Reclaiming the Periphery: Automated Kinetic Perimetry for Measuring Peripheral Visual Fields in Patients With Glaucoma

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PURPOSE. Peripheral vision is important for mobility, balance, and guidance of attention, but standard perimetry examines only <20% of the entire visual field. We report on the relation between central and peripheral visual field damage, and on retest variability, with a simple approach for automated kinetic perimetry (AKP) of the peripheral field.

METHODS. Thirty patients with glaucoma (median age 68, range 59–83 years; median Mean Deviation –8.0, range –16.3–0.1 dB) performed AKP and static automated perimetry (SAP) (German Adaptive Threshold Estimation strategy, 24-2 test). Automated kinetic perimetry consisted of a fully automated measurement of a single isopter (III.1.e). Central and peripheral visual fields were measured twice on the same day.

RESULTS. Peripheral and central visual fields were only moderately related (Spearman’s ρ, 0.51). Approximately 90% of test-retest differences in mean isopter radius were < ±4 deg. Relative to the range of measurements in this sample, the retest variability of AKP was similar to that of SAP.

CONCLUSIONS. Patients with similar central visual field loss can have strikingly different peripheral visual fields, and therefore measuring the peripheral visual field may add clinically valuable information.

Keywords: glaucoma, visual field, peripheral vision, automated kinetic perimetry

Since the advent of computerized visual field testing in the 1970s, almost all innovations in perimetry have focused either on improving the sensitivity to early visual field damage in glaucoma,1–6 or on increasing either efficiency7–9 or speed10 of the tests. This drive toward high diagnostic performance has led to a situation where almost all visual field tests performed in glaucoma patients are confined to the central 25–30 degrees of the visual field, an area that constitutes less than 20% of the entire field of vision.

Peripheral vision contributes to postural stability11–14 and the guidance of attention,15,16 and it is important for estimating motion from optical flow.16–18 In people with normal vision, eliminating clues from the peripheral visual field decreases postural stability,11 and patients with glaucoma rely more heavily on vestibular and proprioceptive cues to maintain balance than do healthy controls.12–14 Thus, the central visual field alone does not provide a complete picture of the patients’ real-world field of vision, and examinations of the peripheral visual field may help us to more fully understand the impact of the disease on individuals.

The peripheral visual field may also add information relevant to clinical decision making, for example, for diagnosis,19–21 disease phenotyping, and monitoring progression. For example, peripheral visual field damage has been demonstrated in 15% of glaucoma patients with normal central visual fields.22 At the other end of the spectrum, in patients with advanced damage in whom much of the central visual field may be damaged beyond the useful dynamic range of static perimetry,23 tracking peripheral vision may be useful to demonstrate stability or to uncover further deterioration.24–27

A key reason for why peripheral visual fields are not measured more often is the lack of fast and efficient automated tests. Static programs that include the periphery are available on the Humphrey Field Analyzer (HFA), Carl Zeiss Meditec, Jena, Germany) and the Octopus instruments (Haag-Streit, Köniz, Switzerland).20–22 However, threshold examinations, for example with the 60-4 test of the HFA,23,24 usually take more than 10 minutes, in part because they still rely on the classic full-threshold procedures25 rather than the more efficient techniques for threshold estimation and stimulus pacing introduced by the Swedish interactive thresholding algorithm.7 Likewise, the suprathreshold tests of these instruments have scarcely changed since the 1980s. Last, statistical tools for interpretation of peripheral perimetry (such as total- and pattern-deviation probability maps) have not been made available commercially.

Manual kinetic Goldmann perimetry,36 as introduced in 1945, is probably still the most extensively used technique for measuring peripheral visual fields. In the hands of a highly trained examiner, it is a very flexible technique, but it is difficult to standardize, difficult to quantify, and difficult to compare between different examiners. Progressively fewer centers possess the resources to perform this technique, and manufacture of the original Goldmann instrument (Haag-Streit, Küniz, Switzerland) has recently been discontinued. Semi-automated kinetic perimetry (available on the Octopus 900 perimeter, the
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In this paper, we demonstrate the large dissociation between central and peripheral visual fields in a group of patients with moderately advanced glaucoma. We report on a simple approach of repeated kinetic presentations to estimate isopter positions without interactive input from the examiner. We show that the precision of this technique is comparable to that of static perimetry of the central field and suggest further avenues for more efficient perimetry of the peripheral visual field.

METHODS

Participants

Thirty patients with open-angle glaucoma were recruited from participants of previous studies at City University, London. Patients had been recruited from the glaucoma clinics at Moorfields Eye Hospital, and inclusion criteria were a visual acuity of at least +0.30 logMAR (6/12), ametropia within ±5.00 diopter (D) equivalent sphere and ±2.50 D cylinder, and no concomitant ocular or systemic disease. Table 1 provides descriptive statistics on the patients’ age, visual acuity and contrast sensitivity. All patients were experienced in static perimetry, but none had previously performed kinetic perimetry. The study adhered to the Declaration of Helsinki; the protocol was approved by the School of Health Sciences Research Ethics Committee at City, University of London, and all patients provided written informed consent.

Examinations

Of each participant, one study eye was randomly selected and two static examinations were performed of the central visual field, along with two kinetic tests of the peripheral visual field. All tests were carried out during a single session that lasted approximately 2.5 hours including breaks. Visual acuity (Early Treatment Diabetic Retinopathy Study chart, distance 4 m) and contrast sensitivity (Pelli-Robson chart, at 1 m) were measured at the outset of the session.

Kinetic Perimetry of the Peripheral Visual Field.

Kinetic perimetry was performed with Goldmann III.e stimuli at a speed of 5 deg/s. According to Goldmann nomenclature, these stimuli are circular spots subtending a visual angle of 0.43 degrees with a luminance of 10 cd/m² (i.e., a 1.5 log unit attenuation of the 318 cd/m² maximum-intensity stimulus of Goldmann perimetry). In terms of contrast, this luminance increment corresponds to a 25-dB stimulus with the static programs of the Octopus 900 [nominal Δmax = 3185 cd/m² (10,000 asb)] and to a 21 dB stimulus with the static programs of the Octopus 900 [Δmax = 1273 cd/m² (4000 asb)].

Kinetic stimuli started well outside the normal range of visibility and moved at a speed of 5 deg/s from the periphery toward the center. The entire visual field was sampled along 16 meridians (Fig. 1). Three repetitions were performed for each vector, and the final isoper was defined by the median (middle) of the three responses. Stimuli were presented in random order. The mean radius of the isoper (MIR) was used as a global summary measure, and the reproducibility of an individual patient's answers was summarized as the median absolute deviation (MAD) of individual responses from the final isoper.

Unlike in manual kinetic Goldmann perimetry where perimetrists add additional stimuli to define the shape of isopters in areas of visual field damage, estimates that fell within the central 10 degrees of fixation were treated as missing data and would appear as a gap in the isoper (see patient u for example).

False-positive catch trials (n = 6) were stimuli presented in the far nasal periphery where they were invisible while the sound associated with the movement of the perimeter's

| Table 1. Descriptive Statistics of the Patients’ Age, Visual Acuity, and Contrast Sensitivity in the Study Eye |
|-----------------|-----------------|-----------------|-----------------|
| Mean (SD)       | Median (IQR)    | Range           |
| Age (y)         | 69 (6)          | 68 (67, 73)     | 59, 83          |
| VA (logMAR)     | +0.10 (+0.19)   | +0.07 (0.00, +0.14) | –0.20, +0.30 |
| CS (log)        | 1.60 (0.30)     | 1.65 (1.35, 1.95) | 0.60, 2.05     |

VA, visual acuity; CS, contrast sensitivity.

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FIGURE 1. Kinetic automated perimetry: Goldmann III.e stimuli were moved along 16 meridians (green arrows) at a speed of 5 deg/s. Three stimuli were shown on each meridian. Starting points of the arrows represent the start location of the stimuli. If not detected, they moved within 3 degrees of the fixation point. The dashed arrow represents the location of the six false-positive catch trials. The lightly shaded region indicates the normative response range according to Vonthein et al.42
projection system was audible. To acquaint patients with the procedure, three training stimuli were presented at the outset of the tests. The entire examination was programmed as a custom test in the XML language of the EyeSuite software. Altogether, each test consisted of a total of ~60 presentations (3 training stimuli, 48 kinetic stimuli, and 6 false-positive catch trials) and took approximately 11 minutes.

Static Automated Perimetry of the Central Visual Field. Static perimetry of the central visual field was performed with the German Adaptive Threshold Estimation (GATE) strategy with a 24-2 test pattern and a stimulus duration of 200 ms. The GATE strategy has been described previously.42,43 At the outset of the first test, thresholds are determined at four seed locations, and initial intensities at other locations are then adjusted accordingly. In subsequent tests the GATE strategy starts with stimuli slightly brighter than the thresholds estimated during the previous test and varies stimulus intensities according to a 4-2 dB staircase that normally terminates after two response reversals. In contrast to the classic full-threshold strategy,35 a maximum-intensity stimulus (0 dB) is shown if the initial stimulus has not been seen. If this stimulus is not seen, then the procedure terminates; otherwise, a stimulus 4 dB brighter than the initial intensity is presented next. Finally, the threshold is estimated as the intensity midway between the brightest stimulus not seen and the dimmest stimulus seen.

As a summary measure we used the mean deviation (MD), the average difference of all 54 threshold estimates from their age-corrected expected values. During the test ~10 false-negative and ~10 false-positive catch trials were presented to estimate the observer’s reliability. GATE tests involved ~200 stimulus presentations and took ~6 minutes.

Analyses
The relation between central and peripheral visual field damage was examined via Spearman rank order correlation between MD (central field) and MIR (peripheral field).

Retest variability was estimated with a modified version of Bland-Altman analysis,44 which relates the differences between repeated tests to the best available estimate of the underlying “true” value (the mean of the repeated tests). The median of the retest differences indicates systematic changes between the first and second test that can arise from learning effects, and the retest variability is estimated from the dispersion of the differences. Because the standard deviation of the differences is highly affected by outliers, we used the MAD of the retest differences to estimate the limits of agreement. We defined these as the median difference ± 2.2 * MAD, which estimates the range in which 9 out of 10 observations would be expected to fall if the data were normally distributed.

Graphical representations of the visual fields and statistical analyses were performed in R statistical software (version 2.15.1; https://cran.r-project.org/bin/windows/base/old/2.15.1/, provided in the public domain by the R Development Core Team).

RESULTS
Most patients in this sample had moderate to moderately advanced damage in the central visual field, and only one patient had an MD better than −3.0 dB (Table 2).

Relationship Between Peripheral and Central Visual Fields
Our results demonstrated the large dispersion between peripheral and central visual fields (Fig. 2). For example, some patients with deep central losses showed a nearly normal peripheral isopter (see patient e in case examples, Fig. 6),

![Graphical representation](image_url)

**Figure 2.** Relationship between global summary measures of peripheral visual field (MIR) and central visual field damage (MD). Each data point shows the mean of the two repeated tests. The Spearman rank order correlation coefficient was 0.51 (95% CI: 0.18, 0.74).

<table>
<thead>
<tr>
<th>TABLE 2. Summary Statistics of the Central and Peripheral Visual Field Tests</th>
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<tr>
<td><strong>Central visual field (GATE)</strong></td>
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<tr>
<td>Mean deviation (dB)</td>
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<tr>
<td>False-negative response error rate (dB)</td>
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<tr>
<td>False-positive response error rate (dB)</td>
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<tr>
<td>Test duration (min:s)</td>
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<tr>
<td><strong>Peripheral visual field (AKP)</strong></td>
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<tr>
<td>Mean isopter radius (deg)</td>
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<td>Isopter confidence band (deg)</td>
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<td>Test duration (min:s)</td>
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IQR indicates interquartile range.
while others with similar or less-marked central damage showed a much more constricted peripheral isopter (see patient z; Fig. 7). In particular, in patients with severe central damage, the position of the peripheral isopters varied substantially (see patients B and f in the Supplementary Material).

The Spearman rank order correlation coefficient of MIR and MD was $\rho = 0.51$ (95% confidence interval [CI]: 0.18, 0.74; Fig. 2). This correlation is considerably lower than the correlations between the test and retest values of MD ($\rho = 0.89$, 95% CI: 0.78, 0.95) and MIR ($\rho = 0.92$, 95% CI: 0.84, 0.96). This means that the lack of a close relationship between central and peripheral visual field estimates in our data is a true finding and not caused by poor precision of individual examinations.

**Test-Retest Variability of Static and Kinetic Perimetry**

There were no meaningful systematic differences between the results of the two tests, in either the central or the peripheral visual field (median test-retest difference, 0.25 degrees and $-0.1$ dB, $\rho = 0.28$ and 0.78, respectively). The median absolute differences between test and retest were 1.3 degrees with MIR and 0.9 dB with MD, and approximately 90% of test-retest differences were within ±4 degrees (MIR) with kinetic perimetry and within ±2.5 dB (MD) with static perimetry.

A formal comparison of retest variability between static automated perimetry (SAP) and AKP is problematic—after all, different regions of the visual field are measured with different estimation techniques and with different units of measurement. We therefore related the spread of the retest differences to the range of measures obtained in this sample (Fig. 2 for peripheral kinetic visual field tests, Fig. 3 for central static visual field tests). The ratio between the ranges of the data (width of the gray rectangle) to the spread of test-retest differences (height of the rectangle) was similar for central and peripheral examinations. Thus, the precision of AKP of the peripheral field is similar compared to that of SAP of the central visual field.

Most patients completed the session without any problems, but in two patients the initial kinetic tests had to be interrupted to instruct the patients to avoid false-positive...
Obviously erratic “outlier” responses occurred in about two thirds of tests (see single responses in case examples, Figs. 5–7, and Supplementary Material). This confirmed the need to obtain several responses to achieve a precise estimate of isopter position. The width of the confidence interval around the isopters, derived from the MAD of repeated responses, varied between patients by a factor of $>5$ (Table 2).

In 19 patients (65%) our technique resulted in gaps in the isopter (see patients u and z in Figs. 5, 7), because some stimuli could not be detected until they were close to fixation. In one patient with deep and widespread visual field damage, no useful isopter could be estimated with the III.1.e stimulus because more than 75% of responses were located within the central 10 degrees (patient B in Supplementary Material).

### Case Examples

Examples of three individual patients illustrate the relationship between peripheral and central visual fields and the repeatability of the tests (Figs. 5–7). Both central and peripheral visual field examinations are shown by overlaying the grayscale representation of the central visual field with a plot of the kinetic isopter. Single kinetic responses are shown as red dots, and the final isopter is plotted in dark green. Median responses <10 degrees were treated as “missing data” and appear as gaps in the isopter. The MAD measuring the scatter of single responses is shown as a green band surrounding the isopter, and normative values are represented as the light green band.

### DISCUSSION

The objective of this study was to explore differences between central and peripheral visual field damage in glaucoma and to investigate the precision of isopters that are estimated from repeated kinetic stimulus presentations. Our results show that patients with similar central visual field loss may have strikingly different peripheral visual fields, and this suggests that peripheral perimetry may provide an important component of a more complete assessment of patients’ visual field–related functional impairment. Furthermore, our results demonstrate that kinetic perimetry of a single isopter can provide a global estimate of peripheral visual field with precision similar to that of the MD of static perimetry in the central visual field. In contrast to other approaches to automate kinetic perimetry, the simple approach reported here does not aim to reproduce the often complex isopter shapes of manual Goldmann perimetry in damaged visual fields. Rather, it aims to provide a clinically useful summary measure of peripheral visual field extent that can be used to complement information available from static perimetry of the central visual field.
Lynn et al. have previously described “spurious spikes” in isopters from automated kinetic perimetry. As in Lynn’s data, our results revealed obvious “outlier” responses in most of the automated kinetic exams. One approach to reducing the impact of such outliers is to increase the number of obtained responses. Nowomiejska et al., for example, measured along 24 instead of the traditionally recommended 12 meridians. In contrast, we increased the sampling by repeating presentations at the same meridians. By pooling this information on the reproducibility of responses at each position of the visual field, this approach allowed us to estimate a confidence interval for the isopter for each individual patient. The ±4-degree retest interval of the III.1.e isopter compares favorably to data reported previously.

With manual Goldmann perimetry, the peripheral borders of the visual field are traditionally determined with the I.4.e stimulus in healthy visual fields or with the III.4.e or V.4.e stimuli when visual fields have already sustained some damage. In this study, we used the III.1.e stimulus (approximately equivalent to the I.3.e isopter, which is the largest Goldmann isopter not constrained by facial features, in healthy eyes), to keep stimulus size similar to that most often used in static perimetry (0.48 degrees). Given that our technique will almost always be applied to patients with moderate and advanced visual field damage, more intense (larger and/or brighter) stimuli must be considered for future work. However, this does not change our principal conclusion that, with a fully automated kinetic technique, isopters are best derived from repeated rather than single stimulus presentations.

The kinetic approach used in this study was designed for currently available commercial equipment (Octopus 900, with tests fully prespecified in an EyeSuite XML file). The long test times (~11 minutes, on average) would make its clinical application challenging. With the Open Perimetry Interface (OPI), it will now be possible to reduce test time through more efficient sampling strategies. For example, stimulus presentations should start closer to the expected isopter locations, and when two closely spaced responses have already been obtained on a particular vector, a third presentation may not be needed. It may also be useful to confine kinetic perimetry to those parts of the peripheral visual field that are likely of greatest importance to real-world performance (e.g., inferior and temporal visual field) rather than over the entire 360-degree circumference of the visual field. Finally, application of the OPI will make it possible to adapt stimulus speed more interactively to the response latencies of the patient and to the location of the stimulus (faster in the periphery and slower in the center of the visual field).

Our study demonstrated that precise estimates of peripheral isopters can be obtained from a fully automated kinetic approach when repeated presentations are offered. Further work is now being performed in our laboratory and others to improve the efficiency of this approach, to investigate how it can best be used to complement information obtained with static perimetry, and to answer the question of how perimetry of the entire visual field can help to improve clinical decision making in patients with glaucoma.

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