Altitudinal visual field asymmetry is coupled with altered retinal circulation in patients with normal pressure glaucoma

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Abstract

Aim—To compare the effect of altitudinal asymmetric glaucomatous damage on retinal microcirculation in patients with normal pressure glaucoma (NPG).

Methods—In a prospective cross-sectional study patients with NPG (washed out for antiglaucomatous therapy) and altitudinal asymmetric perimetric findings between the superior and inferior hemisphere (Humphrey 24–2) (n=18) were included and compared with 20 NPG patients with symmetrical field defects and 18 healthy subjects. Fluorescein angiograms were performed using a scanning laser ophthalmoscope. Using digital image analysis, arteriovenous passage time (AVP) and vessel diameters were assessed for comparison of corresponding affected and less affected temporal arcades.

Results—Both affected and less affected hemispheres showed significantly prolonged AVP times (p<0.001) when compared with healthy subject data. In hemispheres with more severe glaucomatous field loss the AVP times were significantly (p=0.04) prolonged compared with the less affected hemisphere (AVP affected 3.1 (SD 7) seconds vs AVP less affected 2.61 (1.4) seconds). There was no asymmetry effect on arterial and venous diameter measurements.

Conclusion—Altitudinal visual field defects are linked together with circulatory deficits of the retinal tissue. The attenuated circulation seems to be a considerable factor in the natural course of glaucomatous optic neuropathy.

In the pathogenesis of glaucomatous neuropathy, particularly in normal pressure glaucoma (NPG), disturbances of ocular circulation seem important in the natural course of the disease. Different techniques1–7 are employed to assess vascular dysfunction. Fluorescein angiography has been shown to detect ischaemic lesions of the optic disc.8 9 Circulatory fluorescein angiographic studies showed abnormalities of retinal and choroidal circulation in patients with chronic open angle or normal tension glaucoma.10 11 This may be attributed to the effect of decreased perfusion pressure due to elevated intracocular pressure, also a considerable factor since arteriovenous passage (AVP) was prolonged despite antiglaucomatous treatment.1 However, the correlation of the AVP to the degree of glaucomatous field damage has not yet been examined.

Early glaucomatous visual field defects are often asymmetric showing either superior or inferior nerve fibre bundle defects.12 Furthermore, particularly in NPG, eccentric cupping corresponds well with typical arcuate defects.13 Fluorescein angiography allows measurements of retinal circulation corresponding to visual hemifield defects. Past studies correlating visual function and microcirculation provided important comparative asymmetric data among both eyes on macular14 or retrobulbar circulation.15 16 In this study, we examined retinal microcirculation in patients with NPG and the correlation with altitudinal asymmetric visual field loss. This approach was chosen because a direct comparison of glaucomatous visual field defect and corresponding retinal microcirculation is possible. From existing data both a non-association or association of altered retinal circulation seems possible and the results could underline the significance of such measurement.

Materials and methods

Scanning laser video fluorescein angiograms (SLO; Rodenstock Institute, Ottobrunn, Germany) combined with an image analysis system (Matrox Inspector, Matrox Graphics, Quebec, Canada) allow the evaluation of AVP times and corresponding arterial and venous diameters. The method has been presented in detail elsewhere.17 In short, fluorescein angiograms at 40 degree field mode (20° × 20°) were performed. Dynamic sequences (25 Hz) were analysed using an image processing unit in order to measure the first appearance of the dye in a retinal artery and the first influx in the corresponding vein by dye dilution technique. Measurements were performed in the temporal superior and temporal inferior vessel formation. The AVP time characterises the shortest passage time through the vascular segment from the artery, via the capillary bed, to the corresponding veins.18 19 Past studies have shown that measurements of AVP times are reproducible and of clinical significance.1 16 17

At the location of the selected points for AVP measurements the arterial and venous diameters were quantified from densitometric curves by using a “half height of maximum height” procedure.16 19 The measurement location of the AVP and diameter measurement was situated half to one disc diameter away from the disc margin on the major temporal arteries and veins. Diameter measurements were performed in mid-transit fluorescein pic-
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PATIENTS

Prospectively, in this study 38 normal pressure glaucoma (NPG) patients (intraocular pressure ≤ 21 mm Hg) were recruited and 18 patients showed asymmetric visual field loss qualifying for further analysis. The topical antiglaucomatous medications of all patients were washed out for 3 weeks before examination. Figures 1 and 2 show asymmetric visual field loss and the corresponding fluorescein angiogram. The diagnosis was based on repeated IOP measurements (profile including midnight measurements), glaucomatous optic nerve head cupping (stereoscopic examination and argon laser imaging (SLO)), and visual field defects (Humphrey Instrument) as well as open angles (gonioscopy). Exclusion criteria included history of allergy to fluorescein and a refractive error ≥ -8 dioptres. The lens density was judged from standardised retroilluminated charts (LOCS III) and only subjects with mild nuclear colour changes (NC 1-2) were included.

Informed consent was obtained from each subject including detailed explanations of all procedures before participation in this study. The protocol for the study was reviewed and approved by the institutional review board of the Indiana University, Indianapolis, USA. The tenets of the Helsinki declaration were followed throughout the study.

In 18 patients (seven male, 11 female) asymmetric visual field damage was recorded. The mean visual acuity was 20/25. The field was defined as being asymmetric if the hemifields had different stages according to the classification by Aulhorn and Karmeneyer. The number of patients assigned to stages were: stage 1 three, stage 2 eight, stage 3 one, and stage 4 six. Twelve patients (66%) had asymmetric loss in the superior visual field and six patients (33%) in the inferior visual field. The standard visual field indices (MD = mean deviation, PSD = pattern standard deviation, MT = mean threshold) were derived from Humphrey 24-2 full threshold fields (MD = -7.3 (SD 4.5) dB; PSD = 9.1 (4.8) dB; MT = 21.8 (4.2) dB).

In 20 patients with NPG (six male, 14 female) no altitudinal defects were detected.

Figure 1 (A) Asymmetric visual field loss of the inferior visual field (Humphrey 24-2) with an arcuate scotoma in the inferior hemifield. (B) Corresponding fluorescein angiogram with focal ischaemia of the optic disc temporally superior (AVP time of the affected temporal superior vessel: 2.17 seconds; the less affected temporal inferior vessel formation: 1.67 seconds).
Among visual fields. The standard visual field indices were as following: MD = −3.7 (3.9) dB; PSD = 5.3 (3.9) dB; MT = 23.6 (4.2) dB. The healthy control group consisted of 18 subjects (eight male/10 female). The mean visual acuity was 20/20 and there was no evidence or history of any ocular or systemic disease.

**Statistical Analysis**

Mean value and standard deviation are given for all samples with normal distributions (Kolmogorov–Smirnov test) and non-normal distributions median and percentiles (10% and 90%). The paired Wilcoxon signed rank test was used to assess the significance of the differences between paired groups and the Mann–Whitney U test for unpaired comparisons. For multiple paired group comparisons the Friedman test was used and for unpaired group comparisons the Kruskall–Wallis test was used. Findings smaller than 0.05 were considered to be statistically significant. Pearson correlation coefficients were calculated to evaluate the relation between the variables. p Values were obtained after carrying out Fisher’s r to z transformations.

**Results**

The clinical and demographic data of the examined groups (NPG patients with and without altitudinal field asymmetry) did not differ significantly (Table 1). The average AVP times (mean of the superior and inferior vessel formation) were significantly prolonged (asymmetric NPG: p<0.001; symmetric NPG: p<0.001) when compared with healthy subject data.

In NPG patients with altitudinal asymmetric visual fields the mean deviation of the affected hemifield differed significantly from the less affected (Table 2). The AVP time in the affected field was significantly prolonged (12%) whereas there was no difference of arterial or venous vessel diameter (Table 3).

In patients with NPG and symmetric field defects the mean deviation of the affected hemifield differed significantly from the less affected (Table 2). These patients showed no significant differences between the temporal superior and inferior vessel formation for the AVP time and the arterial and venous diameters (Table 4). Additionally, no asymmetry effect between temporal superior and inferior vessels was observed for healthy subjects for AVP times and vessel diameters (Table 5).

The average AVP time of temporal superior and inferior vessel formation in the asymmetric NPG group showed no significant correlation with intraocular pressure, diastolic, or systolic blood pressure.

**Discussion**

A variety of circulatory disturbances are known in normal pressure glaucoma patients. Current concepts describe an energy deficit as the possible key for the initiation of a cytotoxic cascade. If an ischaemic change is thought to be causative the knowledge of the microcirculation may be important. Although the lamina and prelaminar region of the optic nerve is the area of major damage no existing method for microcirculatory quantification allows di-

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Table 1 Clinical and demographic data (mean (SD)) for the normal pressure glaucoma (NPG) patients with and without asymmetric visual field loss

<table>
<thead>
<tr>
<th></th>
<th>Healthy subjects</th>
<th>Asymmetric NPG patients</th>
<th>Symmetric NPG patients</th>
<th>Significance*</th>
</tr>
</thead>
<tbody>
<tr>
<td>No (male/female)</td>
<td>18 (8/10)</td>
<td>18 (7/11)</td>
<td>20 (6/14)</td>
<td>p=0.65</td>
</tr>
<tr>
<td>Age (years)</td>
<td>54 (15)</td>
<td>60 (15)</td>
<td>60 (15)</td>
<td>p=0.37</td>
</tr>
<tr>
<td>Systolic blood pressure (mm Hg)</td>
<td>135 (21)</td>
<td>123 (17)</td>
<td>119 (12)</td>
<td>p=0.11</td>
</tr>
<tr>
<td>Diastolic blood pressure (mm Hg)</td>
<td>75 (14)</td>
<td>74 (9)</td>
<td>73 (8)</td>
<td>p=0.62</td>
</tr>
<tr>
<td>Intraocular pressure (mm Hg)</td>
<td>16 (2)</td>
<td>17 (4)</td>
<td>16 (3)</td>
<td>p=0.97</td>
</tr>
<tr>
<td>Heart rate (beats/s)</td>
<td>77 (13)</td>
<td>80 (12)</td>
<td>71 (12)</td>
<td>p=0.33</td>
</tr>
</tbody>
</table>

*Kruskal–Wallis test and χ² test.

Table 2 Mean deviation of affected and less affected hemifields in patients with altitudinal asymmetric field defects are presented (median, 10, 90% percentiles). Furthermore the mean defects of the temporal superior and inferior field of NPG with symmetric field defects are given (median, 10, 90% percentiles)

<table>
<thead>
<tr>
<th></th>
<th>Mean deviation affected branch (dB)</th>
<th>Mean deviation less affected branch (dB)</th>
<th>Mean deviation superior field (dB)</th>
<th>Mean deviation inferior field (dB)</th>
<th>Significance*</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPG with altitudinal asymmetric field defects</td>
<td>−0.1 (−25.3; −3.2)</td>
<td>−1.7 (−6.9; 0.5)</td>
<td>−4.1 (−15.1; 0.5)</td>
<td>−3.4 (−15.4; 0.4)</td>
<td>p=0.002</td>
</tr>
<tr>
<td>NPG with symmetric field defects</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>p=0.88</td>
</tr>
</tbody>
</table>

*Wilcoxon signed rank test.
Table 3 Arteriovenous passage times (AVP) and corresponding arterial and venous vessel diameters of the affected and less affected vessel formations (altitudinal visual field loss) (mean (SD))

<table>
<thead>
<tr>
<th></th>
<th>AVP (seconds)</th>
<th>Arterial diameter (µm)</th>
<th>Venous diameter (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n=18</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Affected branch</td>
<td>3.1 (1.67)</td>
<td>99 (19)</td>
<td>135 (28)</td>
</tr>
<tr>
<td>Less affected branch</td>
<td>2.61 (1.44)</td>
<td>101 (23)</td>
<td>140 (30)</td>
</tr>
<tr>
<td>Significance*</td>
<td>p=0.04</td>
<td>p=0.89</td>
<td>p=0.87</td>
</tr>
</tbody>
</table>

*Wilcoxon signed rank test.

Table 4 Arteriovenous passage times (AVP) and corresponding arterial and venous vessel diameters of the temporal superior and inferior vessel formation in NPG patients without altitudinal visual field defects (mean (SD))

<table>
<thead>
<tr>
<th></th>
<th>AVP (seconds)</th>
<th>Arterial diameter (µm)</th>
<th>Venous diameter (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n=20</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Temporal superior</td>
<td>2.80 (1.60)</td>
<td>97 (20)</td>
<td>130 (22)</td>
</tr>
<tr>
<td>Temporal inferior</td>
<td>2.65 (1.33)</td>
<td>106 (18)</td>
<td>143 (26)</td>
</tr>
<tr>
<td>Significance*</td>
<td>p=0.66</td>
<td>p=0.12</td>
<td>p=0.17</td>
</tr>
</tbody>
</table>

*Wilcoxon signed rank test.

Table 5 Arteriovenous passage times (AVP) and corresponding arterial and venous vessel diameters of the temporal superior and inferior vessel formation in healthy subjects (mean (SD))

<table>
<thead>
<tr>
<th></th>
<th>AVP (seconds)</th>
<th>Arterial diameter (µm)</th>
<th>Venous diameter (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n=18</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Temporal superior</td>
<td>1.57 (0.40)</td>
<td>95 (9)</td>
<td>132 (18)</td>
</tr>
<tr>
<td>Temporal inferior</td>
<td>1.62 (0.37)</td>
<td>98 (9)</td>
<td>127 (21)</td>
</tr>
<tr>
<td>Significance*</td>
<td>p=0.24</td>
<td>p=0.45</td>
<td>p=0.44</td>
</tr>
</tbody>
</table>

*Wilcoxon signed rank test.

Rect evaluation of in vivo results. Retinal measurements appear of interest especially since nerve fibre haemorrhages are coupled with disease progression, and retinal nerve fibre loss is a characteristic feature of glaucomatous optic neuropathy particularly in patients with NPG.

In the presented study, the retinal arteriovenous passage (AVP) time of NPG patients is almost doubled compared with healthy subject data, indicating a severe attenuation of retinal circulation. These results are in agreement with previous studies in patients with chronic open angle glaucoma demonstrating prolonged AVP times and decreased mean dye velocities. The question arises, whether prolonged AVP times correspond with decreased perfusion or malnutrition of the tissue. Theoretically, a vaso-occlusive disease could result via decreased resistance and arteriovenous shunts in shortened AVP times. In patients with diabetic angiopathy AVP is prolonged with increasing severity of retinopathy. Furthermore, in NPG patients no retinal capillary closure has been detected.

Thus, AVP measurements representing the passage through from the artery passing the capillary formation to the vein are an indicator of the integrity and regulation of a vascular segment. The presented results are in agreement with other investigations revealing significantly decreased retinal circulation and pulsatile ocular blood flow in NPG patients when compared with healthy subjects.

Studies of visual field asymmetry have demonstrated considerable differences in NPG and high tension glaucoma suggesting different pathogenic mechanisms. Additionally, higher intraocular pressure is not necessarily associated with more severe visual field loss or a major cause resulting in asymmetric field damage. The comparison of psychophysical and circulatory data might provide information about the pathophysiology—for example, the question whether or not circulatory deficits may be a factor in disease progression.

In patients with ocular hypertension and primary open angle glaucoma blue field entoptic simulation demonstrated a significantly decreased macular leucocyte velocity in the more severely affected eye when performing binocular asymmetry analysis. In patients with primary open angle glaucoma, laser Doppler flowmetry showed decreased perfusion of the optic nerve head and juxtapapillary retina in patients with no or only borderline visual field defects. NPG patients with asymmetric visual field damage have decreased retrolubar velocities in the more affected eye and the decrease in retrolubar velocities was correlated with progressive visual field loss. Pulsatile ocular blood flow is reduced in NPG patients, regardless of the visual field loss when compared with healthy subjects, and significantly lower in the eye with more severe field damage. These circulatory studies did not account for regional differences of the superior and inferior portion of the optic nerve head or the corresponding retinal vasculature.

This study comparing superior and inferior altitudinal damage in one eye allows direct quantification of the affected circulatory bed versus the less affected one. In patients with NPG and altitudinal visual field loss the retinal AVP time as measured in the corresponding temporal superior or inferior vessel formation was significantly prolonged. In contrast, NPG patients with symmetric field defects, or healthy subjects, showed no circulatory asymmetry among superior and inferior vessel formations.

The effect observed in this study might represent a secondary effect due to the loss of nerve fibres and the downregulation due to reduced metabolic needs. However, the retinal segment of the less affected visual field also showed significantly prolonged AVP times compared with healthy subjects. Thus, a circulatory deficit might represent an early insult in glaucomatous disease. However, the primary cause remains unclear when visual field integrity does not necessarily reflect nerve fibre health.

Descending neuropathies (for example, retinitis pigmentosa) are typically associated with reduction of vessel diameters attributed to a metabolic downregulation. The arterial and venous diameter data in this study did not provide any significant association with altitudinal field loss. Past studies in patients with chronic open angle glaucoma showed that vessel diameters were correlated with visual field indices. The differences might be explained by the range of visual field data of the included patients; in other words, by including patients with severe field loss one would expect more pronounced narrowing of the vessel diameters. Those patients might be more affected by a downregulation as mentioned above. Thus, in this study with only moderate visual field loss, no significant alterations of vessel diameters were detected compared with the less affected hemifield.
In conclusion, circulatory deficits of the retinal tissue are linked together with visual field loss. Both the affected and less affected branches show prolonged AVP times when compared with healthy subjects. Circulatory changes should be considered in the treatment regimen when the cascade of events leading to loss of visual function is most amenable to being interrupted.

Proprietary interest: none.

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