A new diagnostic model of primary open angle glaucoma based on FD-OCT parameters

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Abstract

AIM: To build a clinical diagnostic model of primary open angle glaucoma (POAG) using the normal probability chart of frequency-domain optical coherence tomography (FD-OCT).

METHODS: This is a cross-sectional study. Total 133 eyes from 133 healthy subjects and 99 eyes from 99 early POAG patients were included in the study. The retinal nerve fibre layer (RNFL) thickness parameters of optic nerve head (ONH) and RNFL3.45 scan were measured in one randomly selected eye of each subject using RTVue-100 FD-OCT. Then, we used these parameters to establish the diagnostic models. Four different diagnostic models based on two different area partition strategies on ONH and RNFL3.45 parameters, including ONH traditional area partition model (ONH-T), ONH new area partition model (ONH-N), RNFL3.45 traditional area partition model (RNFL3.45-T) and RNFL3.45 new area partition model (RNFL3.45-N), were built and tested by cross-validation.

RESULTS: The new area partition models had higher area under the receiver operating characteristic (AROC; ONH-N: 0.990; RNFL3.45-N: 0.939) than corresponding traditional area partition models (ONH-T: 0.979; RNFL3.45-T: 0.881). There was no statistical difference among AROC of ONH-T, ONH-N, and RNFL3.45-N. Nevertheless, ONH-N was the simplest model.

CONCLUSION: The new area partition models had higher diagnostic accuracy than corresponding traditional area partition models, which can improve the diagnostic ability of early POAG. In particular, the simplest ONH-N diagnostic model may be convenient for clinical application.

KEYWORDS: primary open angle glaucoma; optical coherence tomography; ethnic-specific database; diagnostic model

INTRODUCTION

Glaucoma is the second largest cause of bilateral blindness in the world, affecting over 67 million people worldwide[1]. Among glaucoma, the most common form is primary open angle glaucoma (POAG) which is characterized by the loss of retinal ganglion cells and their axons, and affects 33 million individuals worldwide[2-4]. Study has implied that loss of retinal nerve fibre layer (RNFL) and lesion in optic disc may exist even prior to the occurrence of visual defect in glaucoma patients[5]. Therefore, early diagnosis and active treatment are very important in prevention and control of POAG.

Optical coherence tomography (OCT) is a high resolution glaucoma imaging device capable of RNFL and visual function measurements, which has been widely applied in recent years in the diagnosis of many ocular disorders due to its advantages of non-contact and non-invasive[6-8]. With the improvements of the fourth generation of OCT, the resolution and reproducibility of OCT increase gradually, comparing with previous generations[9-13]. Specially, frequency-domain (FD)-OCT has even built an ethnic-specific database of the healthy, based on which a normal probability chart is provided, in favor of the clinical diagnosis of POAG[14]. However, the present normal probability chart can only reflect the normal probability of each selected area instead of the probability of having glaucoma for single individual. Additionally, it is well known that retina nerve fiber in different area has different susceptibility to glaucomatous damage. The change of structural parameters in different locations has quite different weights in glaucoma diagnosis, especially in the early stage.

Our previous study has collected data from 99 early POAG and 133 healthy volunteers to build a diagnostic model of primary POAG using the FD-OCT ethnic-specific database of normal humans[15]. In that study, the diagnostic model was established (ONH=OSTITI:2+CA) based on multi-parameters [RNFL, optic nerve head (ONH), and ganglion cell complex (GCC)], which was relatively complicated.
FD-OCT in POAG

In the present study, we aimed to use two glaucoma scanning programs, ONH and RNFL3.45, in FD-OCT (RTVue-100, software version 6.1, Optovue, Fremont, CA, USA) to detect the RNFL thickness of the same patients and healthy volunteers. We proposed two area partition strategies and grading strategy for the parameters based on the normal probability chart provided by the ethnic-specific normative database to build a more effective and simple diagnostic model for early POAG.

SUBJECTS AND METHODS

Subjects and Samples This is a cross-sectional study. From July 2013 to March 2014, the patients diagnosed as early POAG in the follow-up examination at the specialty outpatient department in Peking University First Hospital (PUFH) were included in this study, including POAG patients with high pressure glaucoma (HPG) and normal tension glaucoma (NTG). The normal control group consisted of volunteers recruited from July 2013 to March 2014 in Beijing. This study followed the principles of the Declaration of Helsinki and was approved by Ethics Committee of PUFH.

The inclusion criteria for healthy volunteers were: 1) the best corrected visual acuity ≥0.8; 2) diopter: sphere -6.00S to +4.00S and cylinder -3.00C to +3.00C, and an isometropia ≤2D; 3) with normal chamber depth, transparent dioptric media, and normal fundus; 4) with healthy optic disc appearance, cup-to-disc ratio (C/D) <0.6, no evidence of diffuse or focal rim thinning, cupping, optic disk hemorrhage, or RNFL defects, interocular asymmetry of C/D<0.2; 5) intraocular pressure tested by applanation tonometer ≤21 mm Hg (1 mm Hg = 0.133 kPa), and central corneal thickness (CCT) between 520 μm and 580 μm; 6) with normal visual field, and determined as “within normal limits” by glaucoma hemifield test (GHT); 7) without family history of glaucoma; 8) without history of diabetes, optic neuropathy, uveitis, eye trauma and intraocular surgery; 9) without diabetes, high blood pressure or any other systematic diseases which might influence the outcome of measurement; 9) age >18y.

A total of 133 eyes from 133 healthy subjects and 99 eyes from 99 early POAG patients were included in the study. We randomly selected 90 normal eyes and 60 early POAG eyes to construct modeling samples. The remaining samples, including 43 normal eyes and 39 early POAG eyes, were used as test samples to verify the diagnostic model.

Disease History Inquiry and Routine Ophthalmological Examination Routine ophthalmological examinations included eyesight and optometry examination (mydriatic optometry was performed for subjects <40 years old), slit-lamp examination, fundus examination with direct ophthalmoscope, Goldmann applanation tonometry, and gonioscope examination.

Visual Field Examination Visual field examination was performed using a Humphrey perimeter (Humphrey Field Analyzer model 750i; Carl Zeiss Meditec, Dublin, CA, USA) with the SITA rapid 24-2 program. The results were considered reliable if they were consistent with the following criteria: 1) fixation losses <20%; 2) both false positive rate and false negative rate <30%.

Optical Coherence Tomography Examination Optovue RTVue100 (software version 6.1; Optovue, Fremont, CA, USA) FD-OCT was used for the OCT examination. The scanning wavelength of RTVue-100 was 840±10 nm, with the depth of the scanning area of 2-2.3 mm, length of 2-12 mm, longitudinal resolution of 5 μm, transverse resolution of 15 μm and scanning speed of 26 000 A-scans/s.

The scanning was performed under non-mydriatic condition in internal fixation, and “China” was selected for the race. Three scanning models, including MM6/Radial Slicer, ONH, and RNFL3.45, were used for each subject. Images scanned were stored if they were complete, with signal strength indicator (SSI) ≥35 and clear fundus image, and without strong reflection.

Fundus Color Photography A Topcon TRC-ss fundus color stereo camera was used for vectograph of fundus in patients with early POAG. Mydriasis was induced by the instillation
of 0.5% compound tropicamide by an experienced technician. The photos were read by a glaucoma specialist.

**Determination of Colors for the Scanning Parameter**

When the RNFL thickness of patients determined by OCT were beyond the normal range compared with the database, the result would be marked with a specific color by the software in the report. Those with normal distribution probability less than 5% were marked as yellow (critical status), and those with normal distribution probability less than 1% were marked as red (abnormality). A database of Chinese race was used in this research, and a correction had been made on the age and size of optic disc in this database.

**Area Partition Strategies**

Area partition was performed for RNFL thickness parameters of RNFL3.45 and ONH scan. They were divided into 16 sectors and combined into 6 regions (with 4 sectors for each): the superior temporal and the inferior temporal (STIT) included IT1, IT2, ST1 and ST2; the superior nasal and inferior nasal (SNIN) included SN1, SN2, IN1 and IN2; the superior temporal and superior nasal (STSN) included ST1, ST2, SN1 and SN2; the inferior temporal and the inferior nasal (ITIN) included IT1, IT2, TL1 and TL2; the nasal upper and the nasal lower (NUNL) included NU1, NU2, NL1 and NL2. The traditional partition included STSN, ITIN, TUTL and NUNL. The new partition included: STIT, SNIN, TUTL and NUNL (Figure 1).

**Statistical Analysis**

In the control group, one eye of each volunteer was randomly selected for statistics. In POAG patients, the eye with less severe disease was selected for statistics. SPSS 16.0 was used for statistical analysis, and \( P<0.05 \) was considered significant. Number of cases and percentages were used for enumeration data. Rank sum test was used for the comparison of difference in measurement data, and Chi-square test was used for the comparison of difference in enumeration data. Receiver operating characteristic (ROC) curves and area under the receiver operating characteristic (AROC) curve were used to evaluate the capability of each parameter and diagnostic model of FD-OCT in distinguishing early-stage POAG eyes from normal eyes. The method for comparing confidential interval was used to compare AROC. Total 90 healthy people and 60 early-POAG patients were randomly selected to form a modeling sample, then a combination of parameters was made with logistic regression to establish diagnostic models for early POAG. The established diagnostic models were validated by test sample that formed with the remaining healthy people and early-POAG patients.

**RESULTS**

**Demographics of Participants**

The demographics of participants were shown in Table 1. For diopter, there was no significant difference between glaucoma patients and normal subjects in modeling sample (\( P=0.795 \)) and test sample (\( P=0.640 \)). For visual field MD and pattern standard deviation, significant differences were found between glaucoma patients and normal subjects with \( P=0.000 \) for both modeling group and test group.

**Analysis of Partitioned Parameters in the Nerve Fibrous Layer**

Quantification of scanning parameters was necessary for the establishment of the diagnostic models. We used a scoring method for quantification of each parameter, with higher probability of abnormality in higher scores. It was scored 0 for green parameter (representing normality), and 1 for yellow parameter (representing critical status). The red parameter represented abnormality and its score should be higher than yellow, but the weight ratio between red and yellow should be calculated. Because of the difference in diagnostic values among the 6 regions, the weight ratio of the scores might differ between the yellow squares and the red ones in each region. Given 4 sectors (i.e. four small squares) in one region, there were five possibilities of the weight ratio between the red squares and the yellow ones, i.e. the weight ratio of one red square was equivalent to 1, 2, 3, 4 or more than 4 yellow squares. Therefore, when partitioned scoring was performed, it was scored 1 for the yellow squares, and 1, 2, 3,
4 or 5 for the red ones respectively, and the score of 5 in the red square meant the weight ratio of the red square was larger than 4 yellow squares.

Each of the two types of scans including ONH and RNF3.45 comprised of 6 regions, and each region might have 5 different weight ratios when it was scored for the red squares and the yellow ones, i.e. the five different scoring methods. In order to define the most appropriate scoring method for each region, AROC was made for five scoring methods in each region. The results revealed that the scoring methods with the highest AROC in the 6 regions scanned with RNFL3.45 were RSTIT12, RSNIN12, RSTIT12, RTUTL12, RSTSN12, and RITIN12 respectively, and that in the 6 regions scanned with ONH were ONUNL12, OSNIN13, OSTIT12, OTUTL15, OSTSN13 and OITIN13. Table 2 showed the AROC of the best scoring method in each region, and Table 3 presented the nomenclature.

The scoring method with the highest AROC in each region was used. The 4 regions based on the traditional partition (including STSN, ITIN, NUNL and TUTL) and the new partition (including STIT, SNIN, TUTL and NUNL) were combined respectively using logistic regression model to establish two area partition diagnostic models for early-stage POAG. The models of RNFL3.45 and ONH for the traditional partition were RNFL3.45-T=RTUTL12×0.37+ RSTSN12×0.65+RITIN12×0.95 and ONH-T=ONUNL12× -0.66+OSTSN13×1.51+OITIN13×1.3, and those for the new partition were RNFL3.45-N=RSTIT12 and ONH-N=OSTIT12. Table 4 presented AROC and its sensitivity, specificity, positive predictive value, negative predictive value, positive likelihood ratio and negative likelihood ratio for the 4 area partition diagnostic models.

Model Validation The 4 established models were respectively validated using the test sample. The general information of the test sample was given in Table 1. Table 5 showed the results of AROC and its sensitivity, specificity, positive predictive value, negative predictive value, positive likelihood ratio and negative likelihood ratio for the 4 models.

DISCUSSION Glaucoma is characterized by a combination of structural and functional damage\(^{[13]}\). The structural changes in glaucoma patients usually precede functional changes\(^{[16-17]}\). Therefore, the most important challenge in glaucoma research is to find...
Table 4 The AROC, sensitivity, specificity, PV+, PV-, LR+ and LR- for the 4 area partition diagnostic models established using the modeling samples

<table>
<thead>
<tr>
<th>Models</th>
<th>AROC</th>
<th>P</th>
<th>95% confidential interval</th>
<th>Critical value</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>LR+</th>
<th>LR-</th>
<th>PV+ (%)</th>
<th>PV- (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RNFL3.45-T</td>
<td>0.956±0.018</td>
<td>0.000</td>
<td>0.921-0.990</td>
<td>2.33</td>
<td>90.0</td>
<td>90.0</td>
<td>9</td>
<td>0.11</td>
<td>95.7</td>
<td>93.1</td>
</tr>
<tr>
<td>ONH-T</td>
<td>0.990±0.006</td>
<td>0.000</td>
<td>0.978-1.000</td>
<td>3.14</td>
<td>98.3</td>
<td>94.4</td>
<td>17.55</td>
<td>0.02</td>
<td>92.2</td>
<td>98.8</td>
</tr>
<tr>
<td>RNFL3.45-N</td>
<td>0.966±0.017</td>
<td>0.000</td>
<td>0.933-0.998</td>
<td>2</td>
<td>94.9</td>
<td>93.3</td>
<td>14.16</td>
<td>0.05</td>
<td>90.5</td>
<td>96.6</td>
</tr>
<tr>
<td>ONH-N</td>
<td>0.990±0.007</td>
<td>0.000</td>
<td>0.977-1.003</td>
<td>2</td>
<td>98.3</td>
<td>96.7</td>
<td>29.79</td>
<td>0.02</td>
<td>95.2</td>
<td>98.9</td>
</tr>
</tbody>
</table>

PV+: Positive predictive value; PV-: Negative predictive value; LR+: Positive likelihood ratio; LR-: Negative likelihood ratio.

Table 5 The AROC and its sensitivity, specificity, PV+, PV-, LR+ and LR- of the 4 models established using the test samples

<table>
<thead>
<tr>
<th>Models</th>
<th>AROC</th>
<th>P</th>
<th>95% confidential interval</th>
<th>Critical value</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>LR+</th>
<th>LR-</th>
<th>PV+ (%)</th>
<th>PV- (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RNFL3.45-T</td>
<td>0.881±0.039</td>
<td>0.000</td>
<td>0.805-0.958</td>
<td>2.33</td>
<td>84.6</td>
<td>79.1</td>
<td>4.048</td>
<td>0.195</td>
<td>78.6</td>
<td>85.0</td>
</tr>
<tr>
<td>ONH-T</td>
<td>0.979±0.013</td>
<td>0.000</td>
<td>0.954-1.000</td>
<td>3.14</td>
<td>94.9</td>
<td>86.0</td>
<td>6.77</td>
<td>0.059</td>
<td>86.0</td>
<td>94.9</td>
</tr>
<tr>
<td>RNFL3.45-N</td>
<td>0.939±0.028</td>
<td>0.000</td>
<td>0.884-0.993</td>
<td>2</td>
<td>89.7</td>
<td>90.7</td>
<td>9.65</td>
<td>0.114</td>
<td>89.7</td>
<td>90.7</td>
</tr>
<tr>
<td>ONH-N</td>
<td>0.990±0.007</td>
<td>0.000</td>
<td>0.977-1.000</td>
<td>2</td>
<td>94.9</td>
<td>95.3</td>
<td>20.19</td>
<td>0.054</td>
<td>94.9</td>
<td>95.3</td>
</tr>
</tbody>
</table>

PV+: Positive predictive value; PV-: Negative predictive value; LR+: Positive likelihood ratio; LR-: Negative likelihood ratio.

the early glaucomatous structural damage. Recent developed FD-OCT technology not only has high resolution and good repeatability, but also provides both an ethnic-specific normal database with parameters (age and optic disc size) corrected and a corresponding diagnostic probability graph[18-19]. At present, we preliminarily discussed the application value of database and probability graph of Optovue RTVue100 FD-OCT in early diagnosis of POAG. Compared with our previous diagnostic model that was established based on multi-parameters, the present study focused on the different AROC between two diagnostic models based on two different area partition strategies on ONH and RNFL3.45 parameters, obtaining a more simple and practical model.

**Comparison with Single Parameter** The AROC had been used to evaluate the diagnostic accuracy and compare the diagnostic performance of single and combined anatomic variables in many studies. Some studies suggested that combined parameters could optimize the glaucoma diagnosis[20-21]. Lu et al[20] performed a study to identify the best combination of Stratus OCT RNFL thickness parameters for the detection of glaucoma and found that overall average RNFL thickness had the highest AROC value of all single parameters evaluated, while the 3-parameter combination was significantly better than the overall average alone. Recent research of Wang et al[21] combined time-domain OCT measurements of the optic disc, circumpapillary RNFL, and macular retinal thickness to improve the diagnosis of glaucoma. They found that all combination diagnostic variables had significantly larger AROCs than any single diagnostic variable. Furthermore, some recent studies with FD-OCT also reported similar results. For instance, Fang et al[22] evaluated the diagnostic capability of parameters of the optic disc, RNFL thickness, and GCC using FD-OCT for early POAG patients and suggested that the combined parameters had the highest AROC. In accordance with previous studies, our result showed that the AROC of combined diagnostic model ONH-N was 0.990, which was significantly higher than the RNFL average (AROC=0.912), the best single parameter of GCC, optic disk and RNFL scanning parameters (shown in our previous study)[19]. There simplified that diagnostic capability of our area partition model may be better than single parameter. Application of this model in clinical practice may be able to improve the sensitivity and specificity of FD-OCT in early diagnosis of POAG.

**Comparison Between Diagnostic Models** It has been reported that RNFL defect mostly occurs in inferior temporal or superior temporal in early POAG[20,25-26]. Based on the traditional partition, we introduced the novel partition, i.e. the combined region of STIT and SNIN. Thus points along the circumference scanned by ONH and RNFL3.45 were divided into six regions (with 4 points for each) including STIT. Among the six regions, STIT had the highest AROC. Although there was no statistical difference between AROC among the four diagnostic models, the new partition diagnostic models (i.e. RNFL3.45-N, ONH-N) seemed to be simpler and more convenient in clinic. STIT12 was the only left over models (RNFL3.45-N, ONH-N) seemed to be simpler and more convenient in clinic. STIT12 was the only left over model in the model after regression analysis. STIT12 is a parameter obtained by assigning red squares score 2 and yellow squares score 1 and summing up the scores of four squares in STIT. The sensitivity and specificity of the STIT12 score in diagnosis of early POAG was 98.3% and 96.7% respectively with the cut-off level of 2.
In our previous work, an aggregative model was established combining the parameters of RNFL, optic disc and GCC\cite{24}. The aggregative model was more complex than the new area partition models. Overlapping of the confidential interval between them had no statistical difference. So we hold that the new area partition models are more suitable for clinical application. Being different from previous studies\cite{20,22-23,25,27-29}, which used parameters directly coming from FD-OCT, we divided the 16 FD-OCT parameters into 6 sections and built the scoring system for each section, and then built the logistical diagnostic model using these section parameters and calculated its AROC. Thus, we could get much simpler model which was easy to use in clinic.

**Verification with the Test Sample** When we applied the four models in the test sample, the result showed that except for RNFL3.45-T, the AROCs of the three diagnostic models (ONH-T, RNFL3.45-N, ONH-N) were more than 0.90, suggesting a high diagnostic ability of the scoring system and diagnostic models. This result implied that the diagnostic efficiency of the diagnostic models that was recommended in the modeling sample was validated in the test sample. Overlapping of the confidential interval among the three models had no significant difference. Among them, ONH-N=OSTIT12 seemed to have the highest AROC and might be the most simplified and convenient model. Based on the result, we expected that in clinic, when a case showed 1 red square or 2 yellow squares among the four squares in STIT in the normal probability graph of RNFL parameter under the ONH scanning, doctors could make a primary diagnosis for glaucoma patient with a relative high sensitivity and specificity. It might facilitate the reading of FD-OCT scanning results to some extent and improve the application value of OCT scanning parameters in the early diagnosis of POAG, especially for non-glaucoma specialist.

In spite of no significant difference, our result showed that the AROCs of the new regional diagnostic models (ONH-N and RNFL3.45-N) were higher than that of the traditional regional diagnostic models (ONH-T and RNFL3.45-T) respectively. Besides, the AROCs of some parameters in ONH scanning (i.e. ONH-N) were significantly higher than that in RNFL scanning (i.e. RNFL3.45-T). The results may suggest not only the superiority of ONH scanning to RNFL3.45 scanning, but also the superiority of new partition models to traditional partition models in diagnostic efficiency. More researches are needed to verify the results.

In this study, we attempted to establish a scoring system and diagnostic models, using FD-OCT ethnic-specific database of normal humans, and tried to simplify and facilitate their clinical application. There are still some limitations for our research. The integral multiple was used for indexes when establishing the models, which might reduce the diagnostic efficiency of the models. In addition, the sample size in each group was a little small, especially in preperimetric glaucoma group. An expanded sample will be expected in our future study, especially for patients with preperimetric glaucoma in glaucoma group and physiologic large cup in normal group, to optimize our model for the early diagnosis of early POAG.

**ACKNOWLEDGEMENTS**

**Authors’ contributions:** Conception and design of the research: Zheng YJ; Acquisition of data: Fang Y and Qiao RH; Analysis and interpretation of data: Li M and Cai Y; Statistical analysis: Li XY; Drafting the manuscript: Zheng YJ; Revision of manuscript for important intellectual content: Pan YZ.

**Conflicts of Interest:** Zheng YJ, None; Pan YZ, None; Li XY, None; Fang Y, None; Li M, None; Qiao RH, None; Cai Y, None.

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