Digital imaging of the optic nerve head: monoscopic and stereoscopic analysis

J E Morgan, N J L Sheen, R V North, Y Choong, E Ansari

Aims: To compare monoscopic and stereoscopic assessment of the optic disc using novel software for the digital stereoscopic analysis of optic disc stereopairs.

Methods: Software was developed for the stereoscopic display of digital optic disc images using an interlaced display method. Neuroretinal rim width was determined at 10 degree intervals around the optic disc using a custom (stereoscopic) cursor whose depth was adjusted to that of Elschnig’s rim. Measurements were taken, first viewing the disc monoscopically and at a separate sitting, stereoscopically.

Results: Measurements were made in 35 eyes from 35 patients (1260 estimates for each observer) using three observers. The mean cup to disc ratio (CDR) ranged from 0.57 to 0.66 (SD 0.13–0.14) for monoscopic viewing compared with 0.64 to 0.69 (SD 0.12–0.14) for stereoscopic viewing. Stereoscopic assessments gave higher CDRs in temporal, superior, nasal, and inferior aspects of the optic disc (p<0.001, Mann-Whitney U test). Agreement between observers in estimating CDR was high for monoscopic assessment (intraclass correlation coefficient 0.74 (CI 0.72 to 0.76) increasing to 0.80 (0.78 to 0.82) for stereoscopic assessment.

Conclusion: Digital stereoscopic optic disc assessment provides lower estimates of neuroretinal rim width and higher levels of interobserver agreement compared with monoscopic assessments.

Accurate and reproducible assessment of the optic disc has a key role in the management of the glaucoma patient. Since considerable axon loss can precede the development of clinically detectable visual field loss, the early detection of structural retinal damage should facilitate earlier disease diagnosis and improve the clinical prognosis. Similarly, close monitoring of the optic disc can detect progressive glaucomatous damage before changes appearing in the visual field.

The poor agreement, even among expert observers for subjective assessment of the optic disc, has driven the development of clinical imaging devices such as scanning laser tomography, polarimetry, and optic coherence tomography. These devices show great promise in the diagnosis of glaucoma and in the detection of progressive disease. For example, scanning laser tomography has been reported to detect early glaucomatous optic nerve damage with a sensitivity and specificity approaching 84% and 96%, respectively, and can identify eyes with progressive damage before the onset of visual field loss.

By contrast, stereoscopic analysis of the optic disc images in glaucoma has not been developed as a routine clinical technique even though it remains the “gold standard” by which newer diagnostic technologies continue to be evaluated and has provided valuable insights into the nature of glaucomatous pathophysiology. Studies comparing the efficacy of stereoscopic imaging for the diagnosis of early glaucoma have shown that in expert hands, it can detect early glaucomatous damage with a sensitivity and specificity that compares favourably with that obtained with scanning laser ophthalmoscopic devices.

Several factors have limited the clinical analysis of stereoscopic images. Firstly, the viewing of stereoscopic images can be impractical in the clinical setting where mirror based stereoscopes are required. Secondly, the analysis of the optic discs is usually subjective, and the high diagnostic precision that has been reported with this technique has come from groups with considerable expertise in the clinical assessment of the optic disc; these results may not be applicable in less specialised units. Finally, there is little evidence to show that stereoscopic imaging per se provides sufficient additional clinical information to justify its use when compared with the monoscopic evaluation of optic nerve head images. This issue was highlighted recently by a study analysing sequential stereoscopic images, which suggests that stereoscopic imaging conveys little additional benefit for the assessment of the optic disc.

This study employs software that has been developed for the digital, computer based assessment of stereoscopic optic disc images to address many of the difficulties that have hindered the clinical analysis of these images. We describe the application of this technique, in which we have developed tools for the precise delineation of the pathological features of stereoscopic optic disc images in space. The present study was designed to determine, firstly, whether digital stereoscopic analysis provided different estimate of neuroretinal rim width compared with monoscopic assessment and secondly, whether acceptable levels of interobserver agreement could be obtained for the assessment of neuroretinal rim width. These are important questions that should be addressed before a new imaging modality such as digital stereoscopy is considered for clinical use in the diagnosis of glaucomatous optic neuropathy.

METHODS

The study is based on 35 optic disc images from patients with either glaucomatous or normal discs which were selected from our database of clinical images to provide a range of cup to disc ratios (CDRs). Patients were diagnosed as glaucomatous on the basis of visual field examination (Humphrey 24–2 SITA fast) and the existence of characteristic cupping of the optic disc indicating either diffuse or focal damage to the neuroretinal rim as identified by stereoscopic biomicroscopy.
and, if required, by stereoscopic examination of the optic nerve head images (colour slides). Ocular hypertensive patients had normal optic nerve and visual field examinations with intraocular pressures (IOPs) that were consistently above 22 mm Hg. Patients were classified as normal if they had IOPs of 21 mm Hg or below with normal visual field and optic disc examination. Stereoscopic images were taken in the course of routine patient care following informed and signed consent in accordance with requirements for the use of images in clinical research at the University Hospital of Wales, Cardiff, UK. The observers, who were clinically trained and experienced in optic disc analysis were masked to the diagnosis of the optic discs, which were examined in random order. Sequential high quality stereofundus images were obtained using a Nikon F505 fundus camera using Kodachrome EPN Ektachrome 35 mm slide film (ISO 100). In every case, the pupil was dilated to more than 5 mm using an infrared emitter so that the inner and outer margins of the optic disc, which were examined in random order. Sequential high quality stereofundus images were obtained using a Nikon F505 fundus camera using Kodachrome EPN Ektachrome 35 mm slide film (ISO 100). In every case, the pupil was dilated to more than 5 mm using one drop of tropicamide 1% and phenylephrine hydrochloride 2.5%. Stereoscopic images were taken using established techniques in which the illumination beam of the camera was set at the one part of the corneal limbus before being displaced horizontally to produce a standardised offset. All patients had clear media with a visual acuity of 20/40 or better with correction.

Stereoimages were digitised using a Nikon Coolscan slide scanner to produce 24 bit true colour (1024 × 844 pixel) bit maps. The images were displayed in above and below format using the stereoscopic viewing glasses (Stereographics) to control for the effects of viewing through LC lenses. Not less than 1 week later, the images were viewed stereoscopically again with the inner and outer ends of the line coincided. The length of each line could be adjusted at each 10 degree interval so that the inner and outer ends of the line coincided with the inner and outer margins of the optic disc, respectively. Once the user had completed this adjustment, CDRs were computed automatically and saved for further analysis.

In order to determine whether stereoscopic or monoscopic viewing affected the reproducibility of CDR assessment, the observers first viewed the optic disc monoscopically by viewing the right hand image of the stereoscopic pair. The image was displayed using the same software package and using the stereoscopic viewing glasses (Stereographics) to control for the effects of viewing through LC lenses. Not less than 1 week later, the images were viewed stereoscopically and the CDRs recalculated. Images were analysed in this order so that stereoscopic cues could not influence grading of the monoscopic images.

![Diagram showing the image manipulation required to display stereoscopic images in the above and below format using a synchronous doubling emitter.](image1.png)

**Figure 1** Diagram showing the image manipulation required to display stereoscopic images in the above and below format using a synchronous doubling emitter. h, image height.

![Schematic diagram showing the criterion used for measurement of neuroretinal rim width. The cursor bar is set to the level of the neuroretinal rim (Elschnig’s rim).](image2.png)

**Figure 2** Schematic diagram showing the criterion used for measurement of neuroretinal rim width. The cursor bar is set to the level of the neuroretinal rim (Elschnig’s rim). The inset diagram of the optic disc shows how the bar can be located to any given sector of the disc and adjusted in length to match the width of the neuroretinal rim.

![Distribution of cup to disc ratios (CDR) for the three observers for stereoscopic assessments. Results from the three observers are represented by different symbols.](image3.png)

**Figure 3** Distribution of cup to disc ratios (CDR) for the three observers for stereoscopic assessments. Results from the three observers are represented by different symbols. Each observer made 1260 stereoscopic neuroretinal rim assessments over 35 discs.
Comparison between normally distributed variables was by Student’s *t* test. Non-parametric comparisons were made using the Mann-Whitney test. Differences between CDR estimates were expressed as means and standard deviations. The measure of agreement between the observers was quantified using the intraclass correlation coefficient (two way mixed effect model), which was determined for both monoscopic and stereoscopic images. In each case, statistical analysis was performed using SPSS 10.0.

RESULTS

High quality stereopairs were obtained from 35 eyes in 35 patients. Only one image was taken from each individual with 17 images from glaucomatous eyes, three from ocular hypertensive eyes, and 15 from healthy eyes. The mean age of the patients was 64 years (range 44–83 years). Each observer made 36 estimates of CDR in 35 eyes giving a total of 1260 CDR values. The distribution of stereoscopic CDR estimates for the three observers is shown in figure 3.

For monoscopic assessments, the mean CDRs for the three observers ranged from 0.57 to 0.66 (SD 0.13–0.14) compared with a 0.64 to 0.69 (SD 0.12–0.14) for the stereoscopic assessments. In figure 4 the mean stereoscopic CDRs (all observers, all locations around the disc, 35 discs) have been plotted against the corresponding monoscopic score showing that in the majority of cases the stereoscopic optic disc assessment provided a greater estimate of the CDR. Comparison within each observer showed this difference to be statistically significant at the 0.001 level (Mann Whitney test).

In figure 5 the difference in stereoscopic and monoscopic assessment has been plotted against the mean of the stereoscopic and monoscopic assessment for each sector around the optic disc (Altman and Bland plots, temporal, superior, nasal, and inferior). Positive differences in CDR estimate indicate a greater CDR estimate with stereoscopic compared with monoscopic viewing. It can be seen that for most assessments, the CDR estimate was greater when viewed stereoscopically compared with the monoscopic assessment. Although the monoscopic and stereoscopic assessment were statistically different for each segment (*p*<0.001, Mann Whitney), we did not find a significant difference between segments (ANOVA *p* = 0.40). Linear regression analysis suggests that the CDR assessment (stereoscopic) had minimal effect on the magnitude of the difference between stereoscopic and monoscopic assessment (*r*² = 0.06, *p*<0.001, standardised coefficient beta = 0.27).

The levels of intraobserver and interobserver differences and agreement for monoscopic and stereoscopic observations are shown in tables 1 and 2, respectively. For both monoscopic and stereoscopic assessments, intraobserver variation tended to be less than interobserver variation but this was not a marked effect. The intraclass correlation coefficient (ICC) for intraobserver agreements was calculated by comparison of measurements at the first and second examination sessions. The ICCs for interobserver agreements were based on observations made at the second examination session. For both monoscopic and stereoscopic observations the level of interobserver agreement tended to be higher than interobserver agreement. One notable exception was that observer 3 had a lower level of intraobserver agreement for monoscopic observations compared with interobserver agreement with observer 2 (table 1). The level of interobserver agreement was greater with stereoscopic compared with monoscopic observations. This effect was also seen for the level of agreement between all three observers; for monoscopic assessments the ICC between all three observers was 0.63 (CI 0.35 to 0.77) compared with 0.74 (CI 0.65 to 0.80) for stereoscopic assessments, indicating good agreement with monoscopic assessment and substantial agreement with stereoscopic assessment.20

DISCUSSION

We have described the use of digital stereoscopic optic disc assessment or the estimation of neuroretinal rim width in normal and glaucomatous eyes. The principal findings of the study are, firstly, that we found high levels of interobserver agreement for neuroretinal rim width for both monoscopic and stereoscopic assessments—with agreement being higher for stereoscopic assessment. Secondly, we found that stereoscopic optic disc assessment provided higher estimates of CDR (and therefore a lower estimate of neuroretinal rim width) compared with monoscopic assessment.

The level of agreement for both monoscopic and stereoscopic assessment compares favourably with that obtained by expert observers and exceeds that previously reported for similar grade staff.3 We found that intraobserver agreement was greater than interobserver agreement and in both cases this improved with stereoscopic observations. Several factors may account for the high level of agreement. The stereoscopic cursor enables the precise delineation of the inner margin of the neuroretinal rim with reference to the plane of Elschnig’s rim. The inner border of the neuroretinal rim has conventionally been defined as the point at which the retina surface steepens or changes slope at the margin of the optic disc24 which is highly subjective. To our knowledge this is the first study in which software has been developed so that the cursor can be used at the level of interest within a stereoscopic image. As such, our software allowed the observers to minimise parallax errors in the measurement of neuroretinal rim width. Previous reports of digital stereoscopic disc analysis did not describe the use of stereoscopic cursors. With the Discam system a single cursor is used, which is projected on one of the stereoscopic images and cannot be adjusted to lie within the correct measurement plane. It is possible that the presentation of the cursor to a single eye may introduce effects such as rivalry,25 which could complicate further accurate stereoscopic measurements. These technical differences may account for the findings of Parkin et al.13 who did not report significant difference between monoscopic and stereoscopic assessments of neuroretinal rim width using the Discam stereoscopic disc imaging system but did note a trend for monoscopic rather than stereoscopic measurements to provide a higher estimate of CDR.

Another factor accounting for the high level of agreement between observers is that the observers could adjust their estimate of neuroretinal rim width using cursor bars whose...
length matched the width of the rim. The CDR value was generated by the computer and was not, therefore, a factor in intraobserver or interobserver variation.

The tendency for stereoscopic optic disc imaging to provide higher estimates of CDR (and therefore lower estimates of neuroretinal rim area) has previously been reported. These findings are consistent with the relation between the shape of the optic nerve and the underlying optic nerve head colour. A recent analysis of digital stereoscopic imaging revealed different results with monoscopic estimates for the CDR being greater than stereoscopic estimates is not consistent with this view. It is possible that it relates to the use of monochromatic images (Discam uses a 256 grey scale compared with 24 bit colour resolution used in the present study) which may have obscured subtle variations in optic nerve head pallor at the margin of the neuroretinal rim. Although the clinical relevance of the differences between monoscopic and stereoscopic assessment has yet to be established, it is likely that it will influence the sensitivity with which progressive glaucomatous damage can be detected; for eyes with a CDR of 0.7 a 0.1 difference represents 30% of the remaining neuroretinal rim. Our findings suggest that stereoscopic imaging will permit earlier detection of neuroretinal rim changes compared with monoscopic viewing.

Technical improvements in the quality of the stereoscopic display may also improve sensitivity for the detection of progressive damage. As with others we have employed a synchronised doubled digital stereoscopic system to develop our prototype display because software development is straightforward, does not require specialised graphics cards, and can be implemented on computers and monitors with low specification. Unfortunately, the vertical resolution of the stereoscopic pair images is reduced by 50% and the images

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**Figure 5** Plot of the difference between stereoscopic and monoscopic optic disc assessments against the mean of the monoscopic and stereoscopic assessment. Symbols indicate the data from each observer. (A) Diagram showing angular subtense for each sector around the disc. (B) Temporal segment. (C) Superior segment. (D) Nasal segment. (E) Inferior segment.
are vertically offset by one scan line on the video monitor which results in a subtle degradation of the quality of the stereoscopic image. Higher resolution systems for the analysis of stereoscopic images are now available and are likely to supersede the above and below stereoscopic display format. With a suitable graphics card and a monitor with a refresh rate exceeding 100 Hz, it is possible to display right and left stereoscopic image pairs so that they occupy the same point in visual space on the monitor (so called quad buffered or page flipping technique) without the need for a small vertical offset. Image compression is not required so that images can be displayed at a higher resolution compared with interlaced displays. Another advantage of this technology is that it lends itself to the sequential analysis of aligned stereopairs for the detection progressive damage and will facilitate the clinical application of techniques such as stereochronoscopy.27 28

The final consideration in the assessment of the stereoscopic images is that we used sequential rather than simultaneous stereoscopic images. We selected this option since sequentially acquired views are those that are available in most clinics since they do not require specialised cameras or additional hardware. Our results should, therefore, be applicable to routine clinical practice. Since simultaneous stereoscopic images generally provide higher quality stereoviews29 we predict that these would highlight greater differences between monoscopic and stereoscopic analysis. We anticipate that developments in digital stereoscopy could have a valuable role in the refinement of the diagnosis of glaucomatous optic neuropathy and in the detection of the progressive glaucomatous damage. Given the advances that have been made in other digital imaging modalities, it is likely that digital stereoscopy can be used as a useful adjunctive diagnostic tool.

Table 1 Monoscopic observations

<table>
<thead>
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<th>Observer 1</th>
<th>Observer 2</th>
<th>Observer 3</th>
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<tbody>
<tr>
<td>Mean diffs*</td>
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<td>0.04</td>
<td>0.12</td>
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<tr>
<td>SD diffs†</td>
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<td>0.05</td>
<td>0.08</td>
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<tr>
<td>Range*</td>
<td>0.77</td>
<td>0.62</td>
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</tr>
<tr>
<td>ICC†</td>
<td>0.78</td>
<td>0.85</td>
<td>0.55</td>
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</tbody>
</table>

Summary of the difference between the observers for the estimation of CDR by monoscopic viewing. Mean diffs: mean differences are shown for CDR assessments at matched 10 degree intervals. SD diffs: standard deviation of the difference between measurements. ICC: intraclass correlation coefficient. *Intraobserver comparison. †Interobserver comparisons.

Table 2 Stereoscopic observations

<table>
<thead>
<tr>
<th></th>
<th>Observer 1</th>
<th>Observer 2</th>
<th>Observer 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean diffs*</td>
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<tr>
<td>ICC†</td>
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<td>0.74</td>
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</tbody>
</table>

Summary of the difference between the observers for the estimation of CDR by stereoscopic viewing. Table layout as for table 1.

REFERENCES


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