Evaluation of a System for Automatic Detection of Diabetic Retinopathy from Color Fundus Photographs for Screening Population

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ABSTRACT
Diabetic Retinopathy is an eye disease and a common complication of diabetes that can cause vision loss if left undiagnosed at the initial stage. It is the prime cause of blindness in the working age population of the world. Colour fundus images of eyes are used by ophthalmologists to study eye diseases like diabetic retinopathy. Detection method is proposed to detect dark or red lesions such as microaneurysms and hemorrhages in fundus images. This method comprises of three stages. First, the green channel of the colour retinal image is preprocessed using polynomial contrast enhancement. Second, the candidate dark lesion objects are extracted from the contrast enhanced fundus image. A novel method is developed to extract candidate dark lesions based on matched filtering and local relative entropy thresholding. The performance of this technique is compared to mathematical morphology based dark lesion detection method. A hybrid method that combines both detection schemes is also tested for better segmentation. The final stage classifies true dark lesions from dark non-lesions using knn and svm classifiers. For image based evaluation, this method has resulted with sensitivity, specificity and accuracy of 94.86% and 95.6% 95.38% respectively.

Keywords
Biomedical image processing, retinal Fundus images, microaneurysm detection, morphological operator.

1. INTRODUCTION
We Eye is a very essential and critical organ of the human body which only gives vision. There are huge eye diseases increasing now days due to improper care. Diabetic Retinopathy (DR) is severe and wide diffusion diseases. It has been identified as one of the cause for blindness or vision impairment[1]. Diabetic retinopathy (DR) is a vascular disease of the retina which affects patients with the diabetes. It is the number one cause of blindness in people between the ages of 20-64 in the United States. Early DR detection and timely treatment have been shown to prevent visual loss and blindness in patients with retinal difficulties of diabetes[2]. In the next decade, projections for the United States are that the average age will rise, the number of people with diabetes in each age category will increases, and there will be a restriction of capable eye care providers [5]. The healthcare trends will challenge the public health capacity to care for both patients with DR and people with diabetes at risk for this difficulty. If the previous situation plays out, it will be necessary to either screen large numbers of people with diabetes for DR, ration access to eye care, or both.

The automated algorithms introduced above are improved to recommend recommendation for a patient with any form of DR to an ophthalmologist, and they were enhanced to detect early DR, because in our view, this is the burden[3]. They could be modified to diagnose vision threatening DR, detect those patients with important non proliferative DR, or proliferative DR[4]. This previews but not newly evaluated early detection programs. In addition, the processes were limited to detection of that called microaneurysms and small hemorrhages , to make the comparison though we have previously designed and evaluated systems that also detect exudates and cotton-wool spots[7]. The automated algorithms are thus optimized to recommend transfer for a patient with any form of DR to an ophthalmologist [6].

The retina is a thin multi-layered sensory tissue that covers the inside wall at the back side part of eye is called fundus. It is covered number of. These photo receptors are responsible for receiving light rays, converting them into electrical impulses and then transmitting these signal to the brain where they are convert into images. Figure 1 shows a typical normal retinal fundus image with highlighted regions optic disc, fovea, blood macula and vessels. The outlying parts of the retina are responsible for outlying vision while the central area, that is called macula, is in charge of central vision that allows us to see details and perform tasks that require central vision, like was a reading. The circular area in the central region is a macula and that the retina measuring about 5 mm to 6 mm in diameter. A small gloom in the center of macula measuring about 1.5 mm in diameter is that is called fovea.
Diabetic retinopathy, the most major causes of common diabetic eye disease blindness is occurs when in the retina blood vessels change. Sometimes these vessels swell and leak fluid or block. In some cases, abnormal new blood vessels grow on the surface of the retina its last stage of DR. It is classified into two category first is Non-Proliferative Diabetic Retinopathy (NPDR) and Proliferative Diabetic Retinopathy (PDR).

1.2 Non proliferative diabetic retinopathy

Non proliferative diabetic retinopathy is the most common and may arise at the point in time after the diabetes. Ophthalmologists detect these changes by examining the patient's retina and look for spots of bleeding or swelling areas of retinal. Identification and recording of the following abnormalities (will aid in the accurate assessment of retinopathy severity.

Microaneurysms

The primary abnormality that occurs in the eye due to DR is the Microaneurysm. These are identified as tiny, dark red spots or miniscule haemorrhages, either appearing alone or in clusters, inherent to the light sensitive retina. The Microaneurysm is circular in shape and the size varies from 10-100 microns i.e. 1/12th the diameter of an average optics disc. The disease is not alarming at this phase.

Hemorrhages

These are also termed ‘blot’ microaneurysm, with concern to their round shape. These are found in the deeper layers of the retina. Intra retinal hemorrhages appear when capillaries or microaneurysms rupture and some blood leaks out of these vessels. In Figure 1.3 hemorrhages can be seen as red flame shaped regions.

Hard Exudates

Hard exudates represent leak of fluid that is rich in fat and protein from surrounding capillaries and microaneurysms within the retina. That is main features of diabetic retinopathy and appear as random yellowish patches of changing sizes, and shapes.

Soft Exudates

These are often called cotton wool spots and often seen in advanced retinopathy. These abnormalities usually appear as small round in the retina with a yellowish colour, usually adjacent to an area of hemorrhage. Cotton wool spots come to the swelling of the surface layer of the retina in the non-appearance of normal blood flow through the retinal blood vessels. The nerve fibers are injured and that resulting in swelling and appearance of a cotton wool spot [2].

1.3 Proliferative diabetic retinopathy

Proliferative diabetic retinopathy is the advanced stage retinopathy develops in more than 50 percent cases after about 25 years of onset of the disease. Therefore, it is more common in patients [9]. The hallmark of PDR is the growth of new blood vessels in the areas where normal capillaries have already closed. These new blood vessels are abnormal and fragile. They grow along the retina and along the surface of the clear, vitreous gel that fills the inside of the eye. By themselves, these blood vessels do not cause symptoms or vision loss. However, they have thin, fragile walls that leak blood resulting in severe vision loss and even blindness can be the end result.

2. METHODS

Fundus imaging has a main role in diabetic retinopathy monitoring since incidences of retinal abnormalities are common and consequences serious. However, since the eye fundus seems to be sensitive to vascular diseases, fundus imaging is considered as a candidate for non-invasive screening of diabetes. The details of these databases are forming the ground-truth of each image, which is also available in the DIARETDB0. However, one problem presented in this ground-truth is the low accuracy in the marking of lesions, since ophthalmologists delimited areas with a set of lesions and not each lesion itself. In this new ground-truth, among the 130 images, 109 images were defined with lesions and 21 images were defined no lesions. The technique for automatic detection of microaneurysms and hemorrhages proposed in this paper is based on mathematical morphology four stages. At the first stage, a preprocessing of the fundus images is performed in order to improve the contrast of these images. At the second stage, an improvement of the low intensity structures (blood vessels, fovea and red lesions) is performed. The blood vessels are detected by the sum of morphological operation using linear structuring elements with different angles in third stage. At the fourth stage, occurs the deduction of the blood vessels from the image obtained at the second stage. Each stage is explained in details in the following sections.

2.1 Pre-processing

Pre-processing is to reduce this effect by enhancing the contrast and normalizing the mean intensity. The objective of preprocessing is to attenuate the noise, to improve the contrast and to correct the non-uniform illumination. The colour retinal images taken at screening programs are often poorly contrasted and contain artifacts. To correct uneven illumination of images, a morphological top-hat operator with disk-shaped structuring element and fixed radius of 25 pixels was applied to the green component of the colour image. To reduce noise, a 3x3 median filter is applied to the shade corrected image [4].

![Image](https://example.com/image1.png)

<table>
<thead>
<tr>
<th>(a)</th>
<th>(b)</th>
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Fig. 2 – An image before and after the pre-processing operations. (a) green channel image, (b) pre-processed image.

2.2 Microaneurysm Detection

The extraction of microaneurysms step results in a set of candidate microaneurysm objects. The objective of the microaneurysm candidate classification system is to classify the objects as either microaneurysms or dark non-lesions. Hence to discard dark non-lesion candidates, classifiers are used which are trained with the features derived from the candidates. The best classification requires good features as well as good classifier.

\[ G_2 = \psi^{\frac{1}{2}} (G_1) \]
The image resulting from the local variation operator is denoted by G3 as below:

\[ G_3(x) = \frac{1}{N-1} \sum_{i \in W(x)} (G_2(i) - \mu(x))^2 \]

where \( x \) is a set of all pixels in a sub-window \( w(x) \), \( N \) is the number of pixels in \( w(x) \), \( \mu(x) \) is the mean value of \( G_2(i) \) and \( i \in w(x) \). The selection of window size is relied on the preferred compromise between the sensitivity and precision performance measures. So as the window size is larger the small exudates are more difficult to be detected leading to low sensitivity. Based on the experimental tests, we found that a window size of \( 9 \times 9 \) gives good results.

3. RESULTS AND DISCUSSION

The proposed detection methods are tested and evaluated on DIARETDB0 [13], a publicly available database of colored fundus images and corresponding ground truth images. The entire process of segmenting vessels was performed on Intel PC with 1.66 GHz CPU and 512MB memory using Matlab 2012. The processing of each image including difficulty and thresholding took about 30 seconds. The green component, of the RGB fundus image, was chosen to obtain the microaneurysms. Similar to the exudates detection algorithm, first the prominent structures within retina images, Such as blood vessel tree and optic disc are to be removed. After that a sophisticated sequence of image processing algorithms was used to determine the areas within the fundus images to get microaneurysms [14] and exudates. It is important to have some balance between the values of sensitivity and specificity, parameters such as True Positive (TP), True Negative (TN), False Positive (FP) and False Negative (FN).

TP is the number of abnormal images classified as abnormal by the screening system. TN is the number of images that are really normal and classified as normal by the screening procedure. FP is the number of normal images that are predicted to be abnormal and FN is the number of abnormal images, classified by the procedure to be normal. Using these parameters, Sensitivity, Specificity and accuracy are calculated. For each of the images a corresponding manually segmented image is provided. The instruction of an ophthalmologist are coloured white. Quantitative evaluation of the segmentation algorithm is done by comparing the output image with the corresponding manually segmented image. In the classification stage, a training segmented dark non-lesion objects and dark lesion Table 1. Performance Comparison of Mathematical Morphology Detection. False positives are pixels marked as vessel by the method, but that are actually negatives in the ground truth. True negatives are pixels marked as background in both images. The performance of the method was tested on images with incidence of lesions provided by the DIARETDB1. The accounting of the pixels in TP, FN, FP and TN was performed for each resulting image of the method. Table 1 shows the average values of TP, FN, FP and TN for each fundus image.

![Fig. 3(a) shows the green channel of a fundus image](image1)

![Fig. 3(b) shows the detected microaneurysms](image2)

Lesion based evaluation and image based evaluation are employed to measure the accuracy of the proposed detection method at the pixel level. The recommend that any process used for screening sight threatening diabetic retinopathy should have at least 96.9% sensitivity and 96.1% specificity; Sensitivity means the percentage of abnormal fundus classified as abnormal by the procedure. Specificity means the percentage of normal funduses classified as normal by the procedure. The higher the sensitivity and specificity values, best the procedure.

Figure 4. (a) Green channel of the fundus image. (b) Ground truth image. (c) Microaneurysms and hemorrhages detected. (d) True positive pixels. (e) False negative pixels. (f) False positive pixels. (g) True negative pixels.

<table>
<thead>
<tr>
<th>Databases</th>
<th>Total images</th>
<th>Non-exudates</th>
<th>No. of images with</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DIARETDB0</td>
<td>130</td>
<td>58</td>
<td>72</td>
<td>96.9%</td>
<td>96.1%</td>
</tr>
</tbody>
</table>

Table 2 Comparison of Performance with other technique

<table>
<thead>
<tr>
<th>Authors</th>
<th>Method</th>
<th>Accuracy of classification</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wang et al.</td>
<td>Minimum distance discriminant classifier</td>
<td>70%</td>
<td>83%</td>
<td>89%</td>
</tr>
</tbody>
</table>
Vujosevic et al. 2009

<table>
<thead>
<tr>
<th>Proposed Method</th>
<th>Single lesions</th>
<th>Blood vessel, exudates, microaneurysms</th>
<th>Support Vector Machine classifier</th>
</tr>
</thead>
<tbody>
<tr>
<td>[6]</td>
<td>Not reported</td>
<td>86%</td>
<td>95.38%</td>
</tr>
<tr>
<td>[7]</td>
<td></td>
<td>82%</td>
<td>95.6%</td>
</tr>
<tr>
<td>[9]</td>
<td></td>
<td>86%</td>
<td>94.86%</td>
</tr>
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</table>

The proposed method achieved satisfactory results for the red lesions that were detected. These structures are mainly caused by remains of blood vessels, where there was ineptitude of the method to eliminate some vessels without eliminating lesions close to them. One factor that affected the sensitivity was not classifying some microaneurysms occupant area very small. Some of these lesions were confused with noise and were improperly removed due to its very small size, it may also be mentioned that the un-label brightness of some images generated false positive results, which reduced the specificity of the method.

4. CONCLUSION

The proposed method is adapted to deal with different types and qualities of images through taking all image information into account. The proposed method performance of the measured against clinician hand-labelled images. Comparing with some recent related works, the proposed method indicates an improvement in the specificity and accuracy measures and reasonable sensitivity particularly with the high performance method in. The performance results, demonstrated in this work, indicate that automated processing methods that are based on split-and-merge algorithm can give very competitive results in exudate detection. The strength of the proposed method comes from using computed parameters based on analysis of the image. A limitation in our work is that it occasionally fails to exclude some non-exudate objects particularly those that have similar features to real exudates.

5. REFERENCES


