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# **Automated Beta Zone Parapapillary Area Measurement to Differentiate Between Healthy and Glaucoma Eyes**

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# **Abstract**

**Purpose:** To evaluate whether automated assessment of beta-zone parapapillary atrophy (**ß**PPA) area can differentiate between glaucomatous and healthy eyes of varying axial lengths (AL).

**Design:** Cross-sectional study.

**Methods: ß**PPA was automatically identified in glaucoma and healthy eyes with enhanced depth imaging optical coherence tomography (OCT) optic nerve head (ONH) radial B-scans. Associations with AL and the presence of glaucoma were assessed. Manually-delineated **ß**PPA on individual OCT ONH B-scans of 35 eyes from the Diagnostic Innovations in Glaucoma Study served to validate the automated method.

**Results:** One hundred fifty-three glaucoma eyes (mean ± SD) (visual field mean deviation: −5.0  $\pm$  6.4 dB and mean axial length, 25.1  $\pm$  1.1 mm) and 73 healthy eyes (visual field mean deviation,  $0.1 \pm 1.4$  dB; and mean axial length  $24.1 \pm 1.1$  mm) were included. In multivariable analysis, larger **ß**PPA area was significantly associated with a diagnosis of glaucoma after controlling for age, central corneal thickness, and axial length. Moreover, in multivariable analysis, the odds of having glaucoma were doubled for each 0.2 mm<sup>2</sup> larger **ß**PPA area. The age- and AL-adjusted area under the receiver operating characteristic curve (95% CI) of **ß**PPA area for differentiating

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between glaucoma and healthy eyes was 0.75 (0.68–0.81). Agreement for the location of Bruch's membrane opening and the location of retinal pigment epithelium tips was stronger between the automated technique and each individual observer than it was between the two observers.

**Conclusions:** Larger **ß**PPA area, as determined by automated OCT assessment, is significantly associated with a diagnosis of glaucoma, even after adjusting for age and axial length, and can aid in differentiating healthy from glaucomatous eyes.

# **INTRODUCTION**

Parapapillary atrophy (PPA) occurs adjacent to the optic disc and represents a complete loss of retinal pigment epithelium (RPE), an almost complete loss of retinal photoreceptors and a partial closure of the choriocapillaris.<sup>1,2</sup> It was formerly called "halo glaucomatosus"<sup>3</sup> due to its association with severe glaucomatous damage.<sup>3–8</sup> PPA occurs more often in glaucomatous than non-glaucomatous eyes and is associated with an increase in the risk of glaucoma progression.<sup>9–14</sup> PPA is also associated with axial myopia,<sup>7,11,15,16</sup> which limits its use for the diagnosis of glaucoma.<sup>17</sup>

In the clinical setting, PPA is typically divided into two subgroups. The alpha PPA zone (αPPA) consists of regions of hypopigmentation and hyperpigmentation of the retinal pigment epithelium and mostly intact photoreceptors, and is usually located peripherally to beta PPA zone **ß**PPA), when the latter is also present. Histologically, αPPA is equivalent to irregularities of the RPE overlying an intact Bruch's membrane (BM).1,3,11,16 **ß**PPA is characterized clinically by visible sclera and large choroidal vessels.<sup>1,3,11</sup>,<sup>18</sup>

Recent advances in spectral-domain OCT imaging have facilitated the detection of Bruch's membrane opening (BMO) and has led to the identification of different subtypes of beta PPA: Beta-zone **ß**PPA) and gamma-zone PPA.11,12 **ß**PPA is the PPA area with intact BM but without RPE, while gamma-zone PPA is the PPA area without BM and is located between the optic disc clinical border and the edge of BM  $^{1,3,11,16}$  (Figure 1). Based on manual delineation of the PPA subtypes of OCT images, bot **ß**PPA and gamma-zone PPA area have been shown to be associated with increased axial length, 19,20 while **ß**PPA is consistently larger in glaucoma eyes than healthy eyes.19–21 It has been suggested that **ß**PPA may be an age-related atrophic change and more strongly associated with glaucoma, while gamma PPA may be due to scleral stretching associated with elongation of the globe in myopia. 14,22,23

For these reasons, it has been suggested that distinguishing between these 2 PPA subtypes may help to differentiate between myopic eyes with and without glaucoma and/or identify eyes at risk of glaucomatous progression,  $11,12,14$  There are a limited number of studies evaluating this issue with mixed results. 11,13,19,24,25

The San Diego Automated Segmentation Algorithm (SALSA) was developed to segment retinal layers from OCT images,  $26,27$  and now includes the ability to measure the area of **ß**PPA with intact BM. The purpose of this report is to determine whether the automated measurement of **ß**PPA area can be used to differentiate between glaucoma and healthy eyes with a wide range of axial lengths (AL).

# **MATERIALS AND METHODS**

#### **Participants**

This cross-sectional study included subjects from 2 longitudinal observational cohort studies, the Diagnostic Innovations in Glaucoma Study (DIGS, [clinicaltrials.gov](http://clinicaltrials.gov) identifier NCT00221897); and the African Descent and Glaucoma Evaluation Study (ADAGES, [clinicaltrials.gov](http://clinicaltrials.gov) identifier NCT00221923).28–30 The Institutional Review Boards of the University of California San Diego, University of Alabama, Birmingham, and of the New York Eye and Ear Infirmary approved the protocol, and the methodology adheres to the tenets of the Declaration of Helsinki.

All subjects were at least 18 years old and required to have open anterior chamber angles upon gonioscopy. Healthy subjects were defined as individuals without clinical signs of retinal or glaucomatous pathologies based on a clinical examination including ophthalmoscopy with medically dilated pupils, and a reliable visual field test without evidence of repeatable visual field damage on Standard Automated Perimetry (SAP). Glaucoma patients were defined as individuals who had glaucomatous optic neuropathy on dilated clinical examination and at least two consecutive, reliable (<33% fixation losses and false negatives, and <15% false positives) and repeatable abnormal SAP tests with the Humphrey 24–2 Swedish Interactive Threshold Algorithm with Pattern Standard Deviation or a Glaucoma Hemifield Test results outside the normal limits. Healthy and glaucoma eyes were selected to be of similar ages and to reflect a wide range of axial lengths.

#### **OCT Image Acquisition**

Spectral domain optical coherence tomography (SD-OCT) (Spectralis HRA+OCT; Heidelberg Engineering Inc., Heidelberg, Germany) was used to acquire 48 high-resolution EDI-OCT radial scans centered on the ONH. The scan settings included an automatic real time (ART) mode of 9 frames, 73 sections, a B-scan distance of approximately 7 microns between each A-scan, and 1024 A-scans scans per B-scan. BMO area was measured from Spectralis OCT optic disc scans using the San Diego Automated Layer Segmentation Algorithm (SALSA). 26

#### **Parapapillary Atrophy Zone Identification**

Areas of the **ß**PPA were automatically identified by calculating the area between the BMO and RPE tips in SD-OCT radial B-scans using SALSA. Details of SALSA and its validation have been described previously.<sup>26,27,31,32,33</sup> Two ophthalmologists (PICM, MHS), masked to patient diagnosis, independently identified the borders of **ß**PPA by manually marking the 2 RPE tips and the 2 BMO locations on each EDI-OCT radial B-scans of 35 eyes (Fig. 1) using a custom software program in MATLAB.<sup>34</sup> The EDI radial scans consisted of 48 individual radial scans. Every other B-scan was selected for manual delineation resulting in delineation of 24 equidistant scans. **ß**PPA borders identified manually were then compared to SALSA identification of the location of the BMO and RPE tips.

#### **Statistical Analysis**

The associations between the automatically calculated **ß**PPA area with age, AL, intraocular pressure (IOP), BMO area and central corneal thickness (CCT) in healthy subjects and glaucoma patients was evaluated using STATA. 35 Ordinary least squares regression with the cluster option to adjust for the correlation between eyes within subjects was utilized. Univariate and multivariable linear and logistic regressions were completed to evaluate the association of **ß**PPA area and diagnosis, respectively after controlling for age and other ocular parameters. Semipartial correlation coefficients  $(R^2)$  were calculated to adjust for age in the correlations between **ß**PPA area and axial length in healthy and glaucoma eyes. The diagnostic accuracy of **ß**PPA area for differentiating between healthy and glaucoma eyes was assessed by calculating the age- and axial length-adjusted area under the Receiver Operating Characteristic Curve (AUROC), as previously described.36 Agreement between observers and the automated detection of the location of the BMO and RPE (in pixels) was assessed using Bland-Altman plots and summarized as the 95% confidence interval (CI) limits of agreement of the mean difference in pixels.

# **RESULTS**

Two hundred and twenty-six eyes from 88 glaucoma patients and 38 healthy subjects were included (Table 1). Glaucoma participants had a mean age of  $64.2 \pm 9.4$  years, (median = 66.9 years), a mean visual field (VF) mean deviation (MD) of  $-5.0 + 6.4$  dB (median =  $-2.6$ dB), and mean axial length (AL) of  $25.1 + 1.1$  mm (median =  $25.0$  mm). Healthy participants had a mean age of  $60.6 + 10.4$  years (median = 59.0 years), a mean VF MD of  $0.1 + 1.4$  dB (median = 0.5 dB), and mean AL of 24.1 + 1.1 mm (median = 24.0 mm). Glaucoma patients had significantly larger **ß**PPA area (p<0.001), longer axial lengths  $(p<0.001)$ , and worse VF MD  $(p<0.001)$  than healthy eyes.

In univariate analysis of healthy and glaucoma eyes, the automatically delineated **ß**PPA area increased with longer axial length ( $R^2 = 30.1\%$ ; p<0.001) (Table 2). The age-adjusted association between **ßPPA** area and axial length was stronger in glaucoma eyes ( $R^2 = 24.6\%$ )  $p<0.001$ ) than healthy eyes ( $R^2$  = 19.9%,  $p=0.009$ ) (Figure 2).

We also evaluated the univariate and multivariable association of age, AL, BMO area, IOP, CCT, and glaucoma diagnosis with **ß**PPA area as a dependent variable (Table 2). Older age, longer AL, larger BMO area, diagnosis of glaucoma, and higher intraocular pressure (IOP) were each significantly associated with **ß**PPA area in univariate analysis. In multivariable analysis, larger **ß**PPA area was significantly associated with a diagnosis of glaucoma and longer AL. This multivariable model explained 36.9% (R<sup>2</sup>) of the variance in **ßPPA** area.

In addition, we completed multivariable logistic regression analysis with glaucoma diagnosis as the dependent variable and controlling for age, axial length, BMO area, IOP, and CCT. We found that the presence of glaucoma was significantly associated with larger **ß**PPA area. Specifically, for each 0.2 mm<sup>2</sup> higher **ßPPA** area, the odds of having glaucoma were doubled (odds ratio (95% CI:)) 2.1 (1.2 – 3.6)  $p = 0.007$ , Table 3).

The diagnostic accuracy of **ß**PPA area for differentiating between healthy and glaucoma eyes was also evaluated (Figure 3). The age- and axial length-adjusted AUROC (95% CI) for **ß**PPA area was 0.75 (0.68–0.81).

Bland-Altman plots were constructed to assess the agreement between the 2 observers and automated assessment of the RPE tips and BMO (Figures 4 and 5). Better agreement was found between automated detection by SALSA and each of the individual graders for both the RPE tip and BMO identification (95% CI for grader 1: approximately 90 and 32 pixels, respectively; and grader 2: approximately 40 and 15 pixels, respectively), than the interobserver agreement between the 2 graders (95% CI approximately 120 pixels and 38 pixels, respectively).

### **DISCUSSION**

Our results suggest that automated calculation of **ß**PPA area has good agreement with manual expert assessment and has potential for improving our ability to differentiate between healthy and glaucoma eyes. Specifically, larger **ß**PPA area was significantly associated with a diagnosis of glaucoma, even after adjusting for age and axial length. Moreover, the odds of having glaucoma were doubled for each 0.2 mm<sup>2</sup> larger **ß**PPA area.

Dai et al.11 reported that the width and area of the **ß**PPA was significantly associated with longer axial length and the presence of glaucoma, while the width and area of the gamma zone was significantly associated with myopia, longer axial length, and the absence of glaucoma. Although not investigated in this study, larger **ß**PPA area has been shown to be associated with an increased risk of progression of VF damage in patients with glaucoma, while gamma zone is more strongly correlated with increased axial length. <sup>19,37</sup>

With an age and axial length adjusted AUROC of 0.75 (Figure 3), our results suggest that **ß**PPA area may improve our ability to detect glaucoma. These results are in contrast to a report by Vianna et al.24 that found no significant association between **ß**PPA area and glaucoma in myopic eyes ( $AUROC = 0.60$ ). There are several possible explanations for the differences between the current study and the investigation performed by Vianna and colleagues. First, over 90% of their study subjects were of European descent, whereas less than 50% of our subjects were of European descent (Table 1). Second, Vianna et al also limited their study population to myopic eyes with **ß**PPA, while our study included eyes with a range of axial lengths. As the authors suggest, including eyes that already had increased **ß**PPA would likely result in a weaker association of beta-PPA and glaucoma than including the general population in the analysis.

Our study demonstrated that **ß**PPA can be automatically detected from radial EDI ONH Bscans by calculating the area between the BMO and RPE tips. Automatic delineation can facilitate the routine assessment of **ß**PPA area and width in the nasal area, as well as the temporal area. Manual delineation of the RPE tip and BMO margin in the nasal area is challenging. For this reason, some studies focused on manual assessment of **ß**PPA width only in the temporal area.<sup>12,37</sup> The agreement between automated detection of both structures and each individual grader was similar or better than the inter-grader agreement of

manual BMO and RPE detection. BMO detection had better inter-grader agreement than RPE tip detection due to the easier identification of BMO compared to RPE in most eyes. <sup>26</sup> BMO identification with SALSA has been shown to have good reproducibility in a previous study (coefficient of variation  $<$  4%). <sup>26</sup> These results are also consistent with other studies suggesting that the detection of the BMO is easier than detection of the RPE tip, due to factors such as disc ovality, blood vessel shadowing and poor image quality that makes detection of RPE challenging.<sup>12,38</sup>

It has been suggested that **ß**PPA and gamma zone have different underlying pathophysiologies and that the probability of development and progression of glaucoma may differ in eyes with **ß**PPA and gamma PPA.12 Myopia-related PPA versus PPA related to aging or glaucoma may also have different underlying mechanisms.<sup>39</sup> While it is unclear which factors lead to an increased susceptibility of myopic eyes to glaucoma,  $40-46$ individuals with high myopia are 2.5 times more likely to have glaucoma, based on a metaanalysis of 7 studies (odds ratios (95% CI): 2.46 (1.93 to 3.15). <sup>44</sup> Other studies have been suggested that glaucoma may be over-diagnosed and overtreated in individuals with high myopia.<sup>47–52</sup> While recent advances in technology have contributed to faster and more efficient ways of diagnosing eye disease,  $53$  diagnosis of glaucoma in eyes with myopia has remained challenging. Automatic differentiation of the **ß**PPA zone in its new OCT-based definition may facilitate better detection of glaucoma in myopic eyes, as myopic optic nerve heads present significant challenges to diagnosing this disease correctly,<sup>49</sup> and facilitation of identifying eyes at risk of progression. <sup>25</sup>

There are several limitations to our study. First, the sample size, although similar to previously published studies on **ßPPA** area,<sup>11,12,24,54</sup> was relatively small. Second, as in other reports, our glaucoma group was older than the normal controls,  $^{11,24,37}$  though the difference did not reach statistical significance. For this reason, age was included in the multivariable analysis and the values of age-adjusted area under the receiver operating characteristic curve are reported. Future studies should include a larger sample of agematched patients with higher myopia or longer AL, in order to better evaluate **ß**PPA in patients with longer axial lengths in both healthy and glaucoma groups, as well as the impact of AL on the diagnostic accuracy of **ß**PPA. Third, we did not include automated analysis of gamma PPA area. While SALSA was able to detect **ß**PPA in all eyes that were manually segmented, automated detection of gamma PPA detection was found to be more challenging as the detection of the optic disc margin, defined as the border of the lamina cribrosa or as the intraocular projection of the optic nerve pia mater, is more difficult. We are currently working on methods to automate the detection of gamma PPA area.

In conclusion, larger **ß**PPA area was associated with an increased risk of having glaucoma in eyes with a range of axial lengths. Automated calculation of **ß**PPA area shows promise for aiding in the differentiation of eyes with glaucoma from those without glaucoma. Further studies are needed to determine the usefulness of automated calculation of **ß**PPA area for detecting glaucoma in the growing population of high myopes

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# **Figure 1:**

Example of manual delineation and automated identification of retinal pigment epithelial (RPE) tips and Bruch's Membrane (BM). Left: Manual delineation on an optic nerve head photograph and on an individual B-scan (middle). Alpha (α) parapapillary atrophy (PPA) is located between the white and blue arrows (RPE tip). Beta (β) PPA is located between blue (RPE tip) and yellow (BM) arrows. Gamma  $(\gamma)$  PPA is located between the yellow and red arrows. Right: Automated delineation of βPPA area is located between the BM (blue dots) and RPE tips (white dots). Clinical βPPA (manual dashed yellow line to white dots) cannot distinguish between βPPA & gamma PPA, as the BM is only visible with OCT.

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# **Figure 2:**

Linear regression for the relationship between beta zone area and axial length for healthy and glaucoma eyes, showing that a larger automatically calculated βPPA zone area was more strongly associated with axial length in glaucoma eyes ( $R^2 = 24.6\%$ ), than in healthy eyes  $(R<sup>2</sup> = 19.9%)$  after adjust for age.



# **Figure 3:**

The receiver operating characteristic curve of **ß**PPA area for differentiating between healthy and glaucoma eyes. The age- and axial length-adjusted area under the receiver operating characteristic curve (AUROC) (95% CI) is 0.75 (0.68–0.81).



## **Figure 4:**

Bland-Altman plots for retinal pigment epithelium (RPE) tip identification, with 95% confidence interval (CI) (top) inter-grader agreement of ~120 pixels, (lower left) automated RPE detection versus grader 1 agreement of ~90 pixels, and (lower right) automated RPE detection versus grader 2 agreement of ~40 pixels. Please note that every B-scan has 2 RPE tip locations.



#### **Figure 5:**

Bland-Altman plots for Bruch's membrane opening (BMO) identification, with 95% confidence interval (CI) (top) inter-grader agreement of  $\sim$ 38 pixels, (lower left) automated BMO detection versus grader 1 agreement of ~32 pixels, and (lower right) automated BMO detection versus grader 2 agreement of ~15 pixels. Please note that every B-scan has 2 BMO locations.

### **Table 1:**

# Patient Demographics<sup>a</sup>



<sup>a</sup>Values represent means and 95% Confidence intervals unless stated otherwise;

 $b$ CCT = Central corneal thickness;

 $c<sub>VF</sub>$  MD = Visual field mean deviation;

d **ß**PPA = beta parapapillary atrophy;

 $e^{i}$ IOP = Intraocular pressure.

\* Statistical analysis by subject

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# **Table 2:**

Association BPPA surface area with age, ocular parameters and glaucoma diagnosis: Univariate and Multivariable Analysis (n = 226) **ß**PPA surface area with age, ocular parameters and glaucoma diagnosis: Univariate and Multivariable Analysis (n = 226) Association



BPPA = beta parapapillary atrophy; AL = Axial length; CCT = Central comeal thickness; IOP = Intraocular pressure. **ß**PPA = beta parapapillary atrophy; AL = Axial length; CCT = Central corneal thickness; IOP = Intraocular pressure.

**Table 3:**

Multivariable Logistic Regression for Diagnosis of Glaucoma Multivariable Logistic Regression for Diagnosis of Glaucoma



AL = Axial length; CCT = Central corneal thickness; IOP = Intraocular pressure; ßPPA = beta parapapillary atrophy area. AL = Axial length; CCT = Central corneal thickness; IOP = Intraocular pressure; **ß**PPA = beta parapapillary atrophy area.