Aging and degeneration of the human macula.
1. Outer nuclear layer and photoreceptors

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SUMMARY  In a light microscopic study of the macula of 104 human eyes obtained at necropsy of patients aged 3 to 96 we found: (1) Displacement of nuclei from the outer nuclear layer into the outer plexiform layer occurred in small numbers early in life and markedly increased after age 30. (2) Displacement of nuclei from the outer nuclear layer to the layer of rods and cones was rare in early life but increased considerably after age 40. (3) Displacement of nuclei is probably secondary to shrinkage of their attached fibres and is associated with aging. (4) Displaced nuclei apparently undergo changes in size, shape, and chromatin content and may go on to necrosis. (5) Twenty-four of the 104 eyes studied had an obvious reduction in the numbers of nuclei in the outer nuclear layer and their photoreceptors in the macular zone. All were in eyes from patients over age 40. No concomitant defect was found in the subjacent pigment epithelium, Bruch’s membrane, or the choriocapillaris. The loss of nuclei of the outer nuclear layer appears to be a primary retinal disorder associated with aging.

We found in the macula a loss of nuclei from the outer nuclear layer (ONL) which increased with age. This loss apparently starts with displacement or migration of nuclei from the ONL into the outer plexiform layer (OPL) and into the layer of rods and cones. This observation has received little or no attention and we could not find any previous systematic study in the literature.

Material and methods

One-hundred-and-four eyes obtained at necropsy from patients aged 3 to 96 were selected for study because sections included the macular region. Fifty-two of these eyes had sections through the fovea. Only eyes without significant inflammatory or degenerative disease or trauma were studied. Eyes with disciform degeneration or with pigment migration into the retina were excluded. The eyes were opened in the horizontal meridian so that sections included the disc and macula. They were prepared with celloidin embedding cut 14 μm thick and stained with haematoxylin and eosin. The macula includes the fovea and parafoveal zone. Its outer boundary was the point where the ganglion cell layer was less than 2 cells thick. Sections were examined by light microscopy and the total numbers of displaced nuclei in the macula counted for each section. At least 3 slides of each eye were examined and the results averaged. The data were graphed (Figs. 1 and 2).

Results

The normal nuclei in the ONL appear uniform with a circular or slightly ovoid shape and contain many fine chromatin granules. Several layers of cone nuclei are concentrated at the fovea. Elsewhere in the macula cone nuclei are near the external limiting membrane and appear slightly larger and paler than the rod nuclei.

The number of nuclei in the ONL does not remain constant during a long lifetime. Significant changes are found with aging. Some nuclei from the ONL become displaced into the OPL (Fig. 1) and others into the layer of rods and cones (Fig. 2).

Displacement of nuclei into the OPL at the macula is evident even early in life but increases after age 30, and is most pronounced after age 50 (Fig. 1). The nuclei displaced into the OPL appear to be mainly rod nuclei from the ONL and perhaps
some nuclei from the inner nuclear layer (INL) (Fig. 3).

Displacement of nuclei from the ONL through the external limiting membrane to the rod and cone layer has been described as a normal variant observed most often in the macula. However, such displacement is rare in youth (Fig. 2). Only after age 40 is it frequent, being most common after age 50. A decrease in such nuclei was noted after age 70. Nuclear displacement is rare in the centre of the fovea and was noted mainly in the rest of the macula (Figs. 4a and 4b).

Nuclei displaced into the layer of rods and cones lose their normal configuration and often appeared enlarged and oval with their long axis parallel to the rods and cones. The nuclei displaced in the OPL varied in shape. A few retained their spherical form, while many became elongated with their long axis parallel to the oblique nerve fibres of Henle's layer. A number were enlarged and pale, some appeared

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**Fig. 1 Nuclei displaced into outer plexiform layer.**

**Fig. 2 Nuclei displaced into rod and cone layer.**
very irregular, and a few seemed necrotic. In several instances we observed similar degeneration of nuclei within the ONL.

Twenty-four of the 104 eyes studied had discrete areas in the macula with an appreciable loss of nuclei in the ONL. The loss was extensive in 9 and moderate in 15 eyes (Table 1 and Figs. 5 and 6). The loss was usually in small focal areas, but some eyes had a broad zone with a diminished number of nuclei. The loss of nuclei was usually found opposite normal or only slightly defective retinal pigment

Fig. 3 Male age 63. Parafoveal zone. There is a large number of displaced nuclei in the OPL. Most of these nuclei are pale, deformed and elongated and lie parallel to the obliquity of the fibres in Henle’s layer. Some nuclei in the ONL are elongated and deformed. Some of the photoreceptors have lost part or all of their outer segments. (H and E, ×15).

Figs. 4 upper and lower. Female age 54. Parafoveal zone. Many nuclei from the ONL are displaced into the inner layer of rods and cones. Some of the displaced nuclei have lost their regularity and are elongated and paler than in the ONL. The pigment epithelium and Bruch’s membrane appear normal. The choriocapillaris has marked thickening of its anterior and lateral walls. (Upper: H and E ×15; lower: H and E, ×38).
epithelium, Bruch's membrane, and choriocapillaris. Accompanying the loss of nuclei in the ONL we noted shrunken and deformed rods and cones and a diminution in the numbers of photoreceptors (Figs. 5 and 6). The reduced population of cells in the ONL resulted in a loss of the axons traversing the outer plexiform layer, and that layer appeared thinner than normal (Fig. 4).

Three of the 24 eyes with marked loss of cells from the ONL had interesting fellow eyes. In a patient aged 85 there was typical senile pigmentary degeneration of the macula in the other eye, and 2 patients, aged 71 and 94, had disciform degeneration of the macula in the second eye.

Table 1  Loss of nuclei in outer layer in 104 eyes

<table>
<thead>
<tr>
<th>Age</th>
<th>Numbers of eyes</th>
<th>Marked loss of nuclei</th>
<th>Moderate loss of nuclei</th>
<th>% Affected</th>
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<tbody>
<tr>
<td>Before age 40</td>
<td>23</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>40-49</td>
<td>10</td>
<td>3</td>
<td>2</td>
<td>50</td>
</tr>
<tr>
<td>50-59</td>
<td>21</td>
<td>0</td>
<td>7</td>
<td>33%</td>
</tr>
<tr>
<td>60-69</td>
<td>20</td>
<td>2</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td>70-79</td>
<td>17</td>
<td>2</td>
<td>2</td>
<td>22</td>
</tr>
<tr>
<td>80-89</td>
<td>9</td>
<td>1</td>
<td>2</td>
<td>33%</td>
</tr>
<tr>
<td>90-100</td>
<td>4</td>
<td>1</td>
<td>0</td>
<td>25</td>
</tr>
<tr>
<td>Total</td>
<td>104</td>
<td>9</td>
<td>15</td>
<td></td>
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</tbody>
</table>
Aging and degeneration of the human macula

Discussion

In 24 of 104 necropsy eyes without significant ocular disease we found a reduced population of nuclei in the ONL along with a loss of their photoreceptors. These observations of human eyes parallel the recent finding that a progressive loss of cells occurs with aging in the ONL in healthy rats.2

A loss of retinal neurons and photoreceptors at the macula has been demonstrated by many authors in eyes that had pigmentary migration into a degenerated retina or a disciform response.3-9 Those who have studied senile macular degeneration have been impressed with the associated changes in the layers adjacent to the retina, and they have picked one or another layer to assign as the primary site with macular degeneration as a secondary consequence. There is no question that with aging a variety of changes occur in the retinal pigment epithelium, Bruch’s membrane, and the choroidal vasculature. These structures are all so closely interrelated that it has been difficult to ascertain the site of a primary defect. However, the present study indicates that a progressive degeneration of cells in the ONL and their photoreceptors may occur without significant changes in the adjacent layers.

The elongation of nuclei found in the rod and cone layer has been explained as a consequence of com-
pression by the adjacent photoreceptors. Compression would not, however, explain the change in shape of the nuclei displaced into the OPL. That layer of fibres (Henle’s layer) in the macular zone is easily separated by fluid in cystoid oedema and by exudates in diabetes.

What is the explanation for nuclear displacement from the ONL? The nuclei in the ONL are attached at one end to the photoreceptors and on the other to their axonal fibres, which cross the OPL and join with dendrites from the INL. We speculate that nuclear displacement in either direction may be due to traction exerted by shrinkage of the attached fibres. A traction phenomenon could explain elongation of the displaced nuclei in the direction of the nerve fibres in the OPL as well as in the layer of rods and cones. Less likely is the possibility of active nuclear migration.

It is noteworthy that Marshall et al.11 have demonstrated by electron microscopy a change with aging in human rods. Starting at age 40 some rods become elongated and convoluted, and by the seventh decade 10 to 20% of the rods are affected. The degenerative changes we found also appear after the fourth decade.

A normal retina has about 130 000 000 nuclei in the ONL with a corresponding number of photoreceptors.12 In our material we could not accurately count the changes in numbers of nuclei with aging. Our report is thus qualitative and notes only a substantial loss that is obvious by light microscopy. We found that an appreciable number of nuclei were displaced from the ONL after age 30. Some had undergone degeneration characterised by a change in their size and shape as well as the density of chromatin; in some only a cellular ghost remained. Nuclear displacement probably accounts for the loss of cells in the ONL. In our study we found no correlation between the loss of nuclei in the ONL and changes in the pigment epithelium, Bruch’s membrane, or the choriocapillaris. It appears that the cellular diminution in the ONL can be primary and is most likely an aging phenomenon. Along with the loss of cells in the ONL there was similar reduction in number of the attached rods and cones and the axons in the outer plexiform layer. The relation of this process to senile macular degeneration is unclear and requires further investigation.

The loss of nuclei in the ONL and their photoreceptors is likely to cause some change in the visual properties of the eye. It is well known that there are parameters of visual function which diminish with aging without obvious clinical alterations in the lens or the fundus. Our findings may help to explain some of this loss of function.

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References