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Diagnostic Sensitivity/Specificity of Preattentive Vision Tests in Glaucoma

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ABSTRACT

Purpose. Damage to the nerve fiber layer or visual pathway might be expected to reduce the efficiency with which the visual system performs analysis of the ever-changing field of vision. The purpose of this article is to provide a further analysis of previously reported data (Loughman J, Davison P, Flitcroft I, Br J Ophthalmol 2007;91:1493–98.) to: (i) determine the sensitivity and specificity of a test of preattentive vision for glaucoma detection and (ii) provide a cutoff performance level that would serve to distinguish glaucoma in early cases.

Methods. Three groups of observers (glaucoma, suspects, and normals) were examined, using computer-generated flicker, orientation, and vertical displacement targets to assess preattentive visual search (PAVS) efficiency. The task required rapid and accurate localization of a singularity embedded in a field of 119 homogenous distractors on either left or right hand side of a computer monitor. All subjects also completed a choice reaction time task.

Results. Receiver operating characteristic curve analysis demonstrates consistently high diagnostic sensitivity and specificity values (significantly above 90% for all tasks) using the raw PAVS data and also for a novel perceptual search index (which improves the diagnostic capacity of the test). Optimal performance cutoff values for each task were also computed.

Conclusions. A test of PAVS efficiency demonstrates high sensitivity and specificity to early glaucoma. Analysis incorporating the perceptual search index confirms the high diagnostic capacity of the test.

Key Words: preattentive visual search, preattentive vision, glaucoma, flicker, motion displacement, orientation, choice reaction time

The ability to process information across the entire visual field simultaneously has long been known1 and has been named preattentive vision; it enables the visual system to detect any stimulus (target) which differs sufficiently from all others (distractors) in the visual field; the target exhibits “pop-out” from the distractors and attention is immediately drawn to the target location. An example of preattentive (parallel) and serial search is shown in Fig. 1 below. The three different elements in the left panel pop out due to an orientation difference. The three elements in the right panel differ significantly in appearance but contain the same orientation and line ending information and are therefore not seen preattentively.

Application of the preattentive visual search (PAVS) paradigm to clinical conditions has been relatively recent and confined to investigation of conditions with secondary visual impact, including Parkinson disease,2 senile dementia, and Alzheimer disease.3 Probably the first attempt to investigate the relevance of visual search to primary visual clinical conditions was that of Flitcroft et al.,4 who devised a set of clinical tests which they found in a preliminary study to correlate with presence of glaucoma. In a recent study, we confirmed the findings of Flitcroft and co-workers in relation to glaucoma.5 The purpose of the current article is to present a further statistical analysis of that data. Receiver operating characteristic (ROC) curves were used to (1) generate sensitivity and specificity results and (2) determine the optimal differentiating cutoff normal values for both perceptual search index (PSI) and PAVS results for each target type.

The current preattentive vision test exploits the parallel processing capabilities of the visual system. A test of preattentive vision is inherently different from conventional psychophysical techniques.
METHODS

The basic test comprises three separate supra-threshold conditions—flicker detection, displacement (motion) detection, and orientation difference detection (an image of the orientation task and full methodological details have been published elsewhere\(^5\)). The subject’s task was to locate the singular target from among the 119 distractors on either side of the monitor using handheld buttons in a two-alternate forced choice paradigm.

All targets were white with mean luminance of 132 cd/m\(^2\); mean background luminance was 2 cd/m\(^2\) giving a Michelson contrast ratio of 0.97. The white targets and distractors subtended 0.92° with a 1.83° gap between stimuli. The flicker target was a white-filled square box of the above dimensions, square-wave modulated at 16 Hz, and surrounded by identical non-flickering boxes as the distractors.

The displacement target was an empty white box (white lines of width 1 mm, subtending 7 min of arc, forming a square with an unfilled black center), surrounded by identical stationary boxes as the distractors. The displacement target was displaced vertically by square-wave oscillation at 16 Hz through an angle of 14 min. The orientation target was the letter N surrounded by the letter Z as its distractor; both target and distractor limb widths also subtended 7 min arc. Monitor resolution exceeded that required to present the lines forming the open boxes and N and Z targets.

FIGURE 1.
An orientation difference facilitates the pop-out of three targets in the left panel which are readily detected without focused attention. Target differences in the right panel do not pop-out and require serial (foveal) attention to be detected.

parallel processing efficiency of the preattentive system is characteristically assessed by quantification of a subject’s capacity to detect feature singularities from a distracting background. As such the subject is presented with multiple targets (in this case 120) and tasked with detection of the single target, which differs from the others in terms of some basic feature such as flicker, motion, or orientation among others. It is reasonable to assume that PAVS requires neural mechanisms across the entire retina to be intact and conditions such as glaucoma may therefore impact on preattentive search efficiency.

RESULTS

Standard ROC curve analysis was used to (1) generate sensitivity and specificity results and (2) determine the optimal differentiating cutoff normal values for both PSI and PAVS results for each target type. The statistics software (Stats Direct) plotted sensitivity vs. 1—specificity for a series of cutoff values of both PAVS and PSI; data were inputted for normal and glaucomatous patients only. Optimal normal cutoff values were determined on the basis of maximum sensitivity \(\times\) specificity with equal importance assigned to both sensitivity and specificity. Figs. 2 to 4 confirms the high diagnostic capacity of the test for all target types. Each figure gives both PAVS and PSI data for one target type.

Table 1 provides a list of sensitivity and specificity values obtained for each task and illustrates that for all target types, PSI retains marginally increased sensitivity and specificity over PAVS across each target group indicating that it may provide a slightly better performance index, thus confirming the merit of producing such an index. The “sensitivity \(\times\) specificity” index in Table 1 shows the orientation PSI task to yield the most accurate discrimination between groups.

Table 2 gives details of the optimum cutoff values as determined by the analysis. These points are represented by the large open circle and square on the ROC curves for PAVS and PSI data, respectively.

DISCUSSION

In a clinical setting, and especially in the older population most typically affected by glaucoma, motor, and/or neural factors could potentially influence the accuracy of any interpretation of reac-
We have previously reported that this simple step highlights differences in performance efficiency not readily identifiable by analysis of the raw PAVS data. Furthermore, in the present analysis, Table 1 shows that both sensitivity and specificity are greater for all three tasks using PSI rather than raw PAVS data. With sensitivity and specificity values consistently well above 90%, the results here compare very favorably with alternative functional and structural technologies. Our use of ROC analysis assumed equal weighting of sensitivity and specificity and therefore of type 1 and type 2 errors; in a glaucoma screening environment it would be a matter of clinical judgment whether to modify the weighting.

Differentiating normals from glaucoma is an important factor in determining the clinical value of the test. Equally important, however, is the identification of those patients classified as glaucoma suspects most likely to develop glaucoma. Although longitudinal analysis is essential to determine the test capacity to successfully identify such patients, comparison of the raw suspect data with the determined optimal cutoffs above might give some indication as to those suspects most likely to progress. In total, 14 suspects exceeded the normal PSI criterion on at least one task. Four suspects met the glaucoma PSI criterion on all three tasks. Longitudinal analysis will, however, be required to determine which subjects eventually progress from suspect to glaucoma and whether PAVS is a good prognostic indicator.

It is our contention that analysis of PAVS efficiency such as that achieved in the current device warrants serious consideration as an addition to conventional perimetric methods. The test fulfills numerous important criteria in terms of essential properties of a clinically viable test for glaucoma, including resistance to blur, simplicity, high diagnostic sensitivity and specificity, as well as a
patient and practice friendly rapid testing time (screening in 1 min per eye, maximum test time 6 min per eye in advanced glaucoma).

The current test may benefit from optimization of stimulus parameters and test design features. Longitudinal analysis of glaucoma suspects, and analysis of the effects of other eye disease which may influence PAVS and complicate the clinical diagnosis, deserve further exploration.

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REFERENCES


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AQ1— Kindly check if the short title is OK as given.

AQ2— The possessive form has been deleted in all diseases as per the journal style. Kindly check.

AQ3— Kindly spell out IOP in the text.

AQ4— Kindly check whether Table 2 is OK as edited.

AQ5— Ref. 7 is not cited anywhere in the text. Kindly cite it in order or we will mark it Deleted in Proof.