Review of the Use of SWAP and FDT for the Early Detection of Visual Field Loss

John McBride¹ and Fiona J Rowe²*

¹Directorate of Orthoptics and Vision Science, University of Liverpool, Liverpool L69 3GB, UK.
²Department of Health Services Research, University of Liverpool, Liverpool L69 3GB, UK.

Authors’ contributions

Author JMB designed the study, performed the review and wrote the drafts of the manuscript. Author FJR provided oversight for the review and contributed to the drafts of the manuscript.

ABSTRACT

Background: To review the literature reporting SWAP and FDT for the early detection of visual field loss in glaucoma.

Methods: A review of literature published on Medline, Scopus and Web of Science between 1966 and present was undertaken with only articles in the English language reviewed.

Results: SWAP has high inter and intra subject variability. There is no uniform decision as to what criteria should be applied to define abnormal visual fields. In comparison with standard automated perimetry (SAP) there is controversy as to whether SWAP is reliable in detecting glaucomatous damage. As a screening tool FDT may have potential due to its short testing time and lower test-retest variability. There is a lack of definitive abnormality criteria to define visual field loss with FDT. First and second generation FDT have been found to be comparable to SWAP in terms of diagnostic sensitivity. There is a lack of studies comparing FDT to SAP with longitudinal follow up of visual fields which make it difficult to determine its reliability in identifying pre-perimetric glaucomatous damage.

Conclusion: There is a need for further longitudinal studies on both SWAP and FDT to determine fully their reliability in detecting early visual field loss in glaucoma before SAP.

*Corresponding author: E-mail: rowel@liverpool.ac.uk;
Keywords: SWAP; FDT; Matrix; SAP; Glaucoma; Visual Field Evaluation.

1. INTRODUCTION

Primary open angle glaucoma (POAG) is a chronic, potentially blinding condition caused by death of ganglion cells and their axons within the optic nerve due to increased pressure within the eye [1]. It affects approximately 3% of people over 65 in the UK and with average life expectancy in the UK rising, more people are now living with glaucoma even compared to the mid-90’s [2,3]. The cost of treating glaucoma in the UK is more than £300 million per year. This includes both direct (medication, outpatient care, routine assessment and surgical procedures) and indirect treatment (cost of low vision services, loss of earnings resulting from vision loss) [4,5]. Glaucoma also affects quality of life with partial sight or registered severely sight impaired being ultimate possibilities. One of the determinants for a poor visual prognosis is signs of glaucomatous field damage upon presentation to the clinician. This means vision and visual field could be potentially saved if damage could be reliably detected before current visual field analysis is able to [6,7]. Glaucoma diagnosis and progression are determined by three factors; intraocular pressure (IOP) changes, optic disc examination and visual field evaluation. There are a number of methods which can be used to evaluate the visual field including standard automated threshold perimetry, frequency doubling technology, critical flicker frequency, short wavelength automated perimetry, high pass resolution perimetry and motion automated perimetry amongst others.

Within the visual pathway there are a number of sub-pathways that subserve different aspects of vision and whose physiological properties are distinct. The M (magnocellular) pathway consists of larger retinal ganglion cells that are mainly linked with visual perception and respond predominantly to motion and coarse outlines. They are regarded as insensitive to colour stimuli in balanced luminance conditions, respond to high contrast sensitivity and resolve higher temporal and lower spatial frequencies [8,9]. The K (konioacellular) pathway consists of smaller, sparse retinal ganglion cells that are sensitive to short wavelengths (S/blue cones) [8-10]. The spatial and temporal resolutions of the K pathway overlap with the M and P pathways [11]. The P (parvocellular) pathway consists of small retinal ganglion cells that are sensitive to colour (particularly red/green), respond predominantly to fine details and to lower contrast sensitivity. They resolve lower temporal and higher spatial frequencies [8,9]. It is possible to isolate responses from the M, P and K pathways by using different testing stimuli. Strategies to isolate the P and K pathways include the use of coloured, high contrast, small stimuli whereas strategies to isolate the M pathway include the use of stimuli that are colour neutral, have high temporal and low spatial frequencies such as large, fast motion reversal, low contrast targets.

It is known that in early glaucoma, larger ganglion cells within the optic nerve are damaged. First the blue-yellow ganglion cells (part of the konioacellular pathway) which are larger than red-green ganglion cells are damaged leading to dyschromatopsia [12]. This effect has been exploited in testing for early glaucoma damage by using short wavelength automated perimetry (SWAP). SWAP is used with blue points (wavelength 440nm) with a narrow band presented for 200msec on a yellow background (100cd/m²) to try and detect early colour vision loss [3]. Studies have shown that SWAP perimetry can detect glaucomatous damage 3-5 years before conventional white on white perimetry [13-15]. With current standard white on white perimetry around 30-50% of ganglion cells have been destroyed before detection [16].
Furthermore, the M pathway can be isolated when assessing for early glaucoma. Frequency doubling technology (FDT) was first described by Kelly over 45 years ago. It is based on the fact that when an achromatic sinusoidal grating of low spatial frequency undergoes flickering at high temporal frequency the apparent spatial frequency of the grating appears to double [17]. The response to this is thought to be mediated by M_y-cells, which are sensitive to low contrast and motion stimulus. M_y cells are again larger than other retinal ganglion cells (part of the magnocellular pathway) and are therefore more prone to damage in early glaucoma [18]. This forms the rationale for using FDT as a potential visual field screening method for early glaucoma detection.

There is an alternative theory as to why SWAP and FDT may be able to detect early glaucomatous damage: as a result of the reduction in the redundancy in the visual system [19]. The function-specific tests isolate a subset of ganglion cells which may lead to an increased sensitivity to early ganglion cell loss.

There has been conflicting evidence regarding the use of SWAP and FDT for early detection of visual field loss. The aim of this review is to evaluate the literature in respect to whether FDT and SWAP perimetry can reliably detect glaucomatous visual field changes before Humphrey SAP.

2. METHODOLOGY

The SCOPUS, Web of Science, PubMed and Medline databases were searched for the following terms: SWAP and glaucoma, Short Wavelength Automated Perimetry and glaucoma, SWAP and ocular hypertension and Short Wavelength Automated Perimetry and ocular hypertension, SWAP and open angle glaucoma, SWAP and closed angle glaucoma, Frequency Doubling Technology and glaucoma, FDT and ocular hypertension, Frequency Doubling Technology and open angle glaucoma, Frequency Doubling Technology and angle closure glaucoma, Frequency Doubling Technology and Short Wavelength Automated Perimetry, Frequency Doubling Technology and screening, Matrix FDT and ocular hypertension, Matrix FDT and glaucoma and Matrix FDT and closed angle glaucoma.

We included international articles but only those written in English were reviewed due to lack of access to translation facilities. The years included were from the year 1966 to 2013. In this review only studies reporting the use of SWAP and/or FDT for early visual field loss in glaucoma detection are considered.

We found 2917 articles from our search, accessed 112 full articles which were assessed as relevant to the review topic and evaluated 79 articles for this review article.

3. RESULTS AND DISCUSSION

3.1 Short Wavelength Automated Perimetry (SWAP)

SWAP is a type of visual field assessment which is based on the principle that larger ganglion cells within the retina are selectively damaged during early glaucoma. Ten per cent of these larger ganglion cells belong to the blue-yellow pathway: part of the koniocellular pathway [12]. The earlier death of this ganglion cell population has also been confirmed by histological evidence [20-22]. The blue-yellow conditions of SWAP isolate the S cone system (short wavelength cones), reduce the participation of other cone systems (red – long
wavelength and green – medium wavelength) and saturate the rods activity through the adaptation to yellow light [23,24]. This phenomenon has been reported to be of use to predict glaucomatous visual field damage up to 4 years before it is detected on standard achromatic perimetry [13,14].

SWAP can be performed on the Humphrey Visual Field Analyser (HVF: Carl Zeiss Meditec Inc., Dublin, CA, USA) and the Octopus perimeter (Haag Streit Int., USA). A Goldman size V target (1.74° diameter) is used with a blue filter placed over the source of light allowing only light with a peak transmission of 440nm through. The white background light is replaced by yellow light illuminated at 100 cd/m² in the perimeter bowl. The targets are presented for 200msec in all test locations. The test is performed within the central 30 degrees of the visual field.

SWAP has not been introduced routinely into clinical practice due to a number of factors: clinicians are not united on a definition of abnormality criteria [25], the effect of the ageing lens on results [26], lengthy test time [24], patient fatigue and patient’s dislike of the test [27] and the large intra and inter-subject variability even with normal subjects [28] which make it difficult to define an abnormal field. Some of the factors causing a barrier to its use in practice will be addressed in more detail: i.e. test time by means of test choice and abnormality criteria.

3.1.1 Test (Algorithm) choice

The lengthy test time for SWAP is an important factor and, as stated above, is one of the reasons why SWAP is not routinely used in clinical practice. One way in which test time for SWAP could be reduced is to consider the choice of algorithm. There are a number of test programmes (algorithms) that can be used around which there is debate as to which should be used. Current versions available for SWAP include a full-threshold option whereby a staircase method is used to plot sensitivity at specific test locations [29]. A SITA-FAST (Swedish Interactive Threshold Algorithm) version of the SWAP algorithm was developed which reduced testing time by up to 70% (11 minutes with full threshold and just over three minutes with SITA-SWAP) with no apparent reduction in sensitivity of specificity [30]. It is also reported that the SITA-SWAP algorithm partially reduces the issue of the large inter-subject variability by approximately 14% [29-31].

3.1.2 Abnormality criteria

The statistical properties of SWAP have been examined extensively, firstly by Wild et al [26] who tried to establish a normal database containing probability levels and analysis synonymous to that for SAP. An increased between level variability was found between subjects compared to SAP with the variability increasing with target eccentricity and increasing age. This is relevant as the risk of developing glaucoma increases with age[32]. The between subject variation is also important to note when looking at probability maps to define an abnormal field. To use the same level of depression required for SAP would be unwise as this will lead to a large number of false positive results. This adds evidence for the need to design bespoke statistical analyses rather than a cross-over with SAP.

This view was also supported by Kwon et al. [33] showing a similar amount of variation upon repeated testing. Both studies had a similar number of participants making their results comparable, however, neither provided an indication of what may be an appropriate criterion for what ‘normal’ may be.
Polo and associates [25,34,35] aimed to suggest and refine abnormality criteria for SWAP by addressing the sensitivity and specificity at different probability levels. They suggested that using cluster point criteria (four points clustered at the P<5% level or three points clustered lower than P<1%), rather than using global indices, provide the best sensitivity and specificity. A criticism of the classification of the criteria they used to define glaucomatous damage was inclusion of changes in the retinal nerve fibre layer as opposed to the gold standard optic nerve head assessment (optic nerve images taken by photography). NICE guidelines suggest optic nerve changes are more indicative of glaucomatous damage than retinal nerve fibre layer assessment as the latter damage is not always present yet the patient can have other characteristics of glaucoma including raised IOP, optic disc cupping and visual field loss [36].

This view is also supported by Johnson et al [37] who suggested four abnormal points at the P<5% level are indicative of glaucomatous visual field loss. Johnson and associates [37] used their own controls in their study to define a normative database rather than the parameters provided in the perimeter. However, they did test the validity of their controls before comparing these with their ‘abnormal’ group and their normative dataset is large and almost double that of Polo and colleagues [25,34,35] Although the normative database is large the visual fields were only performed once on the control subjects who had no prior knowledge of the test whereas the abnormal group had at least four years’ experience on both SAP and SWAP. It has been shown that there is a significant learning effect for both SWAP and SAP so it is possible the control subjects performed worse on the perimetry tests which made them more comparable to the abnormal patients showing signs of reduced sensitivity [38-40].

The work by Polo [25,34,35] and Johnson [37] is valuable and upon repeated testing Reus et al. [41] found that Polo’s [25] criteria for abnormality on SWAP seemed to obtain more positive results compared with Johnson’s criteria [37]. There was still a large variation in the amount of positives obtained depending on which criteria were used. Reus et al. [41] allowed for the learning effect which appears to improve specificity [42]. No indication was provided regarding the end point of the study and subjects were not followed up to determine if SAP defects persisted or developed.

There is little research to date that has specifically addressed the definition of an abnormality criteria set for SWAP. Many studies use their own normative data and their own diagnostic criteria as to what constitutes a defect. This introduces uncertainty as to whether SWAP can be a clinically useful tool. Further work is required to create a “gold standard” diagnostic criterion for SWAP in order to make direct comparisons between studies. A summary of the studies attempting to define a abnormality criteria set is shown in Table 1.

### 3.1.3 SWAP and SAP

The initial SWAP studies followed ocular hypertensive and early glaucoma patients for five years to determine whether SWAP was able to predict those patients who would go on to develop glaucomatous visual field damage. Johnson et al. [13,14] and Sample et al. [43] showed promise in the use of blue-on-yellow perimetry to detect visual field loss before standard white on white perimetry but with a number of limitations. Explicit criteria for a deficit on white on white perimetry was used (<5% probability level for mean deviation (MD) or pattern standard deviation (PSD) or more than three points clustered at the <5% level).
Table 1. Summary of studies attempting to define abnormality criteria for SWAP

<table>
<thead>
<tr>
<th>Study</th>
<th>Sample Size</th>
<th>Control Group Used</th>
<th>Suggested Abnormality Criteria</th>
<th>Follow up Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kwon et al. [33]</td>
<td>31 eyes</td>
<td>33 eyes</td>
<td>None suggested</td>
<td>8 weeks</td>
</tr>
<tr>
<td>Polo et al. [25]</td>
<td>95 eyes</td>
<td>128 eyes</td>
<td>Four clustered points at P&lt;5% level or three clustered points at P&lt;1% level</td>
<td>3 years</td>
</tr>
<tr>
<td>Johnson et al. [37]</td>
<td>479 eyes</td>
<td>348 eyes</td>
<td>Four clustered points at P&lt;5% level</td>
<td>4 years</td>
</tr>
<tr>
<td>Reus et al. [41]</td>
<td>744 eyes</td>
<td>None used</td>
<td>Four clustered points at P&lt;5% level</td>
<td>6 months up to 1 year</td>
</tr>
<tr>
<td>Van der Shoot et al. [45]</td>
<td>416 eyes</td>
<td>None used</td>
<td>None suggested and compared directly with SAP criteria of at least 3 points at P&lt;5% with at least two at P&lt;1% level or corrected PSD &lt;5%</td>
<td>7 to 10 years</td>
</tr>
<tr>
<td>Liu et al. [46]</td>
<td>132 eyes</td>
<td>37 eyes</td>
<td>None defined</td>
<td>Not reported</td>
</tr>
</tbody>
</table>

No such criterion was implemented for blue-on-yellow as to what actually constituted a defect. It was evident there was a lot of variance even between tests with defects presenting and disappearing upon repeated testing. Their initial test conditions did factor in absorption of yellow light which is important as the ocular media can act as a neutral density filter particularly in the presence of cataract [44]. Their sample size for which they base their results on is relatively small for the amount of people potentially available with ocular hypertension. One issue with the results obtained in the early studies was the reporting of defects in terms of total deviations rather than pattern deviation which is important. Total deviation provides an indication of the depression of the hill of vision whereas pattern deviation gives an indication of the abnormality of the shape of the field. They did however age match their patients but this was only against a maximum of 22 subjects. Considering the variability that SWAP produces this may not be an entirely appropriate approximation of glaucomatous field loss. The results from the papers published by Johnson and colleagues [14,15] and Sample and colleagues [43] are what have driven subsequent studies.

Interestingly the subjects recruited by Polo and colleagues [34] to establish their abnormality criteria were recalled three years later [35] to determine how many had developed a defect on SAP. There was a high percentage who were deemed to have a SWAP defect and did not go onto develop glaucomatous damage after three years indicating the specificity of the test was not high and in the region of 30%.

There have been contradictions to the supporting evidence for SWAP. Van der Shoot et al [45] state SWAP is less sensitive than SAP to predict conversion to glaucoma. However, in their study, no correction criteria were provided for cataract which is known to decrease the mean sensitivity of SWAP by absorption of the short wavelength light in the lens. This is important considering that the age range used in their study would include those starting to develop cataract [26]. The conversion to glaucoma, which was the end point of the study, was based purely on visual field analysis. No anatomical correlation (e.g. optic nerve head cupping) was sought that matched with the visual field defect.

Liu et al. [46] also supports that SWAP is not as sensitive as SAP for detecting glaucoma. They used a sub analysis of early glaucoma as defined by the CIRRUS OCT which had
been modified to ‘capture’ early glaucoma at the expense of specificity. With this method they may however have included patients that otherwise would not have been considered with solely visual field assessment.

3.2 Frequency Doubling Technology (FDT)

The frequency doubling phenomenon was first described by Kelly over 40 years ago[17]. When a low spatial frequency sinusoidal grating (<1 cycle/°) undergoes high temporal counter phase flickering at 15Hz or greater, gratings appear to double to twice their actual spatial frequency[47]. Essentially there is a rapid contrast reversal which light bars become dark and vice versa.

The frequency doubling phenomenon is thought to be mediated by a subset (five per cent) of ganglion cells within the magnocellular pathway called M_y cells [18,48]. Similar to the blue-yellow pathway cells the M_y cells have larger diameter axons making them more prone to damage in early glaucoma [21,49]. In order for FDT to be effective the spatial frequency should be between 0.1 and 4 c/deg^1 (cycles per degree) and the temporal frequency greater than 15Hz. Both are dependent upon retinal eccentricity in addition to a mean background luminance of 50cd/m^2[18,24].This forms the rationale for the use of FDT to detect glaucomatous field damage earlier than SAP.

3.2.1 Test (Algorithm) choice

FDT first came into use in 1997[50]. The first generation FDT manufactured by Welch Allyn (Skaneateles Falls, New York, USA) used a 10° square stimuli, which was found to be initially problematic as its large size prevented it from detecting small visual field defects [24,51,52]. Second generation FDT was released in 2005 using a 5° square stimuli to overcome this problem using the Humphrey Matrix visual field instrument [53,54]. The Matrix test pattern was designed to be similar to the Humphrey Field analyser and Octopus perimetry using a 0.43° round target on a 6° square matrix [50].

Screening programmes on the first generation FDT included the central 30 degrees of the visual field. These algorithms used a modified binary search (MOBS) strategy. The algorithm presents a grating at a level where 99% of normal subjects of an aged matched control would see. If this is seen the zone is labelled normal, if this is not seen on a repeated presentation then a grating at the 99.5% level is presented and increased until the stimulus is seen [24,55]. The screening tests are performed in a supra-threshold pattern. Supra-threshold is performed whereby the intensity of the stimulus shown is calculated to be above the patient’s threshold. Threshold is performed where the stimulus is decreased in intensity using an adaptive staircase procedure until no response is given [56].

Subsequent test programmes have been developed to reduce the test time for FDT. They include rapid efficient binary search (REBS) and zippy estimation of sequential testing (ZEST)[55]. REBS is based on the MOBS strategy but requires two response reversals whereas MOBS requires four [55]. ZEST uses a probability density function at each location (the probability that each subject will have that threshold) with patient responses modifying the probability density function until the standard deviation of the probability density function becomes sufficiently narrowed [57]. The results of FDT tests have been compared to a normative database of 700 eyes and can be age matched [58].
Isawe and colleagues [59] proposed that FDT could be an effective method of visual field screening due to the short test time, its portability and relatively low-cost. Studies have reported sensitivity for FDT ranging between 83% to 93% and specificities ranging between 55% to 100%[60-65]. FDT also appears to have lower test-retest variability and less of a learning effect which makes it a promising emerging psychophysical test [66].

### 3.2.2 Abnormality criteria

FDT is similar to SWAP in that definitive criterion for what is defined as normal and abnormal has not been uniformly decided upon. As Matrix FDT was released in 2005 there remains a lack of large population longitudinal studies to assess what criteria would be most discriminatory for a glaucoma diagnosis. Different criteria have been suggested although the view for abnormal criteria appears to be reaching a more uniform decision compared with SWAP.

Fogagnolo et al. [67] propose that the presence of at least one point at the P<5% level on the pattern deviation plot (PDP) on the N-30 or C-20 screening programme is the most suitable criteria for detecting glaucoma. Six different classifications of a defect were used and analysed by receiver operator characteristic curves (ROC). A ROC curve plots true-positive rate against the false positive rate [68]. The area under a ROC curve of 1 gives 100% sensitivity and specificity whereas a ROC of 0.5 would indicate no diagnostic value [47]. The more stringent of the criteria did not significantly have a higher sensitivity or specificity for detecting glaucoma which explains the author’s choice of defect criteria in order to capture most glaucomatous loss. Patients included in the study had POAG to enable the authors to identify the most sensitive and specific criteria for diagnosing glaucoma. Definitions for glaucoma defects on SAP were in accordance with the guidelines recommended by Hodapp-Parish-Anderson [70] which are also currently used by the National Collaborating Centre for Acute Care [36] in the United Kingdom. The results for sensitivity and specificity for visual field loss was comparable to those of others [63, 71-73] and particularly in moderate and advanced glaucoma cases. Notably the authors found that the screening programmes C-20 and N-30 showed a lower sensitivity for detecting nasal step defects; one of the most common glaucomatous defects [70].

Brusini et al. [74] suggest PSD <5% and/or at least two areas on the PDP P<5% should be used as the cut off criteria for the N-30 programme using the first generation FDT. Their results with these diagnostic criteria were similar to those previously reported by Weinreb, Medeiros & Sample [75] for the same programme.

A small cross sectional comparative study by Hong et al. [76] also suggest using more than five points in the PDP at the P<5% level to discriminate between glaucomatous and non-glaucomatous fields. This classification is more stringent than those proposed by Fogagnolo et al. and Brusini et al. [67,74]. Hong and associates allowed for the influence of cataract on visual field testing [77] which had not been accounted for in previous studies. All subjects included in the study had manifest POAG detectable on SAP which were termed as early defects i.e. MD not worse than -6dB. Both the normal and POAG group had a mean age of under 40 which is significant as it is more likely that older individuals than in this age group would be routinely screened for glaucoma [78]. ROC curves were used to analyse the criteria proposed by the authors (MD P<1% and PSD P<5%) and appropriate statistical testing was used. There was however a significant difference in the depth of defect between the control group and the study group suggesting the criteria may be useful but only in the age range of the population studied. Further work is required using these criteria for
predicting visual field loss in at risk groups and for the learning effect to be taken account for when considering a cut-off point.

Choi, Lee and Park [66] examined the use of the Humphrey Matrix perimeter for the detection of pre-perimetric glaucoma and to define a clinically useful cut-off point. A strength of the study is that the learning effect was allowed for both SAP and FDT by giving each patient two practice attempts. Their initial ophthalmic investigation also included all gold standard tests as defined by the National Collaborating Centre for Acute Care [36]. ROC curves were used to determine which parameter (MD or PSD) provided the best discriminating power. PSD was found to be the best discriminator which is in agreement with Hong et al. [76].

Abnormality criteria are essential for defining a glaucomatous visual field when screening. As is the current situation with SWAP, definitive criteria have not been agreed upon. However evidence currently published does not show as much variation as what ‘abnormal’ should be defined as on FDT when compared to defining SWAP abnormality criteria. There is a need for more longitudinal studies taking into account for the variances that occur in the screening population, the age of patients performing the test and where the criteria produce repeatable defects. A summary of the studies attempting to define abnormality criteria is shown in Table 2 for direct comparison.

**Table 2. Summary of studies attempting to define abnormality criteria for FDT**

<table>
<thead>
<tr>
<th>Study</th>
<th>First Generation or Matrix FDT</th>
<th>Sample Size</th>
<th>Control Group Used</th>
<th>Suggested Abnormality Criteria</th>
<th>Follow up Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fogagnolo et al. [67]</td>
<td>First generation</td>
<td>40 eyes</td>
<td>40 eyes</td>
<td>At least one point at P&lt;5% on the PDP</td>
<td>Not reported</td>
</tr>
<tr>
<td>Brusini et al. [55]</td>
<td>First generation</td>
<td>258 eyes</td>
<td>60 eyes</td>
<td>PSD &lt;5% and/or at least two areas on PDP at P&lt;5%</td>
<td>Not reported</td>
</tr>
<tr>
<td>Hong et al. [76]</td>
<td>Matrix</td>
<td>36 eyes</td>
<td>24 eyes</td>
<td>More than 5 points on PDP at P&lt;5%</td>
<td>Not reported</td>
</tr>
<tr>
<td>Choi, Lee and Park [66]</td>
<td>Matrix</td>
<td>99 eyes</td>
<td>122 eyes</td>
<td>PSD of greater than -3.14dB</td>
<td>Not reported</td>
</tr>
</tbody>
</table>

### 3.2.3 FDT and SAP

Few studies exist that longitudinally compare FDT with SAP in respect to whether a subject who presents with a defect on FDT would proceed to develop a defect on SAP. Many of the early studies looking at agreement between FDT and SAP were conducted with subjects who already had detectable manifest glaucoma [79,80].

A small prospective study by Quaranta et al. [81] examined the concordance of first generation FDT with SAP in 25 manifest glaucoma patients and 25 ocular hypertensives using MD plots; a measure of overall sensitivity loss. Although the study indicated a significant correlation between MD in the glaucoma and ocular hypertensive group, they also compare PSD but with no results presented other than ‘it was significant between groups’.
There was no classification given as to what defines a visual field as glaucomatous or not which makes it difficult to draw conclusions from. No follow up was performed in this study to ascertain whether the defects persisted on repeated testing or whether the subjects showing FDT defects in the ocular hypertensive group went on to produce defects on SAP.

In a larger study by Racette et al. [82] on 80 eyes, they found that first generation FDT was better able to detect early glaucomatous damage than SAP using ROC analysis. Mean deviation for SAP and PSD <5% for first generation FDT were found to be the best indicators of early glaucomatous damage. However, there are issues with their definition of their normal pressure. Routinely a pressure under 21mmHg is considered normal yet a cut-off upper pressure of 23mmHg was used. The study has an advantage over the smaller study by Quaranta and colleagues as they provide parameters to classify glaucomatous loss and check for persistence of defects. Furthermore, they acknowledge the limitations of their study and for longitudinal change, further studies are required.

Medeiros and associates [83] reported that first generation FDT defects were predictive of future SAP defects in a longitudinal observational study over 17 months. The subjects were retrospectively taken from a larger study (Diagnostic Innovations in Glaucoma Study) but were not taken at random from the study group: the purpose being to include only subjects at risk of developing POAG and not from secondary causes e.g. pigment dispersion syndrome. This potentially introduces selection bias. It is interesting to note that the age range of the sample taken is not given yet the authors found a statistically significant difference between the age of subjects who converted and those who did not. The lack of age range renders it difficult to consider the results for the population at risk. However, gold standard tests were used to exclude manifest glaucoma as per National Collaborating Centre for Acute Care guidelines [36] and allowances were made for learning effect [84]. More importantly the fields were repeated at follow up to determine if the defects persisted. To reduce the influence of intra test variability and for a defect to be confirmed on SAP the field was repeated three times. A normal database was used from a previous study conducted by the authors in normal individuals and found 94% specificity upon using their definition of an abnormal FDT (two points on MD P<5% level or a PSD of P<0.05%). Fourteen out of the 105 eyes studied progressed to repeatable defects on both SAP and FDT with the locations of the defects correlating. The results of this study show promise in terms of a possible prediction of the development of SAP defects by location of repeatable defects on FDT.

Spry, Hussin and Sparrow [80] compared the use of matrix FDT 24-2 threshold strategy with the 24-2 SITA-Fast programme available on the HVF. The study was larger than that of Quaranta et al. [81] and all patients went through gold standard assessment for detection of glaucoma (SAP, gonioscopy, tonometry and optic nerve head examination) [36]. A strength of the study was the classification of subjects into subgroups based on clinical examination separating out the different types of glaucoma and suspects. In contrast to the Quaranta study [81] the mean age of the study group was more representative of the patients undergoing glaucoma testing. Upon comparison of MD and PSD moderate agreement was found between the two techniques by means of a ROC analysis. The criteria used to define a glaucomatous field were liberal and based upon clinical decision in order for it to be applied to the clinical environment and the authors acknowledge that this may have led to the high sensitivity found. Spry and associates [80] suggest that for the purposes of glaucoma screening sensitivity should be compromised in order to optimise the specificity and positive predictive value of FDT. In order to confirm the results further follow up is
required to consider whether individuals who are screened with matrix FDT with the 24-2 test pattern do develop defects that are synonymous with SAP.

### 3.3 FDT and SWAP

Both SWAP and FDT have been recognised as potential methods for the detection of early glaucoma. Studies have looked at the comparability of FDT with SWAP with respect to sensitivity and specificity and concordance with each other [27,46,85-89].

Landers and associates [85] reported that FDT and SWAP are comparable in terms of sensitivity, specificity and concordance. Spry et al. [80] examined the diagnostic capability of both SWAP and FDT. They reported that FDT had a lower test-retest variability (39% SWAP compared to 34% FDT) but both methods had similar abilities in detecting early glaucoma. One issue with the comparison is that FDT was not performed using the Welch Allyn nor matrix FDT perimeter but on a 'custom device'. Thus it is difficult to compare their results with other studies using standard rather than customised assessments. Specifically, the 24-2 test pattern used for their FDT was a 4° diameter square which is not used in either the Welch Allyn or matrix perimeter [50,51]. Gold standard tests were used however to classify glaucoma and pattern deviation was used to classify defects rather than total deviation; the latter providing a generalised impression of field loss. The use of PD or PSD is also supported by Leeprechanon et al. [27] who found that, for matrix FDT, this provided the best sensitivity and specificity. As for SWAP there is not a definitive definition of ‘abnormal’ as previously discussed.

A prospective study by Horn et al. [87] attempted to overcome the limitations of previous studies in order to determine whether SWAP and first generation FDT are able to detect preperimetric glaucoma, with interesting results. A double-bagging technique was used in the study. The previous studies by Landers et al, Soliman et al. and Spry et al. [47,80,85] relied upon making a clinical judgement regarding the outcome of the visual field testing. By using the double bagging technique this standardises the outcome for all participants with all fields being treated in the same way. Horn et al. [87] recognised that the numbers assessed in their study might not be sufficient to produce the most accurate, reliable or efficient algorithm and therefore requires a larger study. The method itself may facilitate future study and allow this area of research to finally move forward.

Matrix FDT was compared with SWAP in a prospective case-control study by Leeprechanon et al. [7]. The results showed a significant correlation between FDT and SWAP using MD and PSD with FDT showing a higher sensitivity of detecting glaucoma (72% versus 54%). The sample size was small, however the 24-2 test pattern used for both FDT and SWAP allowed for a more direct comparison of results. A strength of the study was that concordance of the location of the defect was investigated with a 72% agreement reported. When considering the indices PSD and PD at the P<5% level they were significantly higher in the early glaucoma group versus the normal group on FDT but not on SWAP suggesting that FDT may be better at filtering out the generalised noise which is known to occur in SWAP.

More recently Liu et al. [46] found matrix FDT to be more sensitive than SWAP in detecting glaucoma at comparable levels of specificity. The criteria used for defining an abnormal visual field for FDT and SWAP was the same as that for SAP for which there is contention as to whether SAP criteria can be used for other visual field assessment methods [26,66]. Furthermore, the confidence intervals overlapped for matrix FDT and SWAP results which may indicate that there is little difference in the diagnostic abilities of SWAP and matrix FDT.
A subgroup analysis was performed with patients with ‘early glaucoma’ but no definition of early glaucoma was provided. The results reported by Liu et al. [46] were in agreement with those of Leeprechanon et al. [27] in their larger sample size.

4. CONCLUSION

With an aging population and the rise in the prevalence of glaucoma worldwide there is a clear need for rapid diagnosis to commence treatment early to preserve both vision and visual field. The need for rapid diagnosis however does need to be considered from an economic perspective i.e. is the time and money spent on diagnosing glaucoma earlier clinically useful providing the best cost: benefit ratio.

There is evidence available to show that SWAP, first generation and matrix FDT have potential as psychophysical tests for earlier glaucoma detection. SWAP has been investigated at a greater length than the more recently emerging FDT methods. Both SWAP and FDT lack defined abnormality criteria like those that exist for SAP. Until this is established it is difficult to determine whether or not the tests have a role in either glaucoma screening in the community or at the first visit to the hospital eye service. SWAP has the issue of a large inter and intra subject variability requiring up to five attempts before the patient becomes sufficiently reliable. This has time and economic implications which need to be considered. FDT shows less variability with more repeatable results and lesser learning effect. The testing time is also short with FDT which again may make it a more favourable test than SWAP.

SWAP and FDT on direct comparison have been found to have similar sensitivity and specificity in predicting glaucomatous visual field loss and show moderate agreement with SAP for detecting field loss. Few studies however consider whether the defects correlate between tests and, in SWAP and FDT, whether the defects develop in the same location as that in SAP. This is a question requiring further research given that SWAP (which uses blue stimuli to isolate the S cones and K pathway) may be more relevant to central perimetry within 10 degrees of fixation whereas FDT (which uses fast reversal, large contrast stimuli to isolate the M pathway) may be more relevant to perimetry peripheral to the central 10 degrees. There is a lack of longitudinal studies addressing persistence of defects and anatomical correlation of defects to be able to definitively ascertain the clinical application of both FDT and SWAP. Further research is thus recommended to answer these outstanding questions.

To conclude both SWAP and FDT could potentially be used as methods for earlier glaucoma detection but require well-defined, consistent abnormality criteria within specified testing algorithms. Currently, standard automated perimetry remains the preferred choice for clinical assessment until such time that evidence shows SWAP and/or FDT techniques to be superior in respect to accuracy and reliability in the detection of early visual field loss in glaucoma.

CONSENT

Not applicable.
ETHICAL APPROVAL

Not applicable.

ACKNOWLEDGEMENTS

Financial Disclosure: No funding was provided for the conduct of this review.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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Peer-review history:
The peer review history for this paper can be accessed here:
http://www.sciencedomain.org/review-history.php?id=369&id=23&aid=2804