Automated detection of dilated capillaries on optical coherence tomography angiography

CHANGLEI DONGYE,1,2 MIAO ZHANG,1,3 THOMAS S. HWANG,1 JIE WANG,1
SIMON S. GAO,1 LIANG LIU,1 DAVID HUANG,1 DAVID J. WILSON1 and YALI JIA1,

1Casey Eye Institute, Oregon Health & Science University, Portland, OR 97239, USA
2College of Information Science and Engineering, Shandong University of Science and Technology,
Qingdao, 266590, China
3OptoVue, Inc., 2800 Bayview Dr, Fremont, CA 94538, USA

jiaya@ohsu.edu

Abstract: Automated detection and grading of angiographic high-risk features in diabetic retinopathy can potentially enhance screening and clinical care. We have previously identified capillary dilation in angiograms of the deep plexus in optical coherence tomography angiography as a feature associated with severe diabetic retinopathy. In this study, we present an automated algorithm that uses hybrid contrast to distinguish angiograms with dilated capillaries from healthy controls and then applies saliency measurement to map the extent of the dilated capillary networks. The proposed algorithm agreed well with human grading.

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1. Introduction

Optical coherence tomography angiography (OCTA) produces 3-dimensional (3D) angiograms of the human retina with capillary-level resolution by measuring signal decorrelation or variance caused by red blood cells in consecutive cross-sectional images (B-frames) taken at the same location [1–5]. Unlike fluorescein angiography, the standard for ocular angiography, OCTA is noninvasive, rapidly acquired, 3D, and amenable to automated analysis.

Projection artifacts, however, which are caused by moving blood cells in the more superficial vessels, hinder the clear visualization of deeper plexuses. This has limited the ability of conventional OCTA to evaluate retinal vasculature in deeper plexuses. The depth-resolution of OCTA has recently been improved with the projection-resolved (PR) OCTA algorithm, which produces clearer images of deeper retinal plexuses by resolving the ambiguity between real flow signal and projection artifacts [6].

In diabetic retinopathy (DR), the ability to observe individual capillary networks may aid in the diagnosis and treatment of the disease [7]. Automation of this feature could enhance clinical practice and be useful for screening treatment-threshold DR.
In this study, we present and validate an automated detection algorithm based on contrast and saliency methods to distinguish dilated capillaries from normal capillaries, and then quantify the extent of dilated capillaries on deep capillary plexus (DCP) angiograms.

2. Methods

2.1 OCTA data acquisition and processing

Participants were recruited at the Casey Eye Institute of the Oregon Health & Science University (OHSU) in accordance to an OHSU Institutional Review Board approved protocol. This study complied with the tenets of the Declaration of Helsinki. Written informed consent was obtained from all participants after explanation of the nature and possible consequences of the study.

Two 3 × 3 mm scans (1 x-fast and 1 y-fast) with 2 mm depth were sequentially obtained in one eye of each participant within a visit using a commercial spectral domain OCT system (RTVue-XR; Optovue, Fremont, CA) with a center wavelength 840 nm, a full-width half maximum bandwidth of 45 nm, and an axial scan rate of 70 kHz [1]. In the fast transverse scanning direction, 304 axial scans were sampled to obtain a single 3 mm B-scan. Two repeated B-scans were captured at a fixed position before proceeding to the next location. A total of 304 locations along a 3 mm distance in the slow transverse direction were sampled to form a 3D data cube. All 608 B-scans in each data cube were acquired in 2.9 seconds.

Flow signal was detected by using the split-spectrum amplitude-decorrelation angiography (SSADA) algorithm [10, 11]. Motion artifacts were removed by aligning and merging the x and y-fast scan volumes [2, 12]. Projection artifacts were suppressed by the projection-resolved OCTA (PR-OCTA) algorithm [6]. A directional graph search algorithm identified structural boundaries [13]. The DCP angiograms were created by the en face projection of the OCTA slab which contained the outer 50% of inner nuclear layer (INL) and whole outer plexiform layer with a minimum thickness of 37 microns.

2.2 Algorithm overview

![Fig. 1. Overview of the developed dilated capillaries detection algorithm. DCP: deep capillary plexus.](image)

The algorithm developed in this work for detection of dilated capillaries is composed of four major steps: vesselness-based denoising, hybrid-contrast-based diagnosis, contrast-and-saliency-based detection, and connectivity-based vessel grading. (Fig. 1)

First, the background noise is removed using a vesselness filter. Next, a hybrid contrast ratio map that combines global and local contrast is generated, differentiating DR eyes with dilated capillaries from normal eyes. Then the DCP angiograms of DR eyes are enhanced by Gabor filter and regions with high saliency and contrast are recognized as potential dilated capillaries region. Finally, connected capillary networks that reside on the potential dilated
capillaries region are graded as dilated capillaries. The following sections describe the above steps in detail.

2.3 Vesselness-based denoising

Vesselness filters have been proven to be successful in enhancing vascular structures in OCTA [14–16]. We have applied it here to remove the background noise. We first measured vesselness, or the likelihood of a pixel belonging to a vessel, by assessing how “tubular” the structure in its vicinity is [17, 18]. For each pixel, the anisotropy ($R_b$) and structure ($S$) were calculated from the eigenvalues ($\lambda_1 \leq \lambda_2$) of its Hessian matrix as:

$$R_b = \frac{|\lambda_1|}{|\lambda_2|}$$  \hspace{1cm} (1)

$$S = \sqrt{\lambda_1^2 + \lambda_2^2}$$  \hspace{1cm} (2)

Then, the vesselness $V$ was calculated by:

$$V = e^{\frac{r_s}{\beta_1}} (1 - e^{\frac{s}{\beta_2}})$$  \hspace{1cm} (3)

Where $\beta_1 = 1$ pixel and $\beta_2 = 10$ pixels are sensitivity constants. Vesselness is low if there is a low intensity change in the vicinity of a pixel (no vessel structure) or the intensity change is large in almost all directions (large anisotropy). Pixels with vesselness less than Otsu’s threshold [19] were determined as noise and removed from the DCP angiogram.

2.4 Hybrid-contrast based diagnosis

Previous experiments of OCTA on microfluidic flow phantoms demonstrated that the decorrelation values were related to both blood flow velocity and channel width [8]. If the velocity is held constant, the flow signal is solely dependent on channel width. In the DCP, which is ordinarily composed of a uniform network of capillaries [9], we assume that abnormally high flow signal is due to the increased channel width. We can then identify dilated capillaries by the flow signal without having to measure the vessel caliber [20], which is difficult to measure given the current transverse resolution of OCTA (15-20µm). In this step, we combined two contrast analyses to identify angiograms with dilated capillaries and locate potential regions with dilated capillaries.

2.4.1 Global contrast

The global contrast [21] preserves the distinctiveness of flow signal intensity by measuring the ratio between the intensity of a given pixel and the average intensity of the angiogram.

$$GC(x, y) = \frac{I(x, y)}{\text{mean}(I)}$$  \hspace{1cm} (4)

where $GC$ is the global contrast matrix; $I$ is the signal intensity on DCP angiogram.

2.4.2 Local contrast based on center-surround mechanism

The local contrast investigates the rarity of an image region with respect to its immediate vicinity [22]. The most obvious dilated capillaries have high local contrast. We defined the local contrast:

$$LC(x, y) = \frac{w^r(x, y)}{w^l(x, y)}$$  \hspace{1cm} (5)
where $w^c$ is the average intensity within a central circle at $(x, y)$ with a diameter of 60 µm, while $w^s$ is the average intensity of the surrounding ring with an inner diameter of 60 µm and outer diameter of 130 µm.

2.4.3 Hybrid contrast

When applied alone, global contrast frequently detects normal branching capillaries on control eyes as dilated capillaries. Local contrast, when applied alone, detects isolated capillary segments as dilated capillaries. The hybrid contrast ratio, which is the product of global and local contrast ratios, can avoid these false positive errors.

2.4.4 Differentiating dilated capillaries from normal

The angiograms with maximum hybrid contrast ratio greater than the empirically derived threshold of 0.3 are identified as those with dilated capillaries.

2.5 Contrast-and-saliency-based detection

The hybrid contrast ratio identifies potential regions with dilated capillaries but cannot identify their specific location. The saliency model (RARE2012), while it is unreliable in distinguishing eyes with dilated capillaries from normal eyes, can effectively localize the regions with dilated capillaries based on brightness, orientation, and position in eyes with known dilated capillaries [23]. We used the RARE2012 model to derive a saliency map, which we then multiplied to the hybrid contrast ratio. We extracted the potential regions of dilated capillaries by applying Otsu’s threshold.

2.6 Connectivity-based capillary grading

The enhanced DCP angiograms were divided into different vascular networks according to connectivity. If the vascular network has an intersection with the potential region, the vascular network was considered as dilated capillaries.

2.7 Evaluation metrics

An experienced masked grader qualitatively examined the en face DCP angiograms [7]. The dilated capillaries were localized using a modified Early Treatment of Diabetic Retinopathy Study (ETDRS) grid [24], dividing the 3 × 3 mm angiogram into a central 500 micron circle, and the remaining area into superior, inferior, nasal, and temporal quadrants. The grid was centered at the center of the image. The grader determined whether dilated capillaries were present or absent in each quadrant in a binary fashion.

The agreement between our automated algorithm and manual grading was evaluated, in term of accuracy ($\text{Acc}$), sensitivity ($\text{Se}$), specificity ($\text{Sp}$) [25]. These metrics are defined as follows:

$$\text{Acc} = \frac{tp + tn}{tp + fp + tn + fn}$$  \hspace{1cm} (6)

$$\text{Se} = \frac{tp}{tp + fn}$$  \hspace{1cm} (7)

$$\text{Sp} = \frac{tn}{tn + fp}$$  \hspace{1cm} (8)

where $tp$, $tn$, $fp$ and $fn$ indicate the true positive (the number of correctly identified dilated capillaries sector), true negative (the number of correctly identified normal vessel sector), false positive (the number of incorrectly identified dilated capillaries sector), and false negative (the number of incorrectly identified normal vessel sector).
3. Results

Eyes from 10 healthy volunteers and 10 participants with DR with dilated capillaries were imaged. One eye from each participant was randomly selected for the study.

Figure 2 illustrates how each step of the algorithm processes the original DCP angiogram (Fig. 2(A)). The vesselness filter first suppressed (Fig. 2(B)) the background noise. By thresholding the hybrid contrast ratio map (Fig. 2(C)), we determined that the angiogram is from a DR eye with dilated capillaries. Next, the denoised DCP angiogram is further enhanced by applying a Gabor filter (Fig. 2(D)). Then, a saliency model detected the dilated capillaries regions. The saliency calculation was performed at multiple scales [26] and combined into a single saliency map (Fig. 2(E)). The potential regions (Fig. 2(F)) were more distinctly identified by multiplying the hybrid contrast ratio map by the saliency map. Finally, the dilated vascular network (Fig. 2(G)) was extracted from the enhanced DCP image based on connectivity.

Fig. 2. Illustration of detecting dilated capillaries by the proposed techniques. (A) Original en face deep capillary plexus (DCP) angiogram. (B) Denoised DCP angiogram. (C) Hybrid contrast ratio map combining global contrast and local contrast. (D) Enhanced DCP angiogram by Gabor filtering. (E) Saliency map. (F) The potential regions of dilated capillaries detected by incorporating the saliency and contrast ratio maps. (G) Extracted dilated capillaries.
Fig. 3. The comparison of DCP angiograms and their hybrid contrast ratio maps between healthy participants and DR participants with dilated capillaries. The first column shows examples of DCP angiograms with normal vessels. The second column shows the corresponding hybrid contrast ratio maps. The third column shows DCP angiograms with dilated capillaries. The fourth column shows the corresponding hybrid contrast ratio maps.

Figure 3 shows 4 DCP angiograms each of normal and severe DR eyes with dilated capillaries paired with their hybrid contrast maps. Figure 4 shows that the maximum hybrid contrast ratios can distinguish normal DCP angiograms from those with dilated capillaries, using a fixed threshold of 0.3.

Fig. 4. The plot of maximum hybrid contrast ratio in one scan of each participant. The cutoff of 0.3 can completely separate the healthy eyes (blue bars) and the DR eyes with dilated capillaries (red bars).

The results from the automated algorithm were compared with the results from expert grading. Five representative cases are shown in Fig. 5. All DR cases in this study showed dilated capillaries in the DCP angiogram. The proposed algorithm agreed well with human grading in sector by sector comparison. The detection accuracy was 0.92 (95% CI: 0.80-0.97); the detection sensitivity was 0.95 (95% CI: 0.80-0.99); and the detection specificity was 0.85 (95% CI: 0.54-0.97).
Fig. 5. Comparison of detection of dilated capillaries between the automated algorithm and human grading. The top row shows the DCP angiograms. The second row shows the dilated capillaries that were extracted by the proposed algorithm. The bottom row shows the results of human grading.

4. Discussion and conclusion

We have proposed an algorithm for detection of dilated capillaries in DCP, which is correlated with severe DR. The vesselness filter removed the background noise. A hybrid contrast ratio map was used to distinguish eyes with dilated capillaries from normal eyes. The contrast ratio map and saliency map were used to locate the regions with dilated capillaries. Morphologic operations were applied to identify the continuous extent of dilated capillaries. This algorithm identified dilated capillaries with 0.92 diagnostic accuracy compared to human grading.

Projection artifacts from large superficial vessels project high flow signals onto the DPC angiograms and can result in false positive detection of dilated capillaries. The removal of these artifacts using PR-OCTA was a critical step for accurate detection of dilated capillaries in the DCP.

Due to the variation of image quality in real world data, significant variation in the absolute signal strength of DCP angiogram was present. For example, the angiogram of the third DR case in Fig. 5 is significantly dimmer than others. The proposed algorithm in this study performed well regardless of low image quality. Further studies are needed to evaluate the performance of the algorithm on scans with signal strength variation due to vignetting and defocusing, etc.

Although this algorithm identified dilated capillaries effectively in the study eyes, it may not perform as well in angiograms that consist mostly of dilated vessels due to the lack of saliency against normal vessels. For example, the algorithm failed to detect some regions in the second DR case in Fig. 5. To avoid the false negative error caused by this issue, one potential solution is to acquire scans with a larger field of view and allow more normal vessels to be included in the processing region.

This study is limited by the lack of the eyes of DR without dilated capillaries. In a future study, it would be useful to investigate the correlation between automatically detected dilated capillary area and the severity of DR. Furthermore, longitudinal studies to monitor the dilated capillaries or track its change in response to treatment would also be valuable. In summary,
this novel algorithm can automatically detect dilated capillaries in retinal deep plexus in DR with good accuracy compared to human grading.

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