Glaucoma With Early Visual Field Loss Affecting Both Hemifields and the Risk of Disease Progression

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Objective: To evaluate whether damage to both hemifields in glaucomatous eyes predicts more rapid disease progression than does single-hemifield involvement.

Methods: We reviewed the medical records of 43,660 consecutive patients. Eyes with glaucomatous optic neuropathy, 10 or more Swedish Interactive Threshold Algorithm standard 24-2 visual fields in at least 5 years, and mean deviation (MD) smaller than −6.0 dB were included. Pointwise linear regression was used to determine progression. Cox proportional hazards analysis was used to calculate risk of progression based on different baseline covariates.

Results: We enrolled 205 eyes (205 patients; mean [SD] age, 64.2 [11.0] years; follow-up, 6.5 [1.8] years; number of visual fields, 12.3 [2.9]). Patients were divided into 3 groups: initial superior defect (group A; n=79; MD, −3.4 [1.9] dB), initial inferior defect (group B; n=61; MD, −3.4 [1.8] dB), and both hemifields affected (group C; n=65; MD, −4.2 [1.5] dB). Group C progressed faster than did groups A and B (P < .02). Multivariate analysis showed significant effect of higher baseline intraocular pressure, thinner central corneal thickness, and initial damage to both hemifields.

Conclusions: Initial damage to both hemifields increases the risk of glaucoma progression. More aggressive therapy should be considered for these eyes.

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was defined as a glaucoma hemifield test result outside normal limits on at least 2 consecutive baseline VF tests and the presence of at least 3 contiguous test points in the same hemifield on the pattern standard deviation plot at $P < .010$ with at least one at $P < .005$, excluding points on the edge of the field and those directly above and below the blind spot. If the same criteria were observed across the horizontal meridian, both hemifields were considered to be affected. The 2 baseline tests required reliability indexes better than 25% to be included. Other inclusion criteria were baseline MD smaller than $-6.0$ dB, minimum follow-up of 5 years, and the absence of ocular disorders other than glaucoma likely to affect the VF. If both eyes met the inclusion criteria, the one with the smallest MD was chosen.

Pointwise linear regression (PLR) analysis was performed using Progressor software (version 3.3; Medisoft Inc, London, England), providing slopes of progression globally and locally for each point and its level of significance ($P$ values). A gaussian filter (based on a $3 \times 3$ test point grid) was also applied to reduce measurement variability without recourse to additional testing or exclusion of “noisy” tests; thus, all available VF tests except the 2 baseline tests were included in the analysis irrespective of reliability criteria. Progression was defined as the presence of a test point with a slope of sensitivity across time greater than 1 dB per year, with $P < .01$. For edge points, a stricter slope criterion of a greater than 2 dB loss per year (also with $P < .01$) was used. All the patients were familiar with automated perimetry and had undergone a minimum of 2 VF tests before study enrollment.

Demographic and ocular characteristics of enrolled patients were recorded at baseline. Central corneal thickness (CCT) (DGH-550; DGH Technology Inc, Exton, Pennsylvania) was calculated using an average of 5 measurements. The MD value and the number of abnormal points ($P < .01$) of the pattern deviation plot from the baseline VF test were used in the analysis to assess baseline functional damage.

Comparisons of continuous and categorical variables between groups were performed using analysis of variance with Bonferroni post hoc analysis and the $\chi^2$ test, respectively. Because eyes with both hemifields affected were more likely to have a larger number of abnormal points at baseline, a general linear model was used to adjust the rate of progression to the MD values and the number of abnormal points. Other variables analyzed as potential risk factors were age, baseline intraocular pressure (IOP), CCT, baseline MD, and the presence of exfoliation syndrome, which have been previously reported to be risk factors for disease progression.

Hazard ratios (HRs) for the association between different variables and VF progression were obtained using Cox proportional hazards models. We used HRs from univariate models, which do not adjust for other covariates, and adjusted HRs from multivariate models. Only variables with $P < .10$ in the univariate analysis were entered into the multivariate analysis.

Because we compared the rates of VF progression between groups using single-hemifield damage (superior or inferior) vs those with damage to both hemifields, a sample size calculation determined that a minimum of 177 eyes (35 eyes per group) was required to detect a 20% difference in rates of progression among 3 groups, with a power of 80% and type I error of 5%. Similarly, selecting a cohort of patients with greater than 10 VF tests, performing a mean of 2 examinations per year, would provide 80% power to detect significant VF change in this population. Statistical analysis was performed using a software package (SPSS version 17.0; SPSS Inc, Chicago, Illinois).

**RESULTS**

Two hundred five eyes (205 patients) met the entry criteria. The mean (SD) patient age was 64.2 (11.0) years; 58.8% were women and 86.0% were of European descent. Mean follow-up was 6.5 (1.8) years, and the mean number of VF tests was 12.3 (2.9).

Patients were divided into 3 groups: initial superior defect (group A, $n = 79$), initial inferior defect (group B, $n = 61$), and damage to both hemifields (group C, $n = 65$) (Figure). Patients received a variety of glaucoma treatments during the study. Table 1 summarizes the baseline characteristics of each group. The MD value and the number of abnormal points in group C were greater than those in the other groups ($P < .02$ for both). There were more eyes with exfoliation glaucoma in the group with superior hemifield damage ($P < .01$).

Group C progressed faster (mean, 0.90 [0.9] dB per year) than did group A (0.52 [0.8] dB per year) and group B (0.33 [0.5] dB per year). Because the MD and the number of points with $P < .01$ in the 3 groups differed, we adjusted the mean slope of the groups to these covariates (general linear model), and the differences remained significant ($P < .025$). Forty-three eyes (54.4%) in group A reached a progression end point compared with 17 (27.9%) in group B and 45 (69.2%) in group C ($P < .01$). Group C had almost twice as many eyes with a fast rate of progression ($> 1.5$ dB per year) as the other 2 groups combined (Table 2).

In the univariate model, damage to both hemifields was associated with an increased risk of VF progression (HR, 1.58; 95% confidence interval, 1.07-2.32; $P = .02$). Elevated baseline IOP and thinner CCT were also associated with a greater chance of reaching a progression end point ($P < .01$) (Table 3). Older age was associated with disease progression, although this was not significant ($P = .10$). After adjusting for these covariates, baseline damage to both hemifields remained a significant predictor of progression (HR, 1.62; 95% confidence interval, 1.09-2.39; $P = .01$) (Table 4). The presence of this pattern of damage increased the risk of future field loss by 62%, whereas each 1-mm Hg higher baseline IOP increased the risk by 7% and each 40-µm decrease in CCT increased the risk by 27%.

**COMMENT**

Glaucoma is a multifactorial disease that results in different patterns and rates of progression for different individuals. An improved understanding of risk factors at all stages of disease is critical to estimating the risk of future disease progression for a specific affected individual. The results of the present study confirm that initial damage to both visual hemifields connotes a worse prognosis than does more localized damage limited to one hemifield, even when there is early VF loss (MD smaller than $-6.0$ dB). This involvement of both hemifields is an independent predictor of more rapid future VF injury in eyes with early functional damage and greatly increases the risk of progression.

In agreement with other studies, we found a significant role for higher baseline IOP and thinner CCT as risk factors for progression. The continued importance of these risk factors in treated patients with glaucoma is particularly important for physicians who must make clini-
cal decisions for patients who may not precisely resemble the individuals who were enrolled in masked, prospective, longitudinal clinical trials. Despite an initial mean baseline IOP of 17 mm Hg in eyes with early VF loss, the risk of progression increased 7% for each additional 1 mm Hg. A thinner CCT was also a risk factor for progression, in-

Figure. Examples of the 3 patterns of baseline visual field damage evaluated in the study. A, Single superior defect. B, Single inferior defect. C, Both hemifields affected, and the points are adjacent. D, Both hemifields affected, and the points are not adjacent. Note that, for all the examples, there was good agreement between the total and pattern deviation plots, and the visual field damage was mild (mean deviation no more than −6.0 dB). GHT indicates glaucoma hemifield test; MD, mean deviation; and PSD, pattern standard deviation.

Table 1. Baseline Characteristics of the Study Population

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Group A (n=79)</th>
<th>Group B (n=61)</th>
<th>Group C (n=65)</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (SD), y</td>
<td>63.0 (10.3)</td>
<td>63.6 (11.1)</td>
<td>66.4 (11.4)</td>
<td>.13a</td>
</tr>
<tr>
<td>European ancestry, No. (%)</td>
<td>69 (87.3)</td>
<td>49 (80.3)</td>
<td>59 (90.8)</td>
<td>.22b</td>
</tr>
<tr>
<td>Diagnosis, No. (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>POAG</td>
<td>43 (54.4)</td>
<td>20 (25.3)</td>
<td>16 (20.3)</td>
<td>&lt;.01</td>
</tr>
<tr>
<td>XFG</td>
<td>25 (31.0)</td>
<td>25 (41.0)</td>
<td>33 (50.8)</td>
<td>.27</td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td></td>
<td></td>
<td>&lt;.01</td>
</tr>
<tr>
<td>MD (SD), dB</td>
<td>−3.4 (1.9)</td>
<td>−3.4 (1.8)</td>
<td>−4.2 (1.5)</td>
<td>&lt;.02a</td>
</tr>
<tr>
<td>No. of points with P&lt;1%, mean (SD)</td>
<td>6.5 (4.2)</td>
<td>7.2 (4.4)</td>
<td>9.5 (4.2)</td>
<td>&lt;.02a</td>
</tr>
<tr>
<td>CCT, mean (SD), µm</td>
<td>540.5 (41.0)</td>
<td>533.8 (41.5)</td>
<td>542.9 (41.5)</td>
<td>.55a</td>
</tr>
<tr>
<td>Baseline IOP, mean (SD), mm Hg</td>
<td>17.5 (4.3)</td>
<td>17.7 (3.5)</td>
<td>18.8 (4.6)</td>
<td>.12a</td>
</tr>
<tr>
<td>Topical medications, mean (SD), No.</td>
<td>1.8 (1.2)</td>
<td>1.7 (1.2)</td>
<td>1.6 (1.1)</td>
<td>.66a</td>
</tr>
</tbody>
</table>

Abbreviations: CCT, central corneal thickness; IOP, intraocular pressure; MD, mean deviation; POAG, primary open-angle glaucoma; XFG, exfoliation glaucoma.

*Analysis of variance with Bonferroni post hoc analysis showed significant differences between group C and the other groups.

bBy χ² test.
increasing the risk by 27% for each 40-µm decrease. These results are consistent with those of the major prospective glaucoma clinical trials. The Early Manifest Glaucoma Trial (EMGT)\(^6\) found a 13% increased risk per each additional 1 mm Hg of IOP and a 23% increased risk per 40-µm decrease in CCT. The Diagnostic Innovations in Glaucoma Study\(^6\) also found a 7% increased risk of progression for each additional 1 mm Hg of IOP and a 62% increase for each 40-µm decrease in CCT in a group of patients with early glaucoma damage. The present study reemphasizes the role of IOP even in a treated population with statistically normal pressures.

Age was not a significant risk factor in the final multivariate model, suggesting that other factors, such as disease stage, treatment, and other covariates, may play stronger roles in predicting progression in this treated population. In its first report, the EMGT\(^7\) also found a positive association between age and progression that was not confirmed in a multivariate analysis. Unlike the EMGT,\(^6\) exfoliation syndrome was not a significant risk factor in the present study, which may be due to the lower IOPs in the cohort.

The nature of glaucoma pathogenesis and functional and structural associations in glaucoma may help explain the increased risk associated with VF loss in both hemifields. Glaucomatous functional damage usually respects the horizontal meridian and the anatomy of the retinal nerve fiber layer.\(^11\) In 1984, Mikelberg and Drance\(^18\) reviewed the pattern of VF progression using static and kinetic perimetry and found that 70% of eyes had initial damage limited to a single hemifield; at the completion of follow-up, 57% still had only single-hemifield involvement, whereas 13% had involvement of both hemifields. The most common pattern of field loss was deepening of an existing scotoma, which was later confirmed using static perimetry.\(^19\) Boden et al\(^10\) found that early glaucomatous field loss rarely crosses the horizontal midline. In a cross-sectional analysis that included patients with mild to severe glaucoma, they found a prevalence of 30% of VF defects across the horizontal meridian, 90% of which could be explained by changes at the optic nerve head (ONH) as assessed using stereophotography. The definition used in their study implied that the superior and inferior affected sectors should be adjacent to the horizontal midline (Figure, C). The present study did not require VF defects to be adjacent to the midline and contiguous, a clinical characteristic that may be found more commonly in practice (Figure, D).\(^18\) Although it has been suggested that worse functional damage (MD) is associated with an increased risk of progression,\(^6\) the present findings suggest that it is also possible to determine different levels of risk based on the extent and location(s) of the defect, even when the VF damage is mild. Despite initially having similar global field damage, damage to both hemifields suggests that more widespread structural and functional abnormality is present, which increases the susceptibility to progression.

Experimental models have been developed to try to clarify the pathogenesis of glaucomatous damage and progression. There is strong evidence that damage to the retinal ganglion cell axons, which ultimately converge to the ONH, is the key cause of vision loss in glaucoma.\(^21\)–\(^23\) Burgoyne and Downs\(^24\) reviewed this issue and proposed that alterations in ONH biomechanics underlie the clinical behavior and likely increased susceptibility of the ONH. That is, a more damaged optic nerve would be more susceptible to future damage. Quigley et al\(^23\) suggested that the structure of the lamina cribrosa is an important determinant of the degree of susceptibility to damage by elevated IOP. Jonas et al\(^25\) showed a correlation between the progression of VF defects and the morphologic features of the lamina cribrosa and suggested that a larger single pore area increased glaucoma susceptibility in the inferior and superior disc regions.\(^26\)

We hypothesized that the presence of damage to both hemifields may reflect greater overall optic nerve susceptibility to glaucoma (in both the superior and inferior poles rather than localized to one location), which resulted in the accelerated rate of progression found in this study. Because we did not address this issue directly, further studies to assess structural characteristics of the ONH and lamina cribrosa in eyes with faster progression rates are necessary to confirm this hypothesis.

To support this hypothesis, Demirel et al\(^27\) recently assessed the role of baseline perimetry data in predicting future progression. They found that depressed VF...
locations close to the midline, mostly in the superior and inferior nasal sectors, are most predictive of future progression. If this is the case, eyes with abnormal points in both hemifields would have a summed effect of susceptibilities from different VF sectors. Similarly, Pascual et al28 showed a spatial relationship between initially defective locations found at baseline and those found on subsequent testing. The authors described superior defects related to progression in the superior field, resembling the nerve fiber bundle patterns, whereas the inferior defects did not show clearly specific patterns of progression. These studies may help explain why fewer eyes with baseline damage limited to the inferior hemifield reached a progression end point in the present study.

We chose to use PLR rather than event analysis in this study for several reasons. First, the commercially available software provides automated values that can be easily determined and reproduced at different medical centers. Second, in contrast to other methods that have been proposed for major clinical trials, PLR can determine the rate of VF loss globally, by sector, or at individual points with decreased subjectivity.29 Finally, the larger number of VFs required for PLR enhances specificity.30

This study has several limitations. Enrolling eyes with mild functional damage limits the conclusions to this specific population. However, eyes with moderate or severe VF damage often have damage to both hemifields, making identification of a comparable group with damage to one hemifield more difficult. The use of PLR may have provided a more sensitive method of progression than is typically used in clinical practice. As in most clinical studies, we analyzed the predictive value of baseline characteristics (level of damage, CCT, age, IOP, and exfoliation glaucoma).6-10 However, reassessment and reevaluation of risk factors during treatment (interrater risk factors) might provide more important information than the initial baseline risk factor assessment. In clinical practice, physicians constantly adjust risk profiles as information becomes available (eg, advancing age and a major cardiovascular event), and the use of a continuous and dynamic measure of progression (expressed as rates) may be particularly helpful in these circumstances. Last, the retrospective nature of this study design and the tertiary care setting create certain inherent biases in patient selection. The long-term follow-up and the similar characteristics among the study groups (age, IOP, CCT, and treatment) serve to mitigate potential bias and suggest that such bias likely did not significantly affect the results. Despite the retrospective nature of this study, the consonance of these results with the major prospective clinical trials serves to confirm the validity of the data set.

In a treated glaucoma population, assessing other variables that could be associated with faster rates of disease progression may help direct future management and aggressiveness of treatment. In these patients with early and similar baseline VF damage, eyes with defects involving both hemifields progressed more quickly and were more likely to reach a predefined end point than were those with a single affected hemifield. For the practicing physician, these findings suggest that a lower target IOP may be warranted for patients with initial, reproducible damage extending to both hemifields.

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REFERENCES


Cavernous Hemangioma of the Retina
Stephen Jae Kim, MD

Figure 1. Composite color fundus photograph of the left eye of a 13-year-old healthy white girl who presented with vitreous hemorrhage. The photograph shows the characteristic appearance of a cavernous hemangioma of the retina with thin-walled, saccular aneurysms partly covered by a fine glial membrane.

Figure 2. Composite fluorescein angiogram demonstrating grapelike clusters of aneurysms, some with sedimented blood in the lower half and clear serum in the upper half (arrow), indicating low perfusion. There was a corresponding ipsilateral cavernous malformation in the corpus callosum.