Choosing two points to add to the 24-2 pattern to better describe macular visual field damage due to glaucoma

Siyuan Chen
Department of Computing and Information Systems, the University of Melbourne, Australia

Allison M. McKendrick
Department of Optometry and Vision Sciences, the University of Melbourne, Australia

Andrew Turpin
Postal:
Department of Computing and Information Systems
The University of Melbourne
Parkville 2010 VIC
Australia
Email: aturpin@unimelb.edu.au
Telephone: +61 3 83441312

Keywords: Visual fields, visual field tests, glaucoma, macular region

Word count: 1956 (excluding title page, abstract, references, figures and tables.)
ABSTRACT

**Background/aims**— A recent study has shown that the paracentral upper visual field in the macular region is often affected in glaucoma and suggested that two test locations within the central 10 degrees should be added to the Humphrey 24-2 visual field test pattern to detect such damage. This study employed data collected using a different visual field test pattern to determine whether the same two test locations are supported as the most informative regarding visual field loss.

**Methods**— A dataset of 62 glaucomatous patients and 48 controls had visual field assessments on the Medmont perimeter M700 (Central Threshold or Glaucoma test). Twelve 24-2 locations within central 10 degrees of visual field were derived by interpolation of the nearest neighbours of the Medmont data. The remaining 24 Medmont locations in the central 10 degrees of the glaucomatous set were labelled as abnormal if their thresholds fell outside the lower 5th percentile of the age-corrected values for same location from the control group. All possible pairs of the 24 locations were then assessed for diagnostic power by counting the number of patients that had 0, 1 or 2 abnormal locations in a pair.

**Results**— Overwhelmingly pairs of locations in the superior macular region were more often abnormal than pairs in the inferior. About 50 pairs of locations had equivalent ability to detect damage, with the best pair having 74% of patients with at least one of the locations as abnormal, and 52% both.

**Conclusions**— Adding a pair of locations to the superior macular region of the Humphrey Visual Field 24-2 pattern increases the number of abnormal locations identified in individuals with glaucoma.
Tomograph II (Heidelberg Engineering, Germany), and all were normal on the Moorfields Regression Analysis or Glaucoma Probability Score Tool. The study was approved by the human research ethics committee of the University of Melbourne and all participants provided written informed consent, in accordance with a protocol consistent with the Declaration of Helsinki.

For the purpose of this study, only one eye per subject was used. If data from both eyes was available, only the right eye was used. If only the left eye data was available, its test locations were mapped to a right-eye format for analysis. Therefore, 62 eyes from the glaucoma group and 48 eyes from the control group were selected from their most recent VF tests. All thresholds were corrected to age 50 using an age-correction factor of -1 dB/decade. The median Average Defect and Pattern Defect scores in the glaucomatous group were -1.80 (range: -7.45 to 2.51) and 9.43 (range: 0 to 20.61) respectively. The Average Defect and Pattern Defect scores on the Medmont are similar in principle to the Mean Deviation and Pattern Standard Deviation of the HFA, but have some differences in the calculation[8].

**Visual Field tests and locations**

Visual field data were collected using the Medmont perimeter M700 (Central Threshold or Glaucoma test), which is a fully automated, hemispheric bowl perimeter (300-mm radius) [9, 10]. In this VF test, stimuli, Goldmann size III (0.43°), are produced by LEDs of 565-nm wavelength that retroilluminate fixed points within the bowl with a background illuminance of 10 apostilbs (3.2 cd/m²)[9]. A ZEST (zippy-estimation by sequential thresholding) procedure is used to determine thresholds. All participants were tested by a trained perimetrists who ensured that fixation was maintained during the test and provided rest breaks during testing as required. Ninety-five percent of participants had fixation losses less than 30%. The other 5% were observed carefully by the perimetrists and showed good fixation via direct observation. The rate of false-positive responses was also estimated in catch-trials, which was less than 21%.

Medmont VF measurements were recorded with test locations distributed on seven circles at eccentricities of 3°, 6°, 10°, 15°, 22°, 30° from the foveal centre. Because only the macula is of interest in this study, measurements from the test locations at eccentricities of 3°, 6°, 10° were analysed. Figure 1 shows the test locations as grey circles. Also shown as black squares are the two test locations, (±1°, 5°), recommended to be added by Hood et al.[6]. Henceforth we refer to these two locations as the Hood Pair.

**Analysis**

As we were interested in adding two locations to the 24-2 pattern, we excluded from our analysis those locations in the Medmont field that would be represented in a 24-2 examination. However, as none of the 24-2 points fell directly on the Medmont locations (one of the attractions of using this dataset for this study), we derived which locations to exclude using the nearest neighbours of the Medmont pattern, which are shown using dashed lines in Figure 1. We then examined all possible pairs of Medmont locations that had not been used to derive the 24-2 pattern (grey circles in Figure 2). There were 276 possible pairs (C_2^1) of locations, and for each pair in the glaucoma data set we counted the number of points below the 5th percentile of the controls for those locations (left panel in Figure 2). Thus each pair can contribute 0, 1 or 2 abnormal points to the 24-2 pattern. We counted the number of patients that have either exactly two abnormal points, or more than zero abnormal points.

To place our results in the context of a 24-2 examination, we also derived threshold values for the 24-2 locations by averaging (after anti-logging) the nearest neighbours, and counted abnormal locations in the 24-2 pattern in the same way as for the Medmont locations (Figure 2).
RESULTS

Figure 2 left hand panel shows the dB values that correspond to the 5th percentile of the control data. Figure 2 right hand panel shows the number of patients that were abnormal in each location. As can be seen, the highest numbers are in the superior field. Figure 3 shows the proportion of patients with abnormal points in each pair of locations. The top panels only show the 44 best performing pairs, while the histograms give data for all 276 pairs. The error bars in the top panels indicate the upper end of a 95% confidence interval derived using bootstrapping. Thus, for all the pairs in the top left panel, the upper end of their 95% range of the number of patients with 1 or 2 abnormal locations includes the mean for the best pair (indicated with the dashed line). For the top right panel, only the first 23 pairs have 95% ranges that include the best mean.

DISCUSSION

Our analysis confirms that the Hood Pair is a reasonable choice of two points to add to the macular region of the 24-2 pattern to improve the detection of macular loss. This confirmation provides confidence to the original findings of the Hood study, as an entirely different dataset was used, collected with a different perimeter, different test pattern, and different test algorithms than the Hood et al. study [6]. Of the 11 patients that showed no macular damage on the twelve 24-2 locations within the macula, 5 had abnormalities at one or both of the Hood Pair (Figure 4, white bars). Of the 4 patients who had one abnormal location on the 24-2 macula test locations, one had a further abnormality in the Hood Pair. While there are other pairs in the superior macular region that perform equally well as the Hood Pair (Figure 3), there is no clear reason to prefer these other pairs.

Other schemes exist for choosing extra locations to add to the 24-2 pattern [11, 12]. Aoyama et al suggested that if the gradient between four test points is large, an additional test location should be placed at the centre of the four points [11]. This is because the VF sensitivity in areas of high gradient is difficult to predict by simple interpolation. By applying this gradient method, we expect the gradient of the four 24-2 locations in the superior macular to be larger than the gradient between the four inferior locations. If this is the case, then the added location would be at (0°, 6°), between and slightly superior to the Hood Pair, but not that far away. Indeed, in our dataset, we found that the average gradient of these four points in the superior field across the control group and across the glaucomatous group are 0.5±0.22 and 0.91±0.68 dB/deg interval, respectively. Not surprisingly, they are larger than the average gradients in the lower VF, 0.43±0.22 and 0.80±0.73 dB/deg interval, respectively.

Another method to determine the distribution of test locations is to maximise an assumed structure-function relationship. For each 12 clock-hour sector of the optic disc, Asaoka et al. [12] chose four test points so that the correlation between VF thresholds and retinal nerve fibre layer thickness was maximised. Interestingly, two of the selected test points in their scheme are close to the Hood Pair, namely, (3°, 5°) and (-1°, 4°), and the Hood Pair is far from those test points that have the weakest structure-function relationship in the macula. This provides further evidence that the Hood Pair may be a good choice for adding to commonly used test patterns.

A further issue to consider that falls outside the scope of the current study is to determine the optimal number of test points to be added to 24-2 tests to maximise benefits. Here we specifically chose two test points to be added in order to compare with existing studies, with the specific aim of determining whether there was spatial concordance with the two test locations previously predicted as most useful. The key aim was to verify these locations using an independent dataset, collected using a different visual field algorithm and pattern. Future work may address the question of the relationship of the
number of additional test points and the cost benefit trade-off between time taken for testing and information gain.

In summary, in this study, we confirmed that test locations around the Hood pair, (±1°, 5°), can improve the detectability of glaucomatous functional damage in the macula over a basic 24-2 pattern.

ACKNOWLEDGEMENTS
This work was supported by Australian Research Council (LP130100055).

Competing interests
AT and AMM receive research support from Heidelberg Engineering GmBH in association with Australian Research Council Linkage Project LP130100055. SC: None

Patient consent
Obtained.

Ethics approval
This study was conducted with the approval of the University of Melbourne Human Research Ethics Committee, Melbourne, Australia.

Contributors
All authors listed have been involved in the conception and design, acquisition of data or analysis and interpretation of data, drafting the article or revising it critically for important intellectual content, and final approval of the version published.

REFERENCES
1. Hood DC, Raza AS, de Moraes CGV, Liebmann JM, Ritch R. Glaucomatous damage of the macula. Progress in Retinal and Eye Research 2013;32(0):1-21
INTRODUCTION

There is considerable evidence for macular involvement in glaucoma[1, and references therein]. Clinically, glaucomatous damage in the macular is observed using ocular coherence tomography, and results in spatially localised[2, 3] as well as diffuse[4, 5] visual field loss. However, the most common visual field test used clinically for glaucoma (Humphrey Visual Field Analyzer 24-2 pattern, Carl Zeiss Meditec, Dublin CA, USA) has only four test points within the central 8 degrees. An alternate current strategy is to use the 10-2 test pattern, which tests on a 2 degree grid, however this requires an additional test to be performed by the patient. Typically 10-2 visual fields are not conducted routinely on people at risk of glaucoma, but are utilised when glaucoma is seen to be threatening fixation. Recent work by Hood and colleagues demonstrates that macular scotomata can exist in the absence of peripheral visual field loss in glaucoma and can be missed entirely by the 24-2 visual field pattern[1, 6].

Performing a detailed test of both central and mid-peripheral vision might be advantageous for detecting and monitoring field loss (for example, both the 10-2 and 24-2); however, it increases the test duration and has logistical limitations. Hence, it has been suggested that a practical solution is to add several additional test locations to the 24-2 pattern in the macular region. Hood and colleagues[6] determined that if only two points were to be added, the most useful locations would be at (±1°, 5°). These locations were determined from 10-2 data from 31 people with glaucoma.

In this paper we take data collected on the Medmont M700 perimeter (Medmont International Pty Ltd, Nunawading) which collects thresholds in arcuate rings and samples more densely than the 24-2 test pattern in the macular area (see Figure 1). The thresholding algorithms between the Humphrey Field Analyzer (HFA) and the Medmont differ, including spatial post-processing of the threshold data for the Swedish Interactive Thresholding algorithm (SITA)[7] and the spatial order in which the points are tested. Consequently, algorithmically derived spatial dependencies within the measured visual fields may vary between these two measurement techniques, and these might influence the apparent best test points for detecting damage. Our aim was to determine which two locations could be added to the central 10 degrees of the 24-2 pattern to improve the detection of macular damage due to glaucoma and to confirm whether the locations were the same as previously derived using visual fields measured with the HFA.

METHODS

Participants
Our dataset includes 110 participants who underwent between one and three Medmont visual field (VF) tests during a three year period while enrolled in other studies in our laboratory. The dataset includes 62 people with glaucoma (median age: 72.1; range: 52.8 to 87.1) and 48 visually normal controls (median age: 65.5; range: 48.8 to 84.8). Individuals with glaucoma had an ophthalmological diagnosis of primary-open-angle glaucoma based on clinical findings, visual fields (typically 24-2 HFA fields) and optic disc appearance. All had prior experience in taking visual field tests before visiting our laboratory. All were currently being treated, had refractive errors of no greater than +/- 6 diopters spherical and no more than 2D of cylinder, and had visual acuity of better than 6/9. Control participants had the same refractive error and visual acuity criterion, had normal findings on a comprehensive ocular examination including slit-lamp biomicroscopy, optic nerve head evaluation and applanation tonometry. Optic disc evaluation was also conducted using the Heidelberg Retinal


Figure 1 Grey circles show the Medmont M700 test locations, numbered for later reference. Black circles show the 24-2 test locations, with dashed lines showing which Medmont locations were interpolated to get the threshold values for these locations. The two black squares indicate the test locations recommended by Hood et al.[6].

Figure 2. Grey dots show the candidates for adding to the 24-2 pattern (black circles). The numbers in the left panel give the lower 5th percentile of the control data for each location in dB. The numbers in the right panel give the number of patients that had that location as abnormal.

Figure 3. For the best 44 pairs of locations, the proportion of patients that had at least one abnormal point in the pair (A) and both locations abnormal in the pair (B) are shown. Bars show the mean, and error bars 1.96 times the standard deviation of 100 bootstrap samples for each pair. The dark grey bar in each plot represents the pair (24, 26), the light grey (11, 24) and the black the Hood Pair (11, 12). The frequency distribution of the proportion of patients with 1 or 2 abnormal in the pair (C) and exactly 2 abnormal in the pair (D) is also shown. The shaded areas of the histogram indicate the number of pairs whose mean plus 1.96 times standard deviation does not drop below the best performing mean.

Figure 4. Each group of three bars represents a number of abnormal points in the set of 12 macular 24-2 locations. Within each group, the number of patients that had 0, 1 or 2 of the Hood Pair as abnormal is shown.
## Proportion of patients with at least one abnormal location in a pair

<table>
<thead>
<tr>
<th>Prop. of patients with abcount 1 or 2</th>
<th>0.0</th>
<th>0.2</th>
<th>0.4</th>
<th>0.6</th>
<th>0.8</th>
<th>1.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of pairs</td>
<td>11</td>
<td>12</td>
<td>24</td>
<td>21</td>
<td>26</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>24</td>
<td>33</td>
<td>33</td>
<td>33</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
</tr>
<tr>
<td></td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
</tr>
</tbody>
</table>

## Proportion of pairs detecting at least one abnormal point

<table>
<thead>
<tr>
<th>Prop. of patients with abcount 1 or 2</th>
<th>0.0</th>
<th>0.2</th>
<th>0.4</th>
<th>0.6</th>
<th>0.8</th>
<th>1.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of pairs</td>
<td>20</td>
<td>15</td>
<td>10</td>
<td>5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
</tbody>
</table>

## Proportion of patients with two abnormal locations in a pair

<table>
<thead>
<tr>
<th>Prop. of patients with abcount = 2</th>
<th>0.0</th>
<th>0.2</th>
<th>0.4</th>
<th>0.6</th>
<th>0.8</th>
<th>1.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of pairs</td>
<td>11</td>
<td>11</td>
<td>12</td>
<td>12</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>23</td>
<td>24</td>
<td>26</td>
<td>26</td>
<td>24</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td>11</td>
<td>11</td>
<td>11</td>
<td>11</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>26</td>
<td>26</td>
<td>26</td>
<td>26</td>
<td>26</td>
<td>26</td>
</tr>
</tbody>
</table>

## Proportion of pairs detecting two abnormal points

<table>
<thead>
<tr>
<th>Prop. of patients with abcount = 2</th>
<th>0.0</th>
<th>0.2</th>
<th>0.4</th>
<th>0.6</th>
<th>0.8</th>
<th>1.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of pairs</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>