Pattern Electroretinogram Parameters and their Associations with Optical Coherence Tomography in Glaucoma Suspects

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Abstract

Aim: To investigate whether steady state pattern electroretinogram (ssPERG) could identify retinal ganglion cell (RGC) dysfunction, and to assess the relationship between ssPERG with optical coherence tomography (OCT) measurements in glaucoma suspects (GS).

Materials and methods: This was a prospective cohort study of GS, identified based on suspicious optic disk appearance and glaucoma risk factors. Complete eye exam, Standard automated perimetry, OCT, and ssPERG were performed. Magnitude (Mag), Magnitude D (MagD), and MagD/Mag ratio were subsequently used in the correlation and linear regression analyses between ssPERG parameters and the RNFL, GCL/IPL, and macular thicknesses measurements.

Results: Forty-nine eyes of 26 patients were included. Mag and MagD were significantly correlated with the superior, inferior, and average RNFL thicknesses (avRNFLT). All ssPERG parameters were significantly correlated with the average and minimum GCL/IPL thicknesses and the inner macular sector thicknesses. Mag and MagD significantly predicted the superior, inferior, and avRNFLT in the regression analysis. All ssPERG parameters were predictive of GCL/IPL thickness in all sectors as well as the average and minimum GCL/IPL thicknesses. All ssPERG parameters were predictive of all inner macular sector thicknesses and MagD was also predictive of some outer macular sector thicknesses as well.

Conclusion: ssPERG has significant correlations with and is predictive of RNFL, GCL/IPL, and macular thicknesses in glaucoma suspects.

Clinical significance: ssPERG may serve as a useful objective functional tool for identifying and following the progression of disease in glaucoma suspects.

Keywords: Ganglion cell layer, Glaucoma, Glaucoma suspect, Inner plexiform layer, Macula, Optic nerve, Optical coherence tomography, Retinal ganglion cell, Retinal nerve fiber layer, Steady state pattern electroretinogram.

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Introduction

Glaucoma is characterized by progressive visual loss secondary to irreversible retinal ganglion cell (RGC) dysfunction and death secondary to cellular and environmental stressors.1 The inner layers of the retina, the ganglion cell, and inner plexiform layers (GCL/IPL), have been proposed to be the preferred location for damage in glaucomatous eyes.2 It has also been demonstrated that there is a significant positive correlation between GCL/IPL thickness and the amplitude of pattern electroretinogram (PERG) of early glaucoma patients and this correlation was apparent prior to any detectable changes in the visual fields (VF) of patients.3,4 These findings suggest that PERG may serve as a more sensitive screening test for patients with early glaucoma compared to current methods, which have limited efficacy and variable accuracy in this population.5

Previous studies have also reported on the associations between PERG and the thicknesses of the retinal nerve fiber layer (RNFL) and macular thicknesses in glaucoma patients.3,5,6 PERG has also been found to predict rim area loss in preperimetric glaucoma patients after controlling for disk area.7 However, to our knowledge no studies have investigated the utility of PERG parameters for predicting structural changes on SD-OCT of all the macular, RNFL, and GCL/IPL sector thicknesses in glaucoma suspects. Whereas VF can be utilized as a functional test that can be correlated with structural damage for glaucoma patients, many glaucoma suspects patients have normal VFs,3,5 and at least 25% of RGC must be lost to detect a significant change on perimetry.10 Therefore, PERG may serve as a promising tool to bridge the gap and allow for the detection of visual dysfunction in glaucoma suspects. In recent years, steady state PERG (ssPERG) with the facilitated acquisition, interpretation, and improved subject comfort, was introduced.11 Additional filters and amplifiers were used to achieve adequate levels of amplitude and...
These studies suggested that the PERGLA protocol introduced by Porciatti et al. earlier could be used as a potential alternative to the classic PERG protocol in a clinical setting, and data interpretation using Fourier analysis in the PERGLA ssPERG is identical to the ISCEV recommendations. In recent years, the new office-based device, the Diopsys NOVA (Diopsys Inc., NJ) using the PERGLA protocol was introduced, with new validated proprietary parameters [Magnitude (Mag), MagnitudeD (MagD) and MagD/Mag ratio] to assess RGC function. They concluded that ssPERG parameters were repeatable, reproducible, and reliable to be used in clinical settings. In another study, Porciatti et al. demonstrated that ssPERG amplitude and phase were essentially uncoupled, implying that these measures reflected distinct different aspects of RGC functional activity.

The purpose of this present study was to investigate the relationship between ssPERG parameters and the SD-OCT thickness measurements of inner retinal layers in glaucoma suspects and to determine their individual contributions to the effects of RGC dysfunction on structural measures.

Materials and Methods
In this prospective cross-sectional study, a total of 26 eligible glaucoma suspects (49 eyes) were recruited from the Manhattan Eye, Ear & Throat Hospital ophthalmology practice and underwent a complete ophthalmologic examination, including slit-lamp biomicroscopy, Goldmann tonometry, standard automated perimetry (Humphrey Field Analyzer II, 24-2 and 10-2 SITA Standard strategy), OCT (Carl Zeiss Meditec Inc., Dublin, CA, USA) and steady state PERG (ssPERG) (Diopsys Inc., Pine Brook, NJ, USA). The study was approved by the Institutional Review Board of Northwell Health System. Written informed consent was obtained from all subjects and the study adhered to the tenets of the Declaration of Helsinki.

Study participants satisfied the following criteria: participants were 40–80 years old and had a best-corrected visual acuity better or equal to 20/40 as measured by Snellen visual acuity testing at the time of enrollment. All participants had a suspicious glaucomatous optic nerve head appearance (increased cup to disk ratio >0.4, or neuroretinal rim thinning, notching, or excavation), and a documented and repeatable normal Humphrey Field Analyzer (HFA) 24-2 at the baseline visit. All participants were not on intraocular pressure-lowering treatment at the time of enrollment. All individuals with prior intraocular or posterior segment intraocular surgery, ocular trauma, or ocular or systemic conditions that may affect optic nerve head structure and/or function, except for uncomplicated cataract extraction with posterior chamber intraocular lens implant and no escape of vitreous to the anterior chamber performed less than a year before enrollment, were excluded from this study.

Spectral-domain Optical Coherence Tomography
Average and in quadrant retinal nerve fiber layer thicknesses (avRNFLT), average, sectoral, and minimum GCCPLT were measured using the Optic Disk Cube protocol of a Cirrus spectral-domain optical coherence tomography (SD-OCT) version 6.0 as described elsewhere.

Visual Field Testing
All patients underwent standard automated perimetry (SAP) testing using the HFA 24-2 and 10–2 protocols. Visual fields with more than 20% fixation losses, false-negative errors, and false-positive errors were excluded. Only participants with visual fields corresponding to stage 0 (no visual field losses) following the Glaucoma Staging System 2 (GSS 2) were considered.

ssPERG Testing
The steady state PERG (ssPERG) was recorded using a commercially available system, Diopsys’ NOVA-PERG (Diopsys, Inc. Pine Brook, New Jersey, USA). ssPERG measurements from Diopsys’ are based on normative data from healthy subjects. Tests were performed in a dark room to standardized environment luminance, free of visual, and audible distractions. The patient’s seat height was adjusted so the tested eye stayed in a horizontal plane with the center of the monitor. The forehead skin was cleaned using NuPerp® Skin Prep Gel (Weaver and Company, CO, USA) and the lower eyelids using OCuSOFT® Lid Scrub Original (OCuSOFT Inc., Rosenberg, TX, USA) to ensure good and stable electrical activity. Disposable hypoallergenic skin sensors Silver/Silver Chloride ink (Diopsys’ proprietary Skin Sensor) were applied on the lower lid of both eyes, close to the lid margin and avoiding eyelashes. One ground sensor (Diopsys’ EEG electrode) was applied in the central forehead area with a small amount of conductive paste (Tens20®, Weaver, and Company) and cables from the Diopsys NOVA device were connected to the electrodes. A total of three electrodes were used per test per patient (two active/reference and one ground electrode). Subjects were fitted with the appropriate correction for a viewing distance of 24 inches and were instructed to fixate on a target at the center of the monitor in front of them.

The stimulus was presented on a gamma corrected Acer V176BM 17-inch monitor, having a refresh rate of 75 frames/second. Luminance output overtime was verified using a luminance meter MavoSpot 2 USB (Gossen, GmbH, Nuremberg; Germany). The pattern stimulus consisted of black/white alternating square bars, reversing at 15 reversals/second (rps) with a duration of 25 seconds for high contrast (HC 85%) and 25 seconds for low contrast (LC 75%) for a total of 50 seconds per eye. The stimulus field subtends a visual angle of 143×90 arc minutes. Each bar will subtend 22.49 arc minutes, for a total of 64 bars. A red target subtending 50.79 arc minutes was used as a fixation target and was centered on the stimulus field. The luminance of the white bars for 85% and 75% contrast was 204 cd/m² and the luminance for black was 20.5 cd/m² and 52.5 cd/m² yielding a mean luminance of 112.3 cd/m² and 128.2 cd/m², respectively. All recorded signals underwent band filtration (0.5–100 Hz), amplification (gain = 20,000), and averaging at least 150 frames. The signal was sampled at 1920 samples per second by an analog to digital (A/D) converter. The voltage range of the (A/D) converter was programmed between −5V and +5V. Sweeps contaminated by eye blinks or gross eye saccades were automatically rejected if they exceeded a threshold voltage of 50 μV, and these sections were identified as artifacts in the report. Synchronized single-channel electroretinography was recorded, generating a time series of 384 data points per analysis frame (200 ms). An automatic fast Fourier transformation was applied to the ssPERG waveforms to isolate the desired component at 15 rps. Other frequencies, such as those originating from eye muscles, were rejected. The ssPERG test results were saved in a Structured Query Language (SQL) database and presented in a report form to be used for analysis. For every subject, four pre-programmed full ‘contrast sensitivity protocols’ were performed sequentially, which consisted of two 25 seconds recordings for each eye: first with high contrast (85%) diffuse retinal
stimulation, then with low contrast (75%) pattern stimulation. The device collected five frames of data per second, totaling 125 frames of data, and the first 10 frames (2 seconds) of data were discarded. For each eye, three ssPERG measurements [Magnitude (Mag), MagnitudeD (MagD), and MagD/Mag ratio] were calculated. Mag (µV) represents the amplitude of the signal strength at the specific reversal rate of 15 Hz in the frequency domain, while MagD (µV) represents an adjusted amplitude of the ssPERG signal impacted by phase variability throughout the waveform recording. A recording where the phase of the response is consistent will produce a MagD value close to that of the Mag, whereas a recording where the phase of the response varies will produce a MagD value lower than that of Mag. This is because averaging responses that are out-of-phase with each other will cause some degree of cancellation. The MagD/Mag ratio is a ratio that is a within-subject representation of the phase consistency of ssPERG. The signal-to-noise ratio (SNR) represents the level of electrical noise compared with the level of the ssPERG signal at 15 Hz.

### Statistical Analysis

For all variables of interest, outliers with values ≥3 standard deviations from the mean were excluded from the analyses. Shapiro-Wilk test was used to determine the normality of the distribution for all important variables. ssPERG parameters were subsequently transformed to achieve normality of the distribution. ssPERG parameters achieved normal distribution through the following transformations: a log10 transformation of Mag (Shapiro-Wilk, p = 0.290), a log10 transformation for MagD (Shapiro-Wilk, p = 0.654), and a cubed transformation for MagD/Mag ratio (Shapiro-Wilk, p = 0.075). Descriptive statistics were used to evaluate continuous and demographic data. Mean and standard deviation values were determined for each ssPERG parameter (Mag, MagD, and MagD/Mag ratio), HFA Swedish Interactive Thresholding Algorithm (SITA) standard (24–2 and 10–2) tests, all RNFLT, GCL/IPL, and macular thickness variables. Associations among continuous variables were analyzed using partial correlations, adjusting for age, sex, intraocular pressure (IOP), central corneal thickness (CCT), and spherical equivalent (SE). Linear regression analyses were used to assess the relationships among ssPERG parameters and OCT derived structural RNFL, GCL/IPL, and macular thickness measures. Statistical Analyses were performed with commercially available software (IBM® SPSS® ver.25.0; SPSS Inc, Chicago, IL, USA).

### Results

#### Cohort Characteristics

This study included 49 eyes of 26 patients with a mean age of 59.86 years. Of these patients, 61.5% were female. Approximately 73.1% of patients identified as white and 15.4% identified as Hispanic. The mean IOP was 17.43 mm Hg and mean deviations (MD) 24–2 and 10–2 MD were 0.00 dB and 0.04 dB, respectively. The mean average retinal nerve fiber layer thickness (avRNFLT) was 90.23 µm. A complete breakdown of cohort characteristics is shown in Table 1.

#### Partial Correlation Analyses among ssPERG Parameters and RNFL Thickness Measurements

After controlling for known glaucoma risk factors such as age, sex, CCT, IOP, and SE, a partial correlation analysis between transformed ssPERG parameters and RNFL quadrants and average were performed (Table 2). Both Mag and MagD had positive correlations with the superior and inferior quadrants. The MagD/Mag ratio only had a positive correlation with the superior quadrant (p = 0.026) and had no significant correlation with the inferior quadrant (p = 0.158). Average RNFL thickness was significantly correlated with Mag (p = 0.005) and MagD (p = 0.007).

#### Partial Correlation analyses among ssPERG Parameters and GCL/IPL Thickness Measurements

After controlling for glaucoma risk factors (age, sex, CCT, IOP, and SE), a partial correlation between transformed ssPERG parameters and GCL/IPL thickness measurements were performed (Table 2). Mag had a positive correlation with superior (p = 0.011) and supertemporal (p = 0.019) and inferonasal (p = 0.043) GCL/IPL sectors. MagD and the MagD/Mag ratio had a positive correlation with all GCL/IPL sectors. All three PERG parameters had significant positive correlations with average GCL/IPL thickness and minimum GCL/IPL thickness.

### Table 1: Clinical features and characteristics of the study cohort (n = 49 eyes of 26 patients)

| Characteristic                      | No. (%) or mean ± SD |
|-------------------------------------|-----------------------|
| Age (years)                         | 59.86 ± 13.1          |
| Sex                                 |                       |
| Male                                | 10 (38.5%)            |
| Female                              | 16 (61.5%)            |
| Race                                |                       |
| White                               | 19 (73.1%)            |
| Black                               | 2 (7.7%)              |
| Asian                               | 3 (11.5%)             |
| Other                               | 2 (7.7%)              |
| Ethnicity                           |                       |
| Hispanic                            | 4 (15.4%)             |
| Non-Hispanic                        | 22 (84.6%)            |
| Central corneal thickness (µm)      | 551.14 ± 32.1         |
| Intraocular pressure (mm Hg)        | 17.43 ± 4.1           |
| Spherical equivalent (D)            | -1.07 ± 2.5           |
| HFA 24-2 visual field index (%)     | 99.22 ± 0.9           |
| HFA 24-2 MD (dB)                    | 0.00 ± 1.1            |
| HFA 24-2 PSD (dB)                   | 1.56 ± 0.4            |
| HFA 10-2 MD (dB)                    | 0.04 ± 0.9            |
| HFA 10-2 PSD (dB)                   | 1.19 ± 0.2            |
| SD-OCT retinal nerve fiber layer (µm)| 90.23 ± 9.8           |
| SD-OCT average GCL/IPL thickness (µm) | 79.19 ± 9.6        |
| SD-OCT minimum GCL/IPL thickness (µm) | 76.36 ± 6.4        |

HFA, Humphrey visual field analyzer; SD-OCT, spectral domain optical coherence tomography; MD, mean deviation; PSD, pattern standard deviation; GCL, ganglion cell layer; IPL, inner plexiform layer.
Hierarchical Stepwise Regression Analysis of ssPERG Parameters and RNFL Thickness Measurements

A hierarchical regression analysis was performed to examine the effect of ssPERG parameters (Mag, MagD, and the MagD/Mag ratio) in predicting RNFL thickness change after controlling glaucoma risk factors (age, sex, CCT, IOP, and SE). Mag significantly predicted superior ($\beta = 20.714$; $p < 0.001$) and inferior ($\beta = 13.024$; $p = 0.013$) RNFL quadrants, as well as avRNFL thickness ($\beta = 9.174$; $p = 0.002$; Table 3). MagD also significantly predicted superior ($\beta = 19.325$; $p < 0.001$) and inferior ($\beta = 11.282$; $p = 0.005$) RNFL quadrants and avRNFL thickness ($\beta = 8.116$; $p = 0.001$; Table 4). Mag and MagD did not significantly predict temporal and nasal RNFL quadrant thickness change. The MagD/Mag ratio was not predictive of avRNFL thickness or thickness of any specific RNFL quadrants change (Table 5). Regression plots for ssPERG parameters predicting average RNFL thickness can be seen in Supplemental Figure 1.

Hierarchical Stepwise Regression Analysis of ssPERG Parameters and GCL/IPL Thickness Measurements

A hierarchical regression analysis was performed to examine the effect of ssPERG parameters in predicting GCL/IPL thickness after controlling for age, sex, CCT, IOP, and SE. Mag ($\beta = 11.226$; $p = 0.001$), temporal ($\beta = 9.368$; $p < 0.001$), inferior ($\beta = 6.497$; $p = 0.007$), and nasal ($\beta = 10.826$; $p = 0.005$; Table 3). Mag was not significantly predictive of any outer macroa sectors. MagD was significantly correlated with all inner macular sectors ($r > 0.451$, $p < 0.007$) and the outer nasal sector ($r = 0.028$, $p = 0.875$), inner nasal sector ($r = 0.425$, $p = 0.012$) and outer nasal sector ($r = 0.302$, $p = 0.082$) sectors. There was no significant correlation between ssPERG parameters and central subfield thickness ($\mu m$), cube volume ($mm^3$), and cube average thickness ($\mu m$).

Hierarchical Stepwise Regression Analysis of ssPERG Parameters and Macular Thickness

A hierarchical regression analysis was performed to examine the effect of ssPERG parameters predicting average RNFL thickness can be seen in Supplemental Figure 1.
Pattern Electroretinogram Parameters and their Associations

It has previously been established that ssPERG parameters Mag and MagD had significant associations with optic nerve head morphology and could be detected on SD-OCT. Additionally, MagD had significant correlations with a greater number of GCL/IPL sector thicknesses than Mag, further indicating that ssPERG could be detecting RGC signaling dysfunction in these sectors prior to RGC death.

Patients with optic disk abnormalities or known glaucoma risk factors with normal perimetry and SD-OCT findings may be regarded as “glaucoma suspects” due to their increased risk for glaucomatous optic nerve changes in the future. All the patients in this study had SD-OCT values for RNFL, GCL/IPL, and macular thickness that fell within the normative data ranges specified by the Zeiss Cirrus OCT manual.22 Previous studies of glaucoma suspects have reported mean avRNFL thicknesses ranging from 74.8–101 μm, which is in agreement with the mean avRNFL thickness in this study.23-25 Patients in this study also experienced no visual field deficits but had clinical features or risk factors that would classify them as glaucoma suspects. Previous studies have identified a reduction in ssPERG amplitude prior to visual field changes in glaucoma suspects.14,26 Bode et al. found that the ssPERG ratio had a sensitivity and specificity of 75% and 76%, respectively and that ssPERG could detect glaucoma in patients 4 years before the onset of any visual field changes.27,28

Table 3: Associations of Magnitude with OCT variables, controlling for age, sex, intraocular pressure, central corneal thickness, and spherical equivalent

| Dependent variable | ΔR² | β  | 95% CI           | p     |
|--------------------|-----|----|-----------------|-------|
| RNFL thickness by quadrants |      |    |                 |       |
| Superior           | 0.244 | 20.714 | [10.076–31.352] | <0.001b |
| Temporal           | 0.016 | 4.025  | [-4.504–12.553] | 0.343  |
| Inferior           | 0.127 | 13.024 | [2.979–23.069]  | 0.013a  |
| Nasal              | 0.003 | 1.011  | [-5.053–7.075]  | 0.736  |
| Average RNFL thickness | 0.158  | 9.174  | [3.488–14.860]  | 0.002b  |
| GCL/IPL thickness by sector |      |    |                 |       |
| Superonasal        | 0.083 | 4.968  | [0.448–9.487]   | 0.032a  |
| Superior           | 0.157 | 6.469  | [2.122–10.816]  | 0.005b  |
| Superotemporal     | 0.147 | 5.371  | [1.545–9.198]   | 0.007b  |
| Inferotemporal     | 0.073 | 3.759  | [0.355–7.163]   | 0.032b  |
| Inferior           | 0.052 | 3.666  | [0.278–7.054]   | 0.035a  |
| Inferonasal        | 0.076 | 4.470  | [0.623–8.317]   | 0.024b  |
| Average GCL/IPL thickness | 0.113  | 5.097  | [0.946–9.249]   | 0.018b  |
| Minimum GCL/IPL thickness | 0.189  | 6.342  | [2.325–10.359]  | 0.003b  |
| Macular thickness by sector |      |    |                 |       |
| Inner superior     | 0.212 | 11.226 | [4.755–17.698]  | 0.001b  |
| Inner temporal     | 0.224 | 9.368  | [4.399–14.337]  | 0.001b  |
| Inner inferior     | 0.144 | 6.497  | [1.933–11.061]  | 0.007b  |
| Inner nasal        | 0.189 | 10.826 | [3.571–18.081]  | 0.005b  |

* p-values <0.05 were considered statistically significant; † p-values <0.01; ‡ Analysis performed using transformed Magnitude; OCT, optical coherence tomography; GCL, ganglion cell layer; IPL, inner plexiform layer; β, unstandardized beta

The ability of MagD, but not Mag, to predict RGC dysfunction suggests that in glaucoma suspects, there may be early signs of RGC synaptic dysfunction in the outer macula prior to any RGC loss that can be detected on SD-OCT. Additionally, MagD had significant correlations with a greater number of GCL/IPL sector thicknesses than Mag, further indicating that ssPERG could be detecting RGC signaling dysfunction in these sectors prior to RGC death.

It has been reported that amplitude, measured in this study with Mag, and latency, measured in this study as MagD, are essentially uncoupled in ssPERG and each represents separate aspects of RGC activity.21 The authors hypothesized that phase delays in the absence of amplitude were secondary to synaptic dysfunction.21
### Table 4: Associations of MagnitudeD with OCT variables, controlling for age, sex, intraocular pressure, central corneal thickness, and spherical equivalent

| Dependent variable | \( \Delta R^2 \) | \( \beta \) | 95% CI | \( p \) |
|--------------------|-----------------|-------------|--------|--------|
| **RNFL thickness by quadrants** | | | | |
| Superior | 0.259 | 19.325 | [11.193–27.458] | <0.001<sup>b</sup> |
| Temporal | 0.004 | 1.528 | [-5.041–8.096] | 0.639 |
| Inferior | 0.127 | 11.282 | [3.652–18.913] | 0.005<sup>b</sup> |
| Nasal | 0.006 | 1.212 | [-3.388–5.812] | 0.596 |
| Average RNFL thickness | 0.156 | 8.116 | [3.773–12.459] | 0.001<sup>b</sup> |
| **GCL/IPL thickness by sector** | | | | |
| Superonasal | 0.107 | 4.779 | [1.435–8.123] | 0.006<sup>b</sup> |
| Superior | 0.190 | 6.200 | [3.042–9.357] | <0.001<sup>b</sup> |
| Superotemporal | 0.149 | 4.671 | [1.785–7.557] | 0.002<sup>b</sup> |
| Inferotemporal | 0.097 | 3.759 | [1.264–6.255] | 0.004<sup>b</sup> |
| Inferior | 0.083 | 3.944 | [1.495–6.393] | 0.002<sup>b</sup> |
| Inferonasal | 0.093 | 4.267 | [1.382–7.152] | 0.005<sup>b</sup> |
| Average GCL/IPL thickness | 0.174 | 5.407 | [2.438–8.376] | 0.001<sup>b</sup> |
| Minimum GCL/IPL thickness | 0.251 | 6.359 | [3.508–9.210] | <0.001<sup>b</sup> |
| **Macular thickness by sector** | | | | |
| Inner superior | 0.210 | 9.261 | [4.424–14.098] | <0.001<sup>b</sup> |
| Inner temporal | 0.196 | 7.077 | [3.224–10.929] | 0.001<sup>b</sup> |
| Inner inferior | 0.132 | 5.283 | [1.781–8.785] | 0.004<sup>b</sup> |
| Inner nasal | 0.201 | 8.984 | [3.535–14.434] | 0.002<sup>b</sup> |

<sup>a</sup> p-values <0.05 were considered statistically significant; <sup>b</sup> p-values <0.01; <sup;c</sup> Analysis performed using transformed MagnitudeD; OCT, optical coherence tomography; GCL, ganglion cell layer; IPL, inner plexiform layer; \( \beta \), unstandardized beta

### Table 5: Associations of MagD/Mag ratio with OCT variables, controlling for age, sex, intraocular pressure, central corneal thickness, and spherical equivalent

| Dependent variable | \( \Delta R^2 \) | \( \beta \) | 95% CI | \( p \) |
|--------------------|-----------------|-------------|--------|--------|
| **RNFL thickness by quadrants** | | | | |
| Superior | 0.054 | 38.260 | [-6.449–82.968] | 0.091 |
| Temporal | 0.001 | 2.668 | [-26.863–32.200] | 0.855 |
| Inferior | 0.034 | 25.548 | [-12.007–63.103] | 0.175 |
| Nasal | 0.005 | 4.822 | [-15.810–25.454] | 0.637 |
| Average RNFL thickness | 0.032 | 15.851 | [-6.692–38.393] | 0.162 |
| **GCL/IPL thickness by sector** | | | | |
| Superonasal | 0.079 | 18.157 | [2.919–33.395] | 0.021<sup>a</sup> |
| Superior | 0.140 | 23.473 | [8.763–38.182] | 0.003<sup>b</sup> |
| Superotemporal | 0.073 | 14.703 | [0.426–28.981] | 0.044<sup>a</sup> |
| Inferotemporal | 0.090 | 16.600 | [4.976–28.223] | 0.007<sup>b</sup> |
| Inferior | 0.072 | 16.400 | [5.875–26.925] | 0.003<sup>b</sup> |
| Inferonasal | 0.060 | 15.321 | [1.972–28.669] | 0.026<sup>a</sup> |
| Average GCL/IPL thickness | 0.190 | 25.127 | [12.278–37.976] | <0.001<sup>b</sup> |
| Minimum GCL/IPL thickness | 0.211 | 25.965 | [12.689–39.241] | <0.001<sup>b</sup> |
| **Macular thickness by sector** | | | | |
| Inner superior | 0.146 | 32.609 | [11.111–54.106] | 0.004<sup>b</sup> |
| Inner temporal | 0.106 | 23.418 | [4.646–42.190] | 0.016<sup>a</sup> |
| Inner inferior | 0.050 | 14.635 | [-2.512–31.781] | 0.092 |
| Inner nasal | 0.109 | 28.510 | [3.227–53.792] | 0.028<sup>a</sup> |

<sup>a</sup> p-values <0.05 were considered statistically significant; <sup>b</sup> p-values <0.01; <sup;c</sup> Analysis performed using transformed MagD/Mag ratio; Mag, Magnitude; MagD, Magnitude D; OCT, optical coherence tomography; GCL, ganglion cell layer; IPL, inner plexiform layer; \( \beta \), unstandardized beta
suggestive early glaucomatous changes despite normal OCT and perimetry.

The correlation between reduced ssPERG parameters and reduced total macular thickness found in this study is also in line with previous studies that have found reduced macular thickness in patients with glaucoma.38,39 It has been suggested that this is secondary to the thinning in the GCL/IPL layers.39 The macular region has been reported to contain approximately 50% of all RGCs, which are reduced in glaucoma, and there is a strong association between macular RGC density and GCL/IPL thickness.40,41 Interestingly, ssPERG had significant correlations with all the inner sector thicknesses and had a much weaker correlation with the outer sector thicknesses of the macula in this study. It is plausible that there may be a higher concentration of RGCs in the inner three-millimeter region of the macula measured by the Zeiss Cirrus OCT machines. A previous study by Hood et al. found that the GCL/IPL layer was thickest at the five degrees radius point from the macula, which appears to align with the inner region measured by the Zeiss Cirrus OCT.42 Therefore, the inner sectors would experience greater decreases in thickness with greater loss of RGCs compared to outer sectors, leading to more significant correlations seen on ssPERG.

Like all retrospective studies, there are limitations to this current study due to potential biases in patient selection. The sample size of this study was also relatively small. Therefore, larger, prospective investigations should be performed to further confirm the findings in this study. There is also a lack of a standardized international

Figs 1A to C: Relationships between pattern electroretinogram parameters and average retinal nerve fiber layer (avRNFL) thickness after adjusting for age, sex, intraocular pressure, central corneal thickness, and spherical equivalent. (A) Magnitude (R^2 = 0.547); (B) MagnitudeD (R^2 = 0.644); (C) MagnitudeD/Magnitude ratio (R^2 = 0.507)
reference range for ssPERG measurements and there are several subjective factors that influence the Mag, MagD, and MagD/Mag ratio including the patient’s ability to concentrate and adequate eye fixation. Despite these limitations, the findings of this study suggest that Mag, MagD, and the MagD/Mag ratio are sensitive markers of early glaucomatous changes and can be predictive of RNFL, GCL/IPL, and macular thinning in glaucoma suspects.

**Conclusion**

Abnormalities in ssPERG parameters, an indication of RGC dysfunction, are associated with structural damage visualized on SD-OCT. However, even when SD-OCT thickness measurements fall within normal ranges, glaucoma suspects can still display functional changes. Functional changes preceded structural damage. These abnormalities on ssPERG in GS suggest a potential increased risk of future glaucoma development. Therefore, it is important to monitor these patients prior to any detectable SD-OCT or VF changes to provide early interventions when the functional damage is possibly reversible.

**Clinical Significance**

In this study of glaucoma suspects, ssPERG had significant correlations with RNFL, GCL/IPL, and macular thicknesses. ssPERG was also predictive of these changes as well in linear regression analysis. Therefore, ssPERG may serve as a valuable tool for early detection of disease and monitoring disease progression in glaucoma suspects.

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