High spatial resolution automated perimetry in glaucoma

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Abstract

Background—Automated perimetry is of fundamental importance in assessing visual function in glaucoma. A technique was evaluated to perform high spatial resolution automated perimetry to allow a more detailed assessment of the luminance sensitivity in selected regions of the visual field than is possible with conventional perimetry.

Method—High spatial resolution perimetry was performed using a Humphrey automated perimeter by measuring luminance sensitivity across a 9 by 9 degree custom grid of 100 test locations with a separation between adjacent locations of 1 degree. Quantitative analysis of the raw and Gaussian filtered thresholds was performed to assess the repeatability of the technique in normals, glaucoma suspects, and glaucoma patients.

Results—The testing protocol was well tolerated by all subjects. High spatial resolution perimetry in glaucomatous eyes demonstrated fine luminance sensitivity loss not suspected with conventional perimetry. High spatial resolution perimetry also demonstrated reproducible areas of sensitivity loss in some glaucomatous eyes in areas of the visual field which appear normal with conventional programmes. The repeatability of the technique correlated with mean threshold sensitivity and was substantially improved to clinically acceptable levels by Gaussian filtering the thresholds.

Conclusion—This technique of high spatial resolution perimetry allows the practical assessment of selected regions of the visual field at higher resolution than conventional perimetry, and may be clinically useful in glaucoma.

Conventional automated perimeter threshold examinations have a relatively low spatial resolution with 6 degrees separating adjacent test locations in the Humphrey 24-2 or 30-2 programs. Perimetry performed at higher resolution may obtain a more detailed picture of the luminance sensitivity in selected areas of the visual field which may be of particular interest to the clinician. We have performed automated threshold perimetry at higher spatial resolution in selected areas of the visual field by using test locations separated by a 1 degree field. This has allowed us to investigate scotomas in much finer detail than is possible with conventional perimetry. Previous investigators have performed perimetry at higher spatial resolution than conventional perimetry, and have identified subtle scotomas beyond the resolution of conventional perimetry.1 2 High spatial resolution perimetry has also been shown to identify scotomas that correspond to retinal nerve fibre layer defects in glaucoma suspects who have normal conventional perimetry.1 3 This suggests that high spatial resolution perimetry may have a number of distinct clinical advantages over conventional perimetry in the detection of the earliest glaucomatous field defects. However, previous techniques of high spatial resolution perimetry have been arduous for patients and have been limited by long testing times.1 4

Another recognised problem of high spatial resolution perimetry has been a high intratest variability, with a corresponding poor repeatability of the threshold measurements.1 We have developed a technique to perform high spatial resolution perimetry using a Humphrey perimeter to generate a fine matrix map (FMM) of a specified region of the visual field within an acceptable test time. This technique of fine matrix mapping has been applied to the study of a number of diseases.5–8

The aim of this study was to investigate the use of fine matrix mapping in glaucoma. We then investigated the use of image processing techniques to improve the repeatability of the high spatial resolution luminance measurements, as these techniques have already been shown to substantially improve the repeatability of conventional perimetry, with no additional cost in test time.1 4

Patients and methods

Subject selection

The study was approved by the Moorfields Hospital ethics committee. We prospectively recruited patients with an established diagnosis of primary open angle glaucoma, Glaucoma suspects, and suitably age matched normal controls. All subjects were phakic and had uncorrected visual acuity of >6/9 in at least one eye. All glaucoma patients had documented intraocular pressure >21 mm Hg on at least one occasion before treatment, and evidence of asymmetrical glaucomatous field loss relative to the horizontal meridian on the most recent Humphrey 24-2 or 30-2 field. Glaucoma suspects had documented intraocular pressure >21 mm Hg on at least one occasion, and a
normal visual field. A field was defined as normal if there were no clusters of significantly depressed locations in either hemifield on the pattern deviation plot of the 24-2 or 30-2 programs of the Humphrey field analyser. According to previously established criteria, a cluster was defined as being significantly depressed if it contained a minimum of three adjacent depressed test locations on the pattern standard deviation plot, with at least one location having a probability of abnormality of $p < 0.01$ and two locations of $p < 0.05$, excluding the peripheral ring of the HFA 30-2. Suitably age matched controls were recruited if they had no significant ocular history, had a normal ocular examination with an IOP less than 21 mm Hg and had normal Humphrey HFA 24-2 fields, according to the above criteria.

TECHNIQUE OF FINE MATRIX MAPPING

To perform fine matrix mapping perimetry the coordinates of four interlaced $5 \times 5$ grids of 25 locations (with a separation between adjacent points of 2 degrees) are entered in the ‘custom grid’ feature of a Humphrey automated perimeter. Each grid is offset relative to the other grids by 1 degree—in the $x$, $y$, or $x$ and $y$ axes. The patient undergoes examination using each of the four grids sequentially, using a target size III on a standard Humphrey bowl illumination of 31.5 apostils. Accuracy of fixation is monitored in the same way as conventional perimetry.

The data are converted into IBM format files which are merged using custom software to produce a single matrix with a separation between test locations of 1 degree. This FMM of 100 locations subtends a visual angle of 9 degrees by 9 degrees, approximately the area occupied by four locations in the 30-2 program. The numerical matrix is then used to generate a surface or contour plot showing the size and location of luminance sensitivity gradients across the grid. In addition to analysis of the raw data, the FMM thresholds underwent spatial processing using a 3 by 3 Gaussian (normal) filter. This technique has been used to filter conventional Humphrey 30-2 field data and FMMs, and has been described in detail previously.

EXAMINATION PROTOCOLS

High spatial resolution perimetry of the blind spot

We investigated the resolution of our technique by obtaining FMMs of the blind spot of a normal control. Luminance contour plots were superposed to the corresponding fundal landmarks, imaged with a confocal laser scanning ophthalmoscope (CLSO), using argon 488 nm light (Zeiss instruments).

High spatial resolution perimetry in patients with retinal nerve fibre layer defects

Two glaucoma patients with clinical evidence of focal retinal nerve fibre layer defects which did not correspond to a scotoma on conventional perimetry underwent high spatial resolution perimetry. Contour plots of the luminance sensitivity were superposed to CLSO images of the retinal nerve fibre layer defects.

Clinical evaluation of high spatial resolution perimetry

Further evaluation of high spatial resolution perimetry was performed in glaucomatous eyes, glaucoma suspect eyes, and normal controls. Each glaucoma patient underwent Humphrey field examination using the 30-2 program. One eye only was chosen for examination. If both eyes satisfied the above inclusion requirements, the eye showing the greatest contrast in defect depth/sizes across the horizontal midline was chosen. The two sites of fine matrix mapping were chosen as mirror image pairs across the horizontal meridian, and were performed at two eccentricities: the supero and inferotemporal site (Fig 3A), or the superior paracentral (Fig 5A) and inferior paracentral sites.

Each subject underwent examination first at one site with each of the four constituent 25 location grids performed successively with a short, standardised rest between each. The patient was then tested at the mirror image retinal location using the same protocol of four successive grids of 25 locations. The patient was then allowed a rest of at least 2 hours and then the whole protocol was repeated at the original retinal locations to assess intertest fluctuation and reproducibility. The time required for each constituent grid varied

![Figure 1](http://bjo.bmj.com/brjophthalcom/first-published-as/104136/bjo.81.6.452.on.1.June.1997.Downloaded.from.http://bjo.bmj.com/)

Figure 1 Contour plots of luminance sensitivity obtained from fine matrix maps (FMMs) performed over a blind spot of a normal control, superposed with a fundal scanning laser ophthalmoscope image (inverted) of the subject. Contour lines represent isoluminant points, in 1 dB increments. Note steep sensitivity gradients at the edge of the blind spot as well as more subtle linear defects corresponding in location and extent to the major retinal vascular trunks exiting the optic disc.
between each patient depending on the number of questions asked, related to defect size and depth. Excluding the brief rest between grids, the test time required on the Humphrey field analyser to obtain a FMM in the glaucoma patients was clinically acceptable and ranged from 8 to 26 minutes, with a mean of 20 minutes.

Glaucoma suspects were tested according to the same protocol using the mirror image pair of FMMs in the supero and inferotemporal site. In the absence of conventional field abnormalities, we chose this site as it has been reported to be one of the locations of early visual field defects in glaucoma suspects. We investigated the repeatability of the FMMs in age matched normal controls, using the same test sites, except that the testing was confined to either the superior or the inferior location of the mirror image pair.

Three dimensional surface plots of the FMMs for matched locations in the visual field were compared between groups. Quantitative analysis of the raw and Gaussian filtered thresholds was performed, and the repeatability of the FMMs was assessed using the technique described by Bland and Altman (see below).

STATISTICAL ANALYSIS
The repeatability of the FMMs was investigated using the technique described by Bland and Altman. This technique has been applied to investigate the repeatability of a number of measurements of ocular components, and has also been used to assess the repeatability of visual field measurements. To assess the repeatability of the first and second FMMs, the numeric difference between the first and second threshold sensitivities is calculated for each test location and plotted versus the mean of the sensitivities for that location. This graphical plot illustrates the spread and distribution of the pointwise differences. The repeatability of the FMMs is defined as the SD of the pointwise differences between the first and second FMM.

Figure 2 (A) Humphrey 24-2 grey scale and STATPAC 2 total deviation plot from glaucoma suspect right eye. Box indicates the site of two adjacent fine matrix maps (FMMs), which reveal an obvious superior arcuate extending from the blind spot, not shown by the Humphrey 24-2. Contour plots (B) have been superposed with a fundal scanning laser ophthalmoscope image (inverted) obtained from the patient. The sensitivity loss corresponds closely with the extent of a retinal nerve fibre layer defect.
Results

HIGH SPATIAL RESOLUTION PERIMETRY OF THE
BLIND SPOT

Figure 1 shows the results of performing high spatial resolution perimetry around the blind spot of a normal subject. A contour map of the luminance sensitivity profile has been superposed to a fundal CFLO image (using argon 488 nm light, Zeiss instruments), obtained from the same eye. Accurate superposition was achieved by aligning anatomical landmarks such as the centre of the disc and fovea with the corresponding perimetric landmarks such as centre of the blind spot and fixation, which are plotted by the Humphrey field analyser. The resulting aligned contour map reveals well defined steep luminance sensitivity gradients at the edge of the blind spot as well as more subtle linear relative defects corresponding in location and extent to the major retinal vascular trunks exiting the optic disc. The diameters of the retinal arterioles at the disc margin are approximately 100 µm, while those of the retinal veins are about 130 µm; 100 µm corresponds to a visual angle of approximately 0.3 degrees. This degree of resolution compares well with other reported techniques of high spatial resolution perimetry, which have also identified angioscotoma associated with the major retinal vascular trunks.16

HIGH SPATIAL RESOLUTION PERIMETRY IN PATIENTS WITH RETINAL NERVE FIBRE LAYER DEFECTS

Figure 2 illustrates the results of high spatial resolution perimetry performed in a glaucoma patient in an area of the visual field corresponding to a clinically visible focal retinal nerve fibre layer defect. A three dimensional surface plot of the luminance sensitivity (Fig 2A) reveals an obvious scotoma which is not apparent on the conventional Humphrey 24-2 test locations.
improvement in repeatability. Lines indicate the least factor of approximately 2, representing a twofold filtering has been to reduce the magnitude of the SD by a factor of approximately 2, representing an improved repeatability by a factor of 2 between the baseline and the second FMMs. The mean pointwise difference remains the same for the raw and Gaussian filtered data, and is close to zero in all the plots.

Figures 3B–E are examples of the Bland–Altman plots of the first and second FMMs performed for one glaucoma patient in two mirror image locations (Fig 3A) of the field. The superotemporal FMM overlaps an area of depressed threshold sensitivity on the Humphrey 30-2, while the inferotemporal FMM overlapped an area of normal Humphrey 30-2 threshold sensitivity.

The repeatability between the first and second FMMs is poorest, as indicated by the higher pointwise SD, in the location of depressed threshold sensitivity (Fig 3B) compared with the location of normal threshold sensitivity (Fig 3C). The effect of Gaussian filtering of the FMMs has been to substantially reduce the SDs (Fig 3D, E) representing an improved repeatability by a factor of 2 between the baseline and the second FMMs. The mean pointwise difference remains the same for the raw and Gaussian filtered data, and is close to zero in all the plots.

Fig 4A shows the repeatability of the first and second FMMs for all the subjects: the SD of the pointwise differences has been plotted against the mean threshold sensitivity of the first and second FMMs for each subject. The SD values are highest for the glaucoma patients, indicating poorer repeatability, compared with the controls. The one outlier was a glaucoma patient who had FMMs performed in a region of absolute sensitivity loss on the Humphrey 30-2. In this patient only one test location had a measurable sensitivity, the remaining test locations had no measurable sensitivity in either baseline or the repeat FMM, with a consequent zero pointwise difference between the majority of points. Figure 4A illustrates the negative correlation between the SD and the mean grid sensitivity which was highly significant (p = 0.0001, R² = 0.91); thus, the repeatability was poorest in the FMMs with greater degrees of luminaire sensitivity loss. Figure 4B shows a similar correlation between the pointwise SD and the mean threshold sensitivity for the Gaussian filtered thresholds. Note that the values of the SD have been reduced by a factor of approximately 2. The limits of agreement for all the Gaussian filtered FMM pairs are below 5 dB; thus, for each FMM pair, the difference between the first and second (repeat) sensitivity thresholds will be less than 5 dB at 95 out of 100 test locations, which we believe is clinically acceptable.

Calculated values for the means of the pointwise differences between the first and second FMMs, representing any overall bias between the first and the second grids were smallest for controls (mean −0.09 dB, range −0.58 dB to 0.50 dB). The means of the pointwise
differences for the glaucoma patients were higher (mean 0.37 dB, range −6.70 dB to 3.00 dB), although they still remained small; 11 of 12 repeated FMMs had a mean difference of less than 4 dB. In addition, there was no systematic departure from zero in any of the groups, indicating that there is no significant bias between the first and second FMMs which we might have expected if there was a significant learning or fatigue effect.

FMMs obtained from high spatial resolution perimetry in glaucoma patients at sites already known to be abnormal from the 30-2 examination revealed a complex profile of sensitivity loss which was not revealed by the low spatial resolution of the conventional Humphrey 30-2 or 24-2 programs. Figures 5A and B show Humphrey 30-2 grey scale and STATPAC 2 total deviation plots from a glaucoma patient who underwent high spatial resolution perimetry at the site indicated. Figures 5C and D show three dimensional surface plots of the first FMM (Fig 5C) and the second (Fig 5D) FMMs which have undergone filtering. Both surface plots show the complex profile of the scotoma, which reveals a steep sensitivity gradient from severely depressed to normal sensitivity. Much of the detail is reproducible.

In two glaucomatous eyes of two patients fine matrix mapping in an area of normal sensitivity on the Humphrey 30-2 revealed an obvious reproducible defect that was not apparent with conventional perimetry. One example is shown by Figures 6A–E. High spatial resolution perimetry identified a localised reproducible scotoma, represented as an elevated area on the first FMM (Fig 6C) and second FMM (Fig 6D). The FMMs were performed in an area of field overlapped by four Humphrey 24-2 test locations which have normal threshold sensitivity (Fig 6A, B). Figure 6E shows a surface plot of a FMM from a normal control for comparison, which shows a uniform luminance profile.

Discussion

A number of investigators have used high spatial resolution perimetry to detect scotomas beyond the resolution of conventional perimetry. Airaksinen et al have used high spatial resolution perimetry to identify scotomas corresponding to retinal nerve fibre layer defects in glaucoma suspects who have normal conventional perimetry. In addition, high spatial resolution perimetry has also been shown to be clinically useful in defining residual small central isles of field in advanced glaucoma, which may be too small for conventional perimetry to map. However, a major disadvantage of some of these studies is that they have been arduous for patients and have required an extremely long test time. We have performed high resolution perimetry in a group of normal, glaucoma, and glaucoma suspect eyes using the technique of fine matrix mapping, performed on a Humphrey automated perimeter. We have identified glaucomatous luminance loss which was not revealed by the lower spatial resolution of the conventional Humphrey 30-2 or 24-2 programs. The testing
A protocol was well tolerated and all subjects performed the test within a test time of 30 minutes, which we believe is clinically acceptable. Previous investigations of high spatial resolution perimetry have demonstrated that one of the earliest perimetric disturbances in glaucoma is an elevated intratest variability, in some regions, even though the mean threshold sensitivity at the location may still remain normal. A number of studies have shown that the intertest variability of conventional threshold perimetry is higher in glaucoma patients than in normals. In addition, the degree of intertest variability has been found to correlate with the degree of sensitivity loss. Our results confirm that a similar correlation exists between intertest variability and sensitivity loss in high spatial resolution perimetry, as has been suggested by other workers. To date, the high intertest variability, and consequently low repeatability, in areas of glaucomatous sensitivity loss has been a significant limitation to the usefulness of high spatial resolution perimetry. Attempts to improve the precision of the threshold measurements by repeating the measurements is unlikely to be practical in high spatial resolution perimetry because of the already lengthy test times. We have investigated an alternative method to improve the repeatability using image processing techniques, which have already been shown to improve the repeatability of conventional automated perimetry. We have been able to obtain a marked improvement (by a factor of 2) in the repeatability of the technique, by applying a Gaussian filter to the high spatial resolution perimetry threshold data. This improvement in repeatability has been achieved at no extra cost in test time. We have obtained good reproducibility of the luminance profiles using this technique, and this has allowed us to obtain finer details of sensitivity loss which were not apparent on the lower spatial resolution of conventional perimetry. We believe that high spatial resolution perimetry fine matrix mapping may be a useful clinical tool, and may have a diagnostic role in the more detailed evaluation of equivocal areas of visual field of glaucoma patients and suspects. Previous studies have suggested that the test point density of conventional perimetry is not a limiting factor for the identification of the earliest glaucomatous field change. However, these studies used conversion criteria based on the development of field defects measured at conventional resolution. It remains to be determined whether the detection of conversion is
improved by the use of high spatial resolution perimetry. In addition, high spatial resolution perimetry may have a role in identifying early progression of glaucomatous visual field defects. There is evidence to suggest that the progression of glaucomatous defects, as detected using conventional perimetry, is slow.\(^{24,25}\) Using conventional perimetric programs with a spacing of 6 degrees between adjacent test points, progression of the advancing edge of a scotoma which is increasing in size must exceed 6 degrees of visual angle before a change is registered at a repeat examination. Perimetry performed using test locations separated by 1 degree offers the theoretical possibility that we may be able to identify advancement at the edge of a scotoma at an earlier stage than is possible with conventional perimetry.

In summary, we have described a technique for performing high spatial resolution perimetry which is clinically practical and can be performed on an unmodified Humphrey automated perimeter. We have identified glaucomatous luminance loss which was not revealed by the lower spatial resolution of the conventional Humphrey 30-2 or 24-2 programs, and have obtained a substantial improvement in the repeatability of the technique by using image processing techniques.

This work was supported by grants from the Friends of Moorfields, the Medical Research Council, the International Glaucoma Association, and the Royal National Institute for the Blind.