Application of an Image Processing Technique for Early Diagnosis and Monitoring of Glaucoma

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Abstrak


Abstract

Early stage glaucoma can be diagnosed by finding retinal nerve fiber defects using a color image processing technique. This method is based on the reflectance of the defective part which has low values in the green and blue wavelengths. The level pixel (picture element) of reflectance is scored from 0 to 255. In early stage glaucoma, the difference of that reflectance is very small. This new method detects that small difference. The method consists of sampling and summation value of pixels in green and blue along the radial lines drawn between two circles whose centers are the same and located at the center of the eye. It was found that the small differences can be detected only by doctors with a sufficient experience. This method is better than the color enhancement. Also, the detection of cup and blood vessels has been shown to assist in automatic processing. It will be necessary to apply this method to many patients to check the of the utility of the method and develop automatic processing system in the future.

Keywords: Image Processing, glaucoma, early diagnosis

INTRODUCTION

The result of eye morbidity survey conducted by Research and Development Institution, Ministry of Health, Republic of Indonesia in 1982 showed that the prevalence of blindness in Indonesia is about 1.2%. There are about 2 million blindness cases in Indonesia among the population of 180 million (1991). Since blindness is not only a health problem, but also a social problem, the Indonesian Minister of Health stated in the official document No.19/Birhup/67 that blindness is a national "disaster". Blindness, or even vision loss before total blindness occurs, causes reductions in working hours, working capacity and working opportunity. Blindness causes reduction in productivity, which may be equal to billions rupiah per year. One of the important causes of blindness in Indonesia is glaucoma.

Glaucoma occurs when the inner pressure of the eye ball increases. In the early stage of glaucoma, some clinical signs can be seen around the optic nerve head and in the retinal nerve fiber layer. The changes in the cup-disc ratio, the change of height in the rim of cup and optic nerve head pallor occurs around the optic nerve head because of the increased pressure. To measure these phenomenon automatically, three
A method for detecting small changes in the retina has been proposed and tried. This procedure involves sampling according to radial lines on a color retinal photograph and summing these data. We also report the experimental results using color retinal photographs of normal and glaucoma eyes.

METHODS

The method consists of cup detection, radial projection, and blood vessel detection. The cup detection is used for deciding the center of the circles for the following processing. The radial projection is the main method for detecting retinal nerve fiber defects. Blood vessel detection is used to classify the vessels and the retinal nerve fiber defect.

1. Radial Projection

The difference between defective part and normal parts of retinal nerve fiber layer is detected as the difference of density in blue and green images of color retina photographs. The quantity of the density difference is very small in the early stage of glaucoma. Here, a new method is proposed to detect this small difference. The ideal method consists of sampling the density data according to each nerve fiber bundle as shown in Figure 1, summation and comparing the summed data for each nerve fiber bundle. By this method, small difference is enlarged and can be seen as big difference. Even if noise is added in images, noise is averaged and has no effect in the final summed data.

But it is not so clear where each fiber bundle runs. It is clear that each fiber bundle starts radial from the rim of the cup. So, the real method proposed here uses the radial sampling around the cup as shown in Figure 2. It may be called the radial projection method. In the computational method, two circles whose centers are the same and are in the cup are drown outside of the cup. Many radial lines are drown from the small circle to the large circle. The data is sampled according to each radial line.
The method is explained using Figure 3. Sampling according to the straight lines in the area with the pixel value of sector 100 and 99 are shown in Figure 3. And the value of summation of data is shown in Figure 4.

![Figure 3. Projection along a line](image)

![Figure 4. Projected data along each line](image)

2. Cup Detection
The cup is detected by finding the brightest part in retinal color photographs. The process is shown in Figure 5. The retinal image is sampled roughly, for example, in ten point intervals to find the brightest point. This point is found in the cup area that looks white in the image. But the position is not the center of the cup.

The four points denoted a, b, c, and d that have large variation in intensity are searched along horizontal and vertical lines drawn from the brightest point A shown in Figure 5. The center of the first circle shown as C is decided by using the 4 points. The first circle is drawn a little bit outside the 4 points. The radius of second large circle is determined by the position of the cup and the image periphery. If the position of the cup is the upper left side shown in Figure 5, the radius of the second circle shown as R2 is the distance between the center of the cup and the periphery of the image.

![Figure 5. The method for automatic detection of the cup. The retinal image is resampled roughly to find the brightest point A. The four point a, b, c, and d have large variation of the intensity. C is the center of the first circle. R2 is the radius of the second circle.](image)

3. Blood Vessel Detection
The blood vessels look red and have lower intensity in green and blue wavelength on images as they run from the cup. Previous methods classify the blood vessels in colored spaces, and are only useful when the retina is uniformly illuminated.\(^1\)\(^6\)\(^7\) By this projection and sampling method, the uniformly illuminated photographs are unnecessary because each pixel (picture element) value of image will be averaged and noises on photograph will be lost.

The new-method that uses around the cup in a green image is shown in Fig.6a and is plotted on a graph in Figure 6b. In Figure 6b, the blood vessel is detected as the dark part or lower value pixel in green or blue wavelength. Finally, in the figures of radial projection (Figures 7-11) the defective part can be detected as the part with lower value in the slow continues change of the summed value due to the lack of flatness in illumination. And also, the blood vessel parts can be detected as the part with sharp decreasing in the same figure. But there two parts are clearly classified by using the information of blood vessel detection.
Figure 6. Blood vessel parts extracted by the method of circle resampling. (a) shows the resampling along the circle. (b) shows the result of resampling and classification using threshold.

RESULTS

Color photographics of retina nerve fiber of glaucoma eyes were used for this experiment. These color photographics were separated into red, green, and blue images with quantity level of 1 to 255 by using a scanner with the aperture of 100×100 micrometer. The green and the blue images were processed by using personal computer NEC98-01 with image buffer and C-Language. Concentric circles are drawn automatically after detecting the center of the cup and the image periphery. Also, the radial lines were automatically drawn. The data is projected along each radial line. The processing time was about 10 seconds.

In Figure 7 to Figure 11, (a) shows the projection lines on the retina images and (b) shows the summed data on each line. In the Figure 7a, the wedge shaped defect can be seen clearly in above and below the cup. In the Figure 7b, the upper graph shows the summation in blue and the lower graph shows the summation in green. From Figure 7b, we find that the values of summation from the line number of 0 to 140 and 300 to 370 are small. Of course, these two parts correspond with the defective part shown in Figure a. The sharp decreasing part painted red at the line number of 50, 150 and 390 is due to the effect of blood vessels.

In Figure 8a, a wedge shaped defect is seen from the upper right side to the lower left side in the center. In Figure 8b, the line number of this defect is 200 to 270. This part looks like a mountain because the illumination was not flat. If the defect were not there, the shape would be a perfect hemisphere. In Figure 9a, it is difficult to find the defective part. At the line number of 100, sharp decreases are seen because of blood vessels noted in the lower graph of Figure 9b. The defective part cannot be found from the summed data in green. At the line number of 150 to 230 and 300 to 350, summed data have low values in the upper graph. This part can be diagnosed as defective part by an experienced physician.

In the Figure 10a, two defective parts are detected. One is clearly observed at line number 100 to 150 in the retinal photograph and in the radial projection map shown in Figure 10b. The other is detected at line number 300 in the radial projection map, but it is difficult to find it in the retinal photography due to its small difference. This part is also detected as the low value at line number 250 in Figure 11b.

Through these experiments, it was found that small differences between nerve fiber defects and the normal parts can be detected, even if the illumination is not flat. If the defective part and the marked changes in blood vessels can be separated, the diagnosis will become easy.

CONCLUSIONS

The radial projection method is proposed to detect the nerve fiber defect in the early stage of glaucoma. The method depends on the fact that the defect has a lower reflectance compared to the normal part. The method consists of sampling and summation along radiating lines drawn between two circles whose centers are the same and located at the center of the cup of eye. Through experiments, it was found that the small difference of the defective and normal parts can be detected only by physicians with experience.

Sometimes, the problem in photography technique and processing is to make the homogen or flat illumination to the object. For these purpose we make compensation illumination. The bad or unflat illuminous will change intensity of the color, and finally will cause misdiagnosis. The compensation of illumination is needed to reduce the error in interpretation of photograph. Therefore this method does not need compensation of illumination. The detection of cup and blood vessels have been shown to assist the automatic processing.
Figure 7. Experimental result. (a) shows the radial projection on the retinal image. The original image is color photograph. The wedge shaped defect is seen clearly in the up and down direction from the cup. (b) the upper graph shows the projection in blue and the lower graph shows the projection in green.

Figure 8. Experimental result. (a) shows the radial projection on the retinal image. The wedge shaped defect is seen from the upper right side to the lower left side in the center. (b) the upper graph shows the projection in blue and the lower graph shows the projection in green.

Figure 9. Experimental result. (a) shows the radial projection on the retinal image. This is the difficult example to find the defect part. (b) the upper graph shows the projection in blue and the lower graph shows the projection in green.
Figure 10. Experimental result. (a) shows the radial projection on the retinal image. (b) the upper graph shows the projection in blue and the lower graph shows the projection in green.

Figure 11. Experimental result. (a) shows the radial projection on the retinal image. (b) the upper graph shows the projection in blue and the lower graph shows the projection in green.

As it is mentioned before, the scanner was used in this experiment to replace the video-camera, due to the budget limitation. This limited the application of this method to the patients. We will apply this method to many patients to check the usefulness of the method and we will develop full automatic processing in the future. Advanced research and experiments are needed to apply and develop the method.

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