole eye with eye lids and lashes which are not necessary in the identification of glaucoma, hence the region of interest is considered before initiating any processing. Glaucoma affects the optic nerve and hence the area inside the eye is necessary for our investigation.

Euclidean distance is used to obtain the first attribute of the classifier. The Euclidean distance between every pixel, and the average pixel, a has been computed individually for the image and also for the individual Red, Green and Blue components using the equation

\[ D(z, a) = \left( (z_a - a_k)^2 + (z_b - a_k)^2 + (z_c - a_k)^2 \right)^{\frac{1}{2}} \]

where \( D(z, a) \) is the Euclidean distance and the subscripts of \( z \) and \( a \) represent Red, Green and Blue components.

Any image pixel \( z \) is said to be similar to \( a \) if the Euclidean distance between them is less than a specified threshold \( D_0 \) [13]. Thresholding is then carried out to extract the high intensity pixels alone and a binary image is thus generated. The white pixels in the binary image represent the ROI. The area is then computed in terms of the number of white pixels. The block diagram for this method is shown in Figure 2.

The thresholded Blue component of the normal and diseased images showed maximum difference and hence the area of the ROI obtained in terms of the number of pixels for the thresholded Blue component has been used as an attribute for classification.

2.2. GLCM Feature Based Classification

The Gray-Level Co-occurrence Matrix is the representation of different combinations of the pixel intensities in an image. Since the relationship between two pixels is analysed in a GLCM, it provides second order statistical information [18]. Features generated by this technique are also known as Haralick features [19]. The images are pre-processed to extract GLCM features as shown in Figure 3. Firstly, the RGB images are converted into grayscale images.

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<td>Homogeneity</td>
<td>( f_1 = \sum_{i,j} \frac{1}{1+(i-j)^2} p(i,j) )</td>
</tr>
<tr>
<td>2</td>
<td>Sum Variance</td>
<td>( f_2 = \sum_{i,j} (i-j)^2 p_{ij}(i) ) where ( p_{ij}(k) = \sum_{i,j} p(i,j) ) if ( i+j = k ), ( i+j = 2k ), ( i+j = 2N_{ij} )</td>
</tr>
<tr>
<td>3</td>
<td>Sum Entropy</td>
<td>( f_3 = \sum_{i,j} p_{ij}(i) \log [p_{ij}(i)] )</td>
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<td>4</td>
<td>Information Measure of Correlation2 (IMC 2)</td>
<td>( f_4 = \left( 1 - \exp\left[ -2.0(\text{IMC2} - 2) \right] \right)^2 ) where ( \text{IMC2} = -\sum_i p(i,j) \log (p(i,j)) ) and ( HXY = -\sum_i \sum_j p(i,j) \log (p(i,j)) ) and ( HXY2 = -\sum_i \sum_j p(i,j) \log [p(i,j) p(j,i)] )</td>
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and inspected have only two input attributes in order to avoid the curse of dimensionality by using too many features for classification. Figure 1 shows the basic block diagram that illustrates the above discussed methodology.

![Figure 1. Basic block diagram.](image1)

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Euclidean distance is used to obtain the first attribute of the classifier. The Euclidean distance between every pixel, which has been computed individually for the image and also for the individual Red, Green and Blue components using the equation

$$D(z,a) = \sqrt{(z_r - a_r)^2 + (z_g - a_g)^2 + (z_b - a_b)^2}$$

where $D(z,a)$ is the Euclidean distance and the subscripts of $z$ and $a$ represent Red, Green and Blue components.

Any image pixel $z$ is said to be similar to $a$ if the Euclidean distance between them is less than a specified threshold $D_h$ [13]. Thresholding is then carried out to extract the high intensity pixels alone and a binary image is thus generated. The white pixels in the binary image represent the ROI. The area is then computed in terms of the number of white pixels. The block diagram for this method is shown in Figure 2.

![Figure 2. Block diagram for segmentation of ROI.](image2)

The thresholded Blue component of the normal and diseased images showed maximum difference and hence the area of the ROI obtained in terms of the number of pixels for the thresholded Blue component has been used as an attribute for classification.

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![Figure 3. GLCM feature based classification.](image3)

Then, image filtering is carried out using median filters. This is followed by image enhancement using histogram equalization which helps in enhancing the contrast of the image. After pre-processing, the Gray Level Co-occurrence Matrix is computed for all the images. Consider the image data shown below on the left. If we use the position operator "1 pixel to the right and 1 pixel down" then we get the gray-level co-occurrence matrix, where an entry $c_{ij}$ is a count of the number of times that $F(x,y) = i$ and $F(x+1,y+1) = j$. For example, the first entry comes from the fact that 4 times a 0 appears below and to the right of another 0.

$$c = \begin{bmatrix} 4 & 2 & 1 \\ 2 & 3 & 2 \\ 1 & 0 & 2 \\ 0 & 0 & 1 \end{bmatrix}$$

The factor $1/16$ is because there are 16 pairs entering into this matrix, so this normalizes the matrix entries to be estimates of the co-occurrence probabilities [20].

Twenty two features are extracted from this GLCM. Four out of these 22 features are selected using Correlation feature selection algorithm [21] for attribute selection using the CFS-Subset Eval tool in a machine learning software called WEKA [22]. These are Homogeneity, Sum Variance, Sum Entropy and Information Measure of Correlation-2 (IMC 2). The features are computed using the formula shown in Table 2.

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<td>2</td>
<td>Sum Variance</td>
<td>$f_2 = \sum_{i,j} (i-f_1)^2 p_{i,j}(i)$ where $p_{i,j}(k) = \sum \sum_{i,j} p(i,j), i+j=k=2,3,...,2N_i$</td>
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<td>3</td>
<td>Sum Entropy</td>
<td>$f_3 = \sum \sum_{i,j} p_{i,j}(i)\log[p_{i,j}(i)]$</td>
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<td>4</td>
<td>Information Measure of Correlation-2 (IMC 2)</td>
<td>$f_4 = \left(1-\exp[-2.0(HXY2-HXY1)] \right)^2$ where $HXY = \sum \sum_{i,j} p(i,j)\log[p(i,j)]$ and $HXY2 = \sum \sum_{i,j} p(i,j)p_j(\log[p_j(i)]p_j(i))$</td>
</tr>
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In this method, temperature mapping is performed on the pseudo-coloured thermal images. The second attribute for the classifier is obtained by computing the statistical moments for the equivalent temperature mapping. Two sam-

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Table 1. Factor examined for clinical analysis of glaucoma.
scales of ocular thermal images with temperature scale are shown in Figure 4. Figure 5 shows a block diagram for the method based on temperature mapping based classification. A thermal camera usually gives the provision to save the recorded thermal image as temperature data or in an image format such as JPEG. Temperature data enables the use of actual temperature data for analysis. However, the thermal images that have been procured for this project are in JPEG format and lack the actual temperature data possessing only a color coding scheme. The white areas shown in Figure 4 indicate the highest relative temperature regions, followed by red and then yellow, while blue corresponds to the least relative temperature regions in the image. To extract the features, temperature mapping is done, assigning temperature ranges from 20°C to 45°C for 0 to 255 pixel intensities. For a pixel value \( f(x,y) \), the equivalent temperature \( T \) [23], is given by

\[
T(x,y) = \left( \frac{f(x,y)}{255} \times (45 - 20) \right) + 20
\]

A histogram of the image is used to select the high temperature values (> 33°C) from the temperature matrix obtained and then, the mean, standard deviation, skewness and kurtosis [24] are computed for each image. They can be extended to non-linearly separable patterns with the help of functions known as kernels. These hyperplanes are constructed after identifying “Support Vectors” which are the data points that lie closest to the decision surface and are the most difficult to classify. SVMs aim at maximizing the margin around the separating hyperplane.

Five SVMs have been constructed with two attributes each. The first attribute is the area and the second is one of the four selected GLCM features. These SVMs have a linear kernel. The fifth SVM has been constructed with the mean temperature and the area of ROI as its two attributes and the uses a quadratic kernel. The choice of non-linear kernel is to ensure better data fitting. The performance of a binary classification test is evaluated by calculating statistical measures such as sensitivity, specificity, accuracy, and by plotting the Receiver Operating Characteristics curves (ROC) [25]. These are given by the following expressions:

- **Sensitivity** = \( \frac{TN}{TP+FN} \)
- **Specificity** = \( \frac{TN}{FP+TN} \)
- **Fall out** = \( \frac{FP}{TP+FN} \)
- **Accuracy** = \( \frac{TP+TN}{TP+TN+FP+FN} \)
- **Error** = 1 – **Accuracy**

where \( TP = \) Number of True Positives, \( TN = \) Number of True Negatives, \( FP = \) Number of False Positives and \( FN = \) Number of False Negatives.

### 3. Results and Discussion

One hundred ocular thermal images have been captured using a FLIR thermal camera and the protocol for thermal image acquisition was followed while capturing the images [26]. Fifty of these images correspond to normal eyes and fifty correspond to eyes of glaucoma patients. Each category has 25 left and 25 right eye images. Samples of images obtained are shown in Figure 6. The images are in JPEG format and their sizes are reduced to 168 × 224 using a standard window so as to eliminate the unnecessary portions of the images. A uniform temperature scale for the color coding in the images is considered as a fair assumption. The proposed two methods have been implemented using MATLAB R2013a.

Features have been extracted as discussed in Section 2, Table 3 shows a sample of training data set after feature extraction and feature selection. The area of ROI is combined with each of the five other features to construct five 2-attribute SVMs. Eighty-four images have been used to train the system, out of which forty-two are images of normal eyes and forty-two are images of the eyes of glaucoma patients. Each of the classifier’s ability to interpolate has been examined by using this very same training dataset for cross validation. In addition, sixteen mutually exclusive images with known labels, i.e., images that have not been used for training are used for testing each of the classifiers and examining their ability to extrapolate.

The performance of each classifier has been evaluated in terms of accuracy, true positive rate, false positive rate, true negative rate, false negative rate and error. These statistics have been listed out in Table 4. The SVMs have been shown in Figure 7(a) – Figure 7(e). The SVM in Figure 7(a) uses a quadratic kernel for better data fitting. The "0" in the legend is to be interpreted as normal and "1" as diseased. Out of the sixteen images tested, one is falsely classified as positive for glaucoma and three diseased images were wrongly classified as normal. The linear SVM in Figure 7(b) has been constructed using Area and Homogeneity (GLCM Feature) as the input attributes. Here, out of the 16 images that were tested, 15 are correctly classified and only one diseased image was incorrectly classified.

**Table 3. Features of sample training data set.**

<table>
<thead>
<tr>
<th>S. No</th>
<th>Normal/Diseased</th>
<th>Area (pixel count)</th>
<th>Mean Temperature</th>
<th>Homogeneity</th>
<th>Sum Variance</th>
<th>Sum Entropy</th>
<th>IMC 2</th>
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<tbody>
<tr>
<td>1</td>
<td>Normal</td>
<td>181</td>
<td>36.1</td>
<td>0.91</td>
<td>63.15</td>
<td>2.52</td>
<td>0.97</td>
</tr>
<tr>
<td>2</td>
<td>Normal</td>
<td>551</td>
<td>36.12</td>
<td>0.90</td>
<td>62.27</td>
<td>2.54</td>
<td>0.97</td>
</tr>
<tr>
<td>3</td>
<td>Normal</td>
<td>147</td>
<td>36.24</td>
<td>0.90</td>
<td>62.59</td>
<td>2.54</td>
<td>0.97</td>
</tr>
<tr>
<td>4</td>
<td>Diseased</td>
<td>2335</td>
<td>37.11</td>
<td>0.87</td>
<td>61.95</td>
<td>2.55</td>
<td>0.96</td>
</tr>
<tr>
<td>5</td>
<td>Diseased</td>
<td>1661</td>
<td>36.90</td>
<td>0.87</td>
<td>61.86</td>
<td>2.55</td>
<td>0.96</td>
</tr>
<tr>
<td>6</td>
<td>Diseased</td>
<td>1920</td>
<td>37.08</td>
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<td>61.81</td>
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<td>0.96</td>
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</tbody>
</table>
ple of ocular thermal image with temperature scale are shown in Figure 4. Figure 5 shows a block diagram for the method based on temperature mapping based classification. A thermal camera usually gives the provision to save the recorded thermal image as temperature data or in an image format such as JPEG. Temperature data enables the use of actual temperature data for analysis. However, the thermal images that have been procured for this project are in JPEG format and lack the actual temperature data possessing only a color coding scheme. The white areas shown in Figure 4 indicate the highest relative temperature regions, followed by red and then yellow, while blue corresponds to the lowest relative temperature regions in the image. To extract the features, temperature mapping is done, assigning temperature ranges from 20ºC to 45ºC for 0 to 255 pixel intensities. For a pixel value \( f(x,y) \), the equivalent temperature \( T \) [23], is given by

\[
T(x,y) = \left[ \frac{f(x,y) \times (45 - 20)}{255} \right] + 20
\]

A histogram of the image is used to select the high temperature values (> 33ºC) from the temperature matrix obtained and then, the mean, variance, standard deviation, skewness and kurtosis [24] are computed for each image. Of these features, the mean has been found to be more distinguishable between normal and diseased images and hence is selected as one of the classifier attributes. An SVM is then constructed and its performance is evaluated.

2.4. Support Vector Machine Classification

Support Vector Machines (SVM) are supervised learning models based on the idea of an optimal hyperplane for linearly separable patterns. They can be extended to non-linearly separable patterns with the help of functions known as kernels. These hyperplanes are constructed after identifying "Support Vectors" which are the data points that lie closest to the decision surface and are the most difficult to classify. SVMs aim at maximizing the margin around the separating hyperplane.

Five SVMs have been constructed with two attributes each. The first attribute is the area and the second is one of the four selected GLCM features. These SVMs have a linear kernel. The fifth SVM has been constructed with the mean temperature and the area of ROI as its two attributes and the uses a quadratic kernel. The choice of non-linear kernel is to ensure better data fitting. The performance of a binary classification test is evaluated by calculating statistical measures such as sensitivity, specificity, accuracy, and by plotting the Receiver Operating Characteristics curves (ROC) [25]. These are given by the following expressions:

Sensitivity = \( \frac{TN}{TP + FN} \)

Specificity = \( \frac{TN}{FP + TN} \)

Fall out = \( \frac{FP}{TP + FN} \)

Accuracy = \( \frac{TP + TN}{TP + TN + FP + FN} \)

Error = 1 – Accuracy

where \( TP = \) Number of True Positives, \( TN = \) Number of True Negatives, \( FP = \) Number of False Positives and \( FN = \) Number of False Negatives.

3. Results and Discussion

One hundred ocular thermal images have been captured using a FLIR thermal camera and the protocol for thermal image acquisition was followed while capturing the images [26]. Fifty of these images correspond to normal eyes and fifty correspond to eyes of glaucoma patients. Each category has 25 left and 25 right eye images. Samples of images obtained are shown in Figure 6. The images are in JPEG format and their sizes are reduced to 168 × 224 using a standard window so as to eliminate the unnecessary portions of the images. A uniform temperature scale for the color coding in the images is considered as a fair assumption. The proposed two methods have been implemented using MATLAB R2013a.

Features have been extracted as discussed in Section 2. Table 3 shows a sample of training data set after feature extraction and feature selection. The area of ROI is combined with each of the five other features to construct five 2-attribute SVMs. Eighty-four images have been used to train the system, out of which forty-two are images of normal eyes and forty-two are images of the eyes of glaucoma patients. Each of the classifier’s ability to interpolate has been examined by using this very same training data-set for cross validation. In addition, sixteen mutually exclusive images with known labels, i.e. images that have not been used for training are used for testing each of the classifiers and examining their ability to extrapolate.

The performance of each classifier has been evaluated in terms of accuracy, true positive rate, false positive rate, true negative rate, false negative rate and error. These statistics have been listed out in Table 4. The SVMs have been shown in Figure 7(a) – Figure 7(e). The SVM in Figure 7(a) uses a quadratic kernel for better data fitting. The “0” in the legend is to be interpreted as normal and “1” as diseased. Out of the sixteen images tested, one is falsely classified as positive for glaucoma and three diseased images were wrongly classified as normal. The linear SVM in Figure 7(b) has been constructed using Area and Homogeneity (GLCM Feature) as the input attributes. Here, out of the 16 images that were tested, 15 are correctly classified and only one diseased image was incorrectly

<table>
<thead>
<tr>
<th>S. No</th>
<th>Normal/ Diseased</th>
<th>Area (pixel count)</th>
<th>Mean Temperature</th>
<th>Homogeneity</th>
<th>Sum Variance</th>
<th>Sum Entropy</th>
<th>IMC 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Normal</td>
<td>181</td>
<td>36.1</td>
<td>0.91</td>
<td>63.15</td>
<td>2.52</td>
<td>0.97</td>
</tr>
<tr>
<td>2</td>
<td>Normal</td>
<td>551</td>
<td>36.12</td>
<td>0.90</td>
<td>62.27</td>
<td>2.54</td>
<td>0.97</td>
</tr>
<tr>
<td>3</td>
<td>Normal</td>
<td>147</td>
<td>36.24</td>
<td>0.90</td>
<td>62.59</td>
<td>2.54</td>
<td>0.97</td>
</tr>
<tr>
<td>4</td>
<td>Diseased</td>
<td>2335</td>
<td>37.11</td>
<td>0.87</td>
<td>61.95</td>
<td>2.55</td>
<td>0.96</td>
</tr>
<tr>
<td>5</td>
<td>Diseased</td>
<td>1661</td>
<td>36.90</td>
<td>0.87</td>
<td>61.86</td>
<td>2.55</td>
<td>0.96</td>
</tr>
<tr>
<td>6</td>
<td>Diseased</td>
<td>1920</td>
<td>37.08</td>
<td>0.87</td>
<td>61.81</td>
<td>2.58</td>
<td>0.96</td>
</tr>
</tbody>
</table>
classified as normal. Similarly, Figures 7(c) –7(e) depict linear SVMs with good interpolation and extrapolation capabilities.

4. Conclusion

In this paper, we have presented a non-invasive procedure to evaluate the presence of glaucoma, the eye disease. Two methods have been proposed for the detection of glaucoma using ocular thermal images analyzed in fifty normal and fifty abnormal subjects. The performance of the proposed methods using SVM classifiers with features obtained from Gray Level Co-occurrence Matrix and the temperature mapping has been evaluated. The simulation results indicate that the classifier in the detection of glaucoma performs well and provides accuracy, sensitivity and specificity of around 96%.

However, extensive clinical trials may further be needed to establish the practical acceptance of our method by medical community.

Table 4. Performance measures of the classifier.

<table>
<thead>
<tr>
<th>SVM No</th>
<th>Attributes</th>
<th>Total Accuracy (out of 1)</th>
<th>TPR (Sensitivity) (out of 1)</th>
<th>FPR (False Alarm) (out of 1)</th>
<th>FNR (Miss rate) (out of 1)</th>
<th>TNR (Specificity) (out of 1)</th>
<th>Error (out of 1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Area + Mean Temp</td>
<td>0.89</td>
<td>0.88</td>
<td>0.12</td>
<td>0.9</td>
<td>0.11</td>
<td></td>
</tr>
<tr>
<td></td>
<td>GLCM Feature based SVMs</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Area + Homogeneity</td>
<td>0.96</td>
<td>0.96</td>
<td>0.04</td>
<td>0.04</td>
<td>0.04</td>
<td>0.04</td>
</tr>
<tr>
<td>3</td>
<td>Area + Sum Variance</td>
<td>0.93</td>
<td>0.9</td>
<td>0.04</td>
<td>0.1</td>
<td>0.96</td>
<td>0.07</td>
</tr>
<tr>
<td>4</td>
<td>Area + Sum Entropy</td>
<td>0.95</td>
<td>0.94</td>
<td>0.04</td>
<td>0.06</td>
<td>0.96</td>
<td>0.05</td>
</tr>
<tr>
<td>5</td>
<td>Area + IMC2</td>
<td>0.96</td>
<td>0.96</td>
<td>0.04</td>
<td>0.04</td>
<td>0.96</td>
<td>0.04</td>
</tr>
</tbody>
</table>

5. Acknowledgement

The authors would like to thank IGCAR, India for the thermal images. Furthermore, the authors would like to thank the management of Sri Sivakumaran Nadar College of Engineering, Kalavakkam, for providing the facilities to carry out this work.

References


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However, extensive clinical trials may further be needed to establish the practical acceptance of our method by medical community.

4. Conclusion

In this paper, we have presented a non-invasive procedure to evaluate the presence of glaucoma, the eye disease. Two methods have been proposed for the detection of glaucoma using ocular thermal images analyzed in fifty normal and fifty abnormal subjects. The performance of the proposed methods using SVM classifiers with features obtained from Gray Level Co-occurrence Matrix and the temperature mapping has been evaluated. The simulation results indicate that the classifier in the detection of glaucoma performs well and provides accuracy, sensitivity and specificity of around 96%.

![Figure 7 (a). SVM with area and mean temperature as attributes.](image1)

![Figure 7 (b). SVM with area and homogeneity as attributes.](image2)

![Figure 7 (c). SVM with area and sum variance as attributes.](image3)

![Figure 7 (d). SVM with area and IMC2 as attributes.](image4)

![Figure 7 (e). SVM with area and sum entropy as attributes.](image5)

Table 4. Performance measures of the classifier.

<table>
<thead>
<tr>
<th>SVM No</th>
<th>Attributes</th>
<th>Total Accuracy</th>
<th>TPR (Sensitivity)</th>
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<td>2</td>
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<td>0.96</td>
<td>0.04</td>
<td>0.04</td>
<td>0.96</td>
<td>0.04</td>
</tr>
<tr>
<td>3</td>
<td>Area + Sum Variance</td>
<td>0.93</td>
<td>0.9</td>
<td>0.04</td>
<td>0.01</td>
<td>0.96</td>
<td>0.07</td>
</tr>
<tr>
<td>4</td>
<td>Area + Sum Entropy</td>
<td>0.95</td>
<td>0.94</td>
<td>0.04</td>
<td>0.06</td>
<td>0.96</td>
<td>0.05</td>
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<tr>
<td>5</td>
<td>Area + IMC2</td>
<td>0.96</td>
<td>0.96</td>
<td>0.04</td>
<td>0.04</td>
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