Nerve fibre layer loss in diseases of the outer retinal layer

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SUMMARY We present examples of nerve fibre layer changes in diseases thought to affect primarily the outer retinal layers. These disease processes include cone-rod dystrophies, rod-cone dystrophies, juvenile macular degeneration (Stargardt's disease) and fundus flavimaculatus, vitelliform macular dystrophy, and Leber's congenital amaurosis. All were associated with alterations in the retinal nerve fibre layer, either diffuse or focal. The presence of these nerve fibre layer changes raises the question of transsynaptic degeneration and of possible primary or associated disease of other retinal cells than the receptors—namely, bipolar, amacrine, Mueller, or ganglion cells—in these diseases of the outer retinal layer. Involvement of the nerve fibre layer also indicates the need to examine patients with defects in the nerve fibre layer by-electroretinograms and other tests for outer retinal layer disease when no obvious optic nerve disease is present.

Clinically manifest damage to the anterior visual afferent pathways, especially the optic nerve, usually causes visible changes in the retinal nerve fibre layer. Not infrequently loss of the retinal nerve fibre layer gives the first clue of optic nerve compromise in glaucoma or multiple sclerosis and precedes the clinical manifestations, visual symptoms, or the visual field deficits. The changes in the retinal nerve fibre layer secondary to optic atrophy are quantitative changes, which appear as diffuse or focal atrophy of the nerve fibre layer. Qualitative changes in the nerve fibres themselves may be seen prior to optic atrophy and are characteristic of some diseases, such as Leber’s hereditary optic atrophy.

A number of ocular disorders known to involve primarily or exclusively the outer retinal layers are also associated with atrophic changes in the nerve fibre layer. To illustrate this point we present below examples of retinal receptor and/or pigment epithelium disorders associated with loss of the nerve fibre layer. Rake and wedge defects and diffuse nerve fibre changes are demonstrated in cone-rod dystrophies, rod-cone dystrophies (retinitis pigmentosa), juvenile macular degeneration, fundus flavimaculatus (Stargardt’s disease), vitelliform macular dystrophy, and Leber’s congenital amaurosis.

Subjects and materials
Patients with various types of retinal degeneration who clearly showed nerve fibre loss were selected from our files. In each case the diagnosis was made after a careful medical and family history and ophthalmic examination consisting of external and slit-lamp examination and direct and indirect ophthalmoscopy. In selected cases fluorescein angiography, electroretinography (ERG), electro-oculography (EOG), and dark adaptometry by standardised techniques were performed. All the findings were analysed to arrive at the most accurate diagnosis. A summary of the diagnoses and pertinent clinical findings can be found in Table 1.

Results
Fig. 1A–H. Patients 1–8 with outer retinal layer disease. All show rake defects (dark striations within grey translucent areas of more normal retinal nerve fibre layer (some arrowed) in temporal arcades and loss of retinal nerve fibre layer detail in the papillomacular bundle. Defects are less distinct than on funduscopy, as the prints are copies of colour transparencies. The diagnoses of the patients are given under the individual Figs.

Patient 1. A 25-year-old male with vitelliform macular dystrophy had best corrected vision acuity of
Table 1  Details of patients

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>Diagnosis</th>
<th>Visual acuity</th>
<th>Electro-retinogram</th>
<th>Goldmann fields</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>25</td>
<td>Vitelliform dystrophy</td>
<td>20/20</td>
<td>Normal</td>
<td>Minimal central depression OD; relative</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>20/50</td>
<td></td>
<td>central and paracentral scotomata OS</td>
</tr>
<tr>
<td>2</td>
<td>26</td>
<td>Simplex cone-rod dystrophy</td>
<td>20/30</td>
<td>1/3 of normal, photopic worse</td>
<td>Baring of blindspot; partial ring scotoma</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>20/40</td>
<td>affected</td>
<td>superiorly, contracted OU</td>
</tr>
<tr>
<td>3</td>
<td>31</td>
<td>Dominant cone-rod dystrophy</td>
<td>20/200</td>
<td>Barely recordable</td>
<td>Pseudoalitudinal defect OU</td>
</tr>
<tr>
<td>4</td>
<td>7</td>
<td>Juvenile Leber's congenital</td>
<td>20/70</td>
<td>Unrecordable</td>
<td>Less than 10° to IV OU</td>
</tr>
<tr>
<td></td>
<td></td>
<td>amaurosis</td>
<td>20/60</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>22</td>
<td>Stargardt's disease</td>
<td>20/40</td>
<td>Normal</td>
<td>Full with central scotomata OU</td>
</tr>
<tr>
<td>6</td>
<td>33</td>
<td>Rod-cone dystrophy</td>
<td>20/20</td>
<td>Barely recordable</td>
<td>Small central fields with peripheral islands</td>
</tr>
<tr>
<td>7</td>
<td>7</td>
<td>Fundus flavimaculatus</td>
<td>20/80</td>
<td>Normal</td>
<td>OU</td>
</tr>
<tr>
<td>8</td>
<td>44</td>
<td>Choroideraemia (carrier)</td>
<td>20/30</td>
<td>Borderline normal</td>
<td>Normal</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>OU</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

20/20 in the right eye and 20/50 in the left (Fig. 1A). ERG and dark adaptation were within normal limits, though the cone portion of the dark adaptation curve was somewhat low. The EOG was abnormal, with light peak to dark trough (Lp/Dt) ratios of 1.2 on the right and 1.35 on the left.

Patient 2 was a 26-year-old woman with cone-rod degeneration. Her visual acuity was 20/30 on the right and 20/40 on the left (Fig. 1B). The ERG showed a barely recordable photopic response, one-third normal scotopic responses, and unrecordable flicker responses. Visual fields showed baring of the blind spot and partial ring scotomata (Figs. 2 and 3).

Patient 3 was a 31-year-old woman with cone-rod degeneration of the autosomal dominant type. She had a visual acuity of 20/200 on the right and 20/100 on the left (Fig. 1C). The photopic ERG was barely recordable, the scotopic ERG one-half normal. Visual fields showed pseudoalitudinal defects with smaller isopters in the right eye (Fig. 4).

Patient 4 was a 7-year-old with juvenile Leber's amaurosis. Visual acuity was 20/70 on the right and 20/60 on the left (Fig. 1D). The ERG was unrecordable. The Goldmann field was less than 10° to a 1-4 target.

Patient 5 was a 22-year-old male with fundus flavimaculatus (Fig. 1E). Visual acuity was 20/40 on the right and 20/50 on the left. The ERG was normal. The EOG showed 155% rise. Visual fields were full, with central scotomata.

Patient 6 was a 33-year-old man with rod-cone degeneration (Fig. 1F). Visual acuity was 20/20 on the right and 20/30 on the left. The photopic and
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Fig. 1C Dominant cone-rod dystrophy.

Fig. 1E Stargardt's dystrophy.

Fig. 1F Rod-cone dystrophy.

Fig. 1D Leber's congenital amaurosis.

Scotopic ERGs were barely recordable. Dark adaptation was 4.25 log units elevated at 40 minutes. The visual field showed small central fields with very large ring scotomata and large peripheral islands (Fig. 5).

Patient 7 was a 7-year-old girl with fundus flavimaculatus (Fig. 1G). Visual acuity was 20/80 OU. The ERGs were normal bilaterally, as were the Goldmann fields.

Patient 8 was a 44-year-old woman who was a carrier of choroideraemia (Fig. 1H). Visual acuity was 20/30 in each eye. The photopic ERG was borderline and the scotopic ERG normal on the right and borderline on the left. The dark adaptation was 1.9 log units elevated at 40 minutes. Visual fields were normal.

Discussion

We present eight examples of atrophy of the retinal nerve fibre layer in patients with outer retinal layer diseases. These changes are very frequently present when sought but may be difficult to detect against a background of generalised atrophy of the nerve fibre layer and/or retinal pigment epithelium and other pigmentary aberrations. In general the changes in the nerve fibre layer correlate well with the severity of the disease process. However, it is often difficult to link individual nerve fibre layer defects with specific field defects where there are also field defects due to the basic outer retinal layer disease process. Once nerve fibre layer atrophy appears, its progression
appears to parallel the progression of the disease, and when advanced stages of disease are reached, in which generalised retinal atrophy is present, the wedge-shaped defects have disappeared.

In previously reported cases of damage to the retinal nerve fibre layer the primary insult has been to the optic nerve or another portion of the ganglion cell axon, such as the chiasm. The consequent changes in the nerve fibre layer represent direct injury to that axon, not evidence of remote effects or transsynaptic degeneration.

Thus the discovery of changes in the nerve fibre layer in disorders of the outer retinal layer raises important questions about the pathogenesis of these changes. Is there transsynaptic degeneration, or do these diseases of outer retinal layers actually directly involve the inner retinal layers as well?

Transsynaptic degeneration has been thought to occur only rarely and to minor degrees in adult humans or animals. For this reason Hoyt and associates were able to point out the significance of one pattern of nerve fibre layer change (homonymous hemianopic hypoplasia) as indicative of congenital or very early injury to the postgeniculate central nervous system.

If the diseases in question truly affect the outer retinal layers predominantly, then an explanation is needed for transsynaptic degeneration occurring after infancy.

Studies in monkeys confirm the clinical impression that transsynaptic degeneration is rare (or hard to detect) in the central nervous system of mature animals. Ablation of the striate cortex of immature monkeys produces approximately 70% loss of retinal ganglion cells, much less ganglion cell loss in adult animals. However, some retrograde transsynaptic degeneration does occur in adult monkeys. Dineen et al. ablated the striate cortex of adult monkeys and found a 30% loss of parafoveal retinal ganglion cells. In addition physiological evidence of transsynaptic transport of neural materials might provide a logical mechanism by which transsynaptic effects are mediated in adult animals.

**Involvement of outer retinal layers**
Electrophysiological studies suggest that some diseases generally considered to affect predominantly inner or outer retinal layers, but not both,
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influence many retinal layers. Kirkham and Coupland found electroretinographic changes in papilloedema due to increased intracranial pressure, and Fazio et al. reported alterations of the single flash electroretinogram in advanced glaucoma. Both of these disease processes are thought to affect the ganglion cell axon primarily. The pattern electroretinogram is considered to reflect ganglion cell (or at least inner retinal) activity, but it is significantly diminished in Stargardt's disease, an outer retinal layer process (and Newman NM, Jackson JL, Latner B, personal communication).

Other aberrations of the ganglion cell axon are not uncommon in diseases of the outer retina. Clinically, waxy pallor of the optic disc has long been recognised as a sign in advanced rod-cone dystrophies, but neural atrophy has gone unnoted and is not discussed as a pathological feature of outer retinal layer disease. Unusual drusen also are noted in retinitis pigmentosa, frequently occurring in the nerve fibre layer away from the disc substance, in contrast to optic nerve head drusen which are confined to the papilla. In addition the temporal portion of the disc and its nerve fibre layer appear to be severely diminished or missing in patients with X-linked congenital stationary night blindness, and similar changes and telangiectasis are extremely common in the cone-rod degenerations.

Thus substantial anatomical, physiological, electrophysiological, and clinical evidence indicates nerve fibre layer involvement in outer retinal layer disease and is consistent with our observations of atrophic nerve fibre layer changes.

The observed changes appear to be signs of disease processes initiated in early, even prenatal, development because many of the changes we illustrate are found in children. Early outer retinal damage may cause transsynaptic changes. In primate studies on the retinal degeneration from striate cortex lesions the severity of transsynaptic degeneration was directly related both to age and to time—for example, the younger the animal and the longer the observation period following the injury, the greater the degeneration. Alternatively, some of the mentioned disease processes may involve metabolic and physiological changes which occur in many retinal layers but appear clinically as changes in the outer retinal layers predominantly.

Awareness of the occurrence of these changes in the nerve fibre layer in outer retinal layer disease has at least one practical implication. The presence of nerve fibre layer changes not readily explained by optic nerve disease should prompt examination of the patient for diseases of the outer retinal layer.

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References


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