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**Original citation**: Artes, P. H., O'Leary, N., Nicolela, M. T., Chauhan, B. C. & Crabb, D. P. (2014). Visual Field Progression in Glaucoma What Is the Specificity of the Guided Progression Analysis?. Ophthalmology, 121(10), pp. 2023-2027. doi: 10.1016/j.ophtha.2014.04.015

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# Visual field progression in glaucoma: What is the specificity of the Guided Progression Analysis?

Short title	Specificity of the Guided Progression Analysis
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Length	2,200 words, 3 figs (+ 3 in supplementary material), & 2 tables
Section Code	Glaucoma
Prev. Pres.	Association for Research in Vision and Ophthalmology meeting 2011
Keywords	glaucoma, visual field, progression, permutation, Guided Progression Analysis, Glaucoma Change Probability
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Support	Glaucoma Research Foundation (PHA & DPC) Grant # MOP-11357, Canadian Institutes for Health Research (BCC)
Disclosures	None
Précis	In patients with glaucoma examined many times over a short period of time, we show that the specificity of the Guided Progression Analysis of the Humphrey Field Analyzer varies substantially between patients.

# 1 Abstract

Purpose: To estimate the specificity of the Guided Progression Analysis (GPA, Carl Zeiss Meditec,
 CA), in individual glaucoma patients.

4 **Design:** Observational cohort study.

5 **Participants:** Thirty patients with open-angle glaucoma.

Methods: In 30 patients with open-angle glaucoma, one eye (median Mean Deviation [MD], -2.5
dB, interquartile range -4.4 to -1.3 dB) was tested 12 times over 3 months (Humphrey Field
Analyzer, Carl Zeiss Meditec; SITA Standard, 24-2). "Possible progression" and "likely progression"
were determined with the Guided Progression Analysis (GPA). These analyses were repeated after
the order of the tests had been randomly re-arranged (1000 unique permutations).

Main Outcome Measures: Rate of false-positive alerts of "possible progression" and "likely
 progression" with the Guided Progression Analysis.

**Results:** On average, the specificity of the GPA "likely progression" alert was high—for the entire 13 sample, the mean rate of false-positive alerts after 10 follow-up tests was 2.6%. With "possible 14 progression", the specificity was considerably lower (false-positive rate, 18.5%). Most importantly, 15 the cumulative rate of false-positive alerts varied substantially among patients, from <1% to 80% 16 with "possible progression", and from <0.1% to 20% with "likely progression". Factors associated 17 with false-positive alerts were visual field variability (standard deviation of MD, Spearman's 18 rho=0.41, p<0.001) and the reliability indices (proportion of false-positive and false-negative 19 responses, fixation losses, rho>0.31,  $p \le 0.10$ ). 20

Conclusions: On average, progression criteria currently employed in the GPA have high specificity, but some patients are much more likely to show false-positive alerts than others. This must be considered when the GPA is used in clinical practice, where specificity needs to be controlled for individuals rather than for large groups of patients.

# 25 Introduction

In patients with glaucoma, accurate decisions on visual field progression are a prerequisite of good clinical management.<sup>1, 2</sup> Visual fields have complex properties, and therefore progression is best judged with the help of software such as the Guided Progression Analysis (GPA) of the Humphrey Field Analyser (HFA, Carl Zeiss Meditec, Dublin, CA). <sup>3 4-6</sup>

The GPA compares each test result, point by point, to values from two earlier baseline tests. Points 30 are highlighted on a probability plot if changes exceed the typical measurement variability derived 31 from a group of stable glaucoma patients. If such changes occur at 3 or more points, and in 2 32 consecutive follow-up tests, the GPA raises an alert of "possible progression"; if they occur in 3 33 consecutive tests an alert of "likely progression" is raised. Criteria similar to "likely progression" 34 were used in the Early Manifest Glaucoma Trial, and the GPA has subsequently been widely 35 adopted in clinical practice and research.<sup>7, 8</sup> Previous studies have shown that the analysis agrees 36 reasonably closely with the subjective judgement of expert clinicians,<sup>9, 10</sup> and some authors have 37 used the GPA as a reference standard for functional change in glaucoma.<sup>11, 12</sup> 38

Owing to the fundamental role of visual field progression in the clinical management of glaucoma, 39 it is important to know how often the GPA raises alerts of "possible progression" and "likely 40 progression" in the absence of genuine change, i.e. false-positives. We previously demonstrated that 41 the GPA is likely to have high specificity, on average.<sup>13</sup> However, the analysis is based on a 42 statistical model of typical variability inferred from a group of stable patients-it does not take into 43 account that some patients are more reliable test-takers than others.<sup>14</sup> Because the reproducibility of 44 visual fields varies more than 2-fold between individuals with the same degree of damage (Artes et 45 al, Invest. Ophthalmol. Vis. Sci. 54: E-Abstract 2630), the limits for significant change of the GPA 46 are likely to be too wide for patients who are have relatively low variability, and too narrow for 47 those with relatively larger variability. 48

In this study, we aim to investigate how the specificity of the GPA varies between individual
 patients. For this purpose, we tested a group of patients multiple times, over a short period of time
 during which a clinically meaningful change was unlikely to have taken place.

# 52 Methods

#### 53 Patients

Thirty patients were recruited from the glaucoma clinics at the Queen Elizabeth Health Sciences 54 Centre in Halifax, Nova Scotia. Inclusion criteria were a clinical diagnosis of open-angle glaucoma, a 55 Mean Deviation (MD) better than -15.0 dB in at least one eye, absence of ocular or systemic 56 pathology known to reduce visual field sensitivity, and the ability and willingness to participate for 57 12 consecutive weekly sessions. All patients were experienced with static automated perimetry and 58 had performed at least 5 visual field tests before the study started. They had well controlled levels of 59 intraocular pressure as judged by their physician (MTN). In accordance with the Declaration of 60 Helsinki, the institutional research ethics board approved the protocol, and all patients gave written 61 informed consent. 62

#### 63 Tests

Patients attended 12 weekly sessions over a period of 3 months. During each session, the study eye
 was examined with program 24-2 SITA-Standard of the HFA.

#### 66 Analysis

#### 67 Guided Progression Analysis (GPA)

The GPA is based on principles previously described as Glaucoma Change Probability.<sup>14</sup> In brief, a visual field baseline is calculated from the first two tests, and each subsequent test is then compared, point by point, to this baseline. If the difference in pattern deviation exceeds the retest variability estimated from a group of stable patients, the corresponding location is flagged on a probability map by an open triangle. Half-filled and solid triangles signify change on two or three consecutive follow-up tests, respectively. The GPA gives alerts of "possible progression" and "likely progression" when there are three or more locations with half-filled or solid triangles, respectively.

#### 76 Permutation

The premise of our study was that a meaningful change was unlikely to have taken place during the 77 short period of 3 months during which the 12 tests were performed. Under this assumption, a GPA 78 alert of "possible progression" or "likely progression" in the series of 12 tests could be regarded as a 79 false-positive event. Furthermore, by assuming that the order of the tests could be treated as 80 arbitrary, a large number of permutations could be generated from the originally observed series, by 81 randomly changing the order of the tests in the sequence. In this way, the probability of observing a 82 false-positive "possible progression" or a "likely progression" alert could be derived specifically for 83 each individual patient. For each patient, we submitted 1000 permuted series to Carl Zeiss Meditec 84 who generated the GPA results as they would appear on the instrument's output. 85

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#### 87 Analysis

Individually for each patient, we determined the proportion of series in which at least one alert of 88 "possible progression" or "likely progression" had been raised, at the 4th through the 12th test, 89 across the 1000 permuted series. Similarly, we determined the cumulative probability of 90 encountering at least one "possible progression" or "likely progression" alert in a patient's series of 91 12 tests (2 baseline and 10 follow-up tests). Confidence intervals for the mean proportion of false-92 positive alerts across the group of patients were determined by bootstrap (n=10,000 samples). We 93 also investigated the association between the cumulative probability of encountering at least one 94 progression alert after 12 tests (2 baseline and 10 follow-up tests) and the MD, the standard 95 deviation (SD) of the MD, and to indices of patient reliability (false-positive and false-negative 96 response errors, fixation losses, averaged across the entire series of 12 tests). All analyses were 97 performed in the open-source programming language R (R Foundation for Statistical Computing, 98 Vienna, Austria; <u>http://www.R-project.org</u>; last accessed 20 January 2014). 99

# 101 **Results**

<sup>102</sup> The median age of the patients was 69.1 years (interquartile range [IQR], 64.4 to 70.7 years).

- <sup>103</sup> Patients had early to moderate visual field damage (median MD, -2.5 dB, IQR -4.4 to -1.3 dB) as
- <sup>104</sup> illustrated in Figure 1 (available at <u>http://aaojournal.org</u>). All patients were experienced test-takers,
- and there were no clinically important learning- or practice effects—the mean MD of the 30
- patients changed by <0.1 dB between the first and last tests (Fig. 2, (available at
- $\frac{\text{http://aaojournal.org}}{\text{model}}$ . However, the variability of the MD varied by a factor >3 between patients

108 (Fig. 3).

The analysis of the randomly re-ordered test results confirmed that, on average, the specificity of 109 the GPA "likely progression" alert was high-after 10 follow-up tests (12 tests in total, including 110 the 2 baselines), the mean false-positive alert rate across the 30 patients was 2.6% (95% confidence 111 interval: 1.2%, 4.4%). The specificity of the "possible progression" alert was considerably lower-112 after 10 tests the mean false-positive rate was 18.5% (95% confidence interval: 11.5%, 26.5%) (Figs. 113 4, 5). Most importantly, however, the false-positive rate of the GPA varied substantially between 114 patients. In 11 patients (37%), no "likely progression" alerts were detected in any of the 1000 115 reordered series, and 4 of these patients also did not have a "possible progression" alert. On the 116 other hand, in one patient 80% of the reordered series contained alerts of "possible progression", 117 and 18% contained alerts of "likely progression". 118

"Possible progression" and "likely progression" alerts were more closely associated with the patient
reliability indices (false-positive and -negative response errors, fixation losses) and with visual field
variability (SD of MD) than with visual field damage as measured with MD and Pattern Standard
Deviation (PSD). However, none of these associations were sufficiently strong to predict to a useful
level of accuracy in which patients the GPA would be prone to false-positive progression alerts.
(Table 1, Figs. 6, 7; available at <a href="http://aaojournal.org">http://aaojournal.org</a>).

# 126 Discussion

The aim of our study was to investigate the specificity of the Glaucoma Progression Analysis, i.e. 127 the likelihood of encountering a "possible progression" or "likely progression" alert in a series of 128 visual fields in which no meaningful change has taken place. Stable series were established by testing 129 patients frequently over a short period during which disease progression was unlikely, such that any 130 GPA progression alert could be regarded as a false-positive event. Under the assumption that the 131 order of the tests could be randomly exchanged, we were able to estimate the rate of false-positive 132 GPA progression alerts from a large number of random permutations of the original visual field 133 series, for each individual patient. 134

Our results corroborate earlier reports of high *average* specificity with the GPA<sup>13, 15</sup> — after 12 tests, the average false-positive rate of "likely progression" alerts was <5%. With tests conducted at intervals of 6 months, a series of 12 tests would translate to approximately 5 years of follow-up, and this level of specificity appears sufficiently high for most clinical applications. However, the large variation in the GPA false-positive rates between individual patients confirmed our hypothesis that some patients are much more prone to show false-positive progression alerts than others. The high average specificity of the GPA observed in a group of patients does not apply equally to all patients.

The GPA uses a statistical model to establish, point by point, whether the differences between a 142 follow-up test and two earlier baseline tests exceed the limits of measurement variability typically 143 observed in patients with glaucoma. This model aims to account for the amount of baseline damage 144 at individual test locations, for the location within the visual field, and for the overall damage of the 145 visual field as measured by the MD index.14 The lack of a relationship between the GPA false-146 positive rate and visual field damage (as measured by MD and PSD, Fig. 6, available at 147 http://aaojournal.org) indicates that the GPA adequately compensates for the larger threshold 148 variability in damaged areas of the visual field. However, the level of damage explains less than half 149 of the variability in visual field measurements.<sup>16, 17</sup> Clearly, there are patient-related factors unrelated 150 to visual field damage that influence variability, for example the ability to sustain attention and to 151 provide consistent responses. Because the GPA uses the variability estimated from a reference 152 group of patients, the analysis is overly conservative (i.e., highly specific, but less sensitive) in 153

patients who are highly reliable test-takers, and not sufficiently conservative (i.e., more sensitive, but
 less specific) in patients with relatively larger between-test variability.

156 While there were statistically significant relationships between overall visual field variability

(measured by the SD of the MD), the reliability indices (false-positive and false-negative response

associations were too weak to be practically useful for predicting in which patients the GPA is most

errors, and fixation losses), and the likelihood of false-positive GPA progression alerts, these

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likely to produce false-positive progression alerts (Fig. 6, 7; available at <u>http://aaojournal.org</u>).

One alternative to the Glaucoma Change Probability model of the GPA is pointwise linear 161 regression (PLR), a method that has been widely discussed elsewhere.<sup>18-20</sup> PLR establishes statistical 162 significance of change at individual visual field locations by least-squares linear regression of 163 sensitivity (or deviation) over time. Other statistical models for deriving rate of change and its 164 statistical significance at single test locations have also been proposed.<sup>21-23</sup> Common to all of these 165 techniques is that the patient's own variability is estimated, obviating the need to rely on variability 166 estimates from other patients. O'Leary et al. have recently introduced a method (Permutation of 167 Pointwise Linear Regression, PoPLR) in which the statistical significance of deterioration over the 168 entire visual field is derived solely from random re-ordering (permutation) of the individual patient's 169 data, without reference to population-based reference values.<sup>24</sup> This method provides an 170 individualised statistical test of the null hypothesis that there is no negative change at any visual field 171 location, removing any between-patient variation in specificity. We believe that this method may 172 provide a useful alternative to the Glaucoma Change Probability model of the GPA, particularly 173 when more than 5 tests are available for analysis and when specificity needs to be controlled at the 174 level of the individual patient, as it must be in clinical practice. 175

Two assumptions of our study are a) that visual fields obtained over a short period of time are representative of those obtained over a longer period, and b) that any re-ordered sequence of tests could have occurred with the same likelihood as the originally observed sequence. It is likely that visual field data violate both assumptions. Variability, for example, may be higher in the long term than observed during the 12-week period of our study, and the differences between two tests obtained one after the other may be smaller than between tests at the start and the end of the sequence (serial correlation). However, while these violations may affect our estimates of specificity,

they are unlikely to have a substantial effect on the finding that the specificity of the GPA varies
considerably between patients.

In summary, we have shown that the GPA criterion of "likely progression" has high specificity on average, but that some patients are much more prone to false-positive alerts than others. Rather than discouraging clinicians from using the GPA, we hope that this report helps to avoid falsepositive decisions on progression in patients with larger-than-average variability and frequent response errors.

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# 191 Acknowledgements

The authors thank Gary Lee and Mary Durbin of Carl Zeiss Meditec for generating the GPA output
for the permuted visual fields.

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