Age related compliance of the lamina cribrosa in human eyes

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Abstract

Aims—To investigate changes in the mechanical compliance of ex vivo human lamina cribrosa with age.

Methods—A laser scanning confocal microscope was used to image the surface of the fluorescently labelled lamina cribrosa in cadaver eyes. A method was developed to determine changes in the volume and strain of the lamina cribrosa created by increases in pressure. The ability of the lamina cribrosa to reverse its deformation on removal of pressure was also measured.

Results—Volume and strain measurements both demonstrated that the lamina cribrosa increased in stiffness with age and the level of pressure applied. The ability of the lamina cribrosa to regain its original shape and size on removal of pressure appeared to decrease with age, demonstrating an age-related decrease in resilience of the lamina cribrosa.

Conclusions—The mechanical compliance of the human lamina cribrosa decreased with age. Misalignment of compliant cribriform plates in a young eye may exert a lesser stress on nerve axons, than that exerted by the rigid plates of an elderly lamina cribrosa. The resilience of the lamina cribrosa also decreased with age, suggesting an increased susceptibility to plastic flow and permanent deformation. Such changes may be of importance in the explanation of age-related optic neuropathy in primary open angle glaucoma.


The lamina cribrosa is a lattice-like structure, consisting of successive perforated cribriform plates across the optic nerve canal, through which pass bundles of nerve fibres. The plates are lined by basement membranes and their cores are filled with significant amounts of fibrillar collagens and elastic fibres. During ageing, the constituents of these cribriform plates are altered.1–4 It is likely that such changes will alter the mechanical behaviour of the ageing lamina cribrosa and therefore compromise its ability to support the nerve axons that pass through it.

The distribution of axonal loss in early glaucoma appears to be related to regional differences in architecture of the lamina cribrosa. The superior and inferior regions of the lamina cribrosa, which exhibit greater pore size and less dense connective tissue, are thought to be more susceptible to axonal damage.5–6 The arcuate retinal ganglion cell axons, which pass through these regions, are damaged first in early glaucoma.

The cause of glaucomatous axonal damage is unknown; ischaemia and mechanical damage as a result of a raised intraocular pressure or rearrangement of the cribriform plates are among the theories that have been proposed.7–9 Whatever the cause, compression of the lamina cribrosa appears to occur before detectable glaucomatous field loss.10

Age-related changes in the lamina cribrosa include increases in the amounts of collagen types I, III, and IV11–14 within the cribriform plates, which lead to an overall increase in total collagen content.15–17 We have also demonstrated alterations in the ratio of collagen types I and III and an increase in non-enzymatic glycation.18 Remodelling of extracellular matrix within the lamina cribrosa has been attributed to the heterogeneity of astrocytes present.19–21

These changes in biochemical composition of the ageing lamina cribrosa may affect the mechanical integrity and therefore the susceptibility of elderly eyes to glaucomatous optic neuropathy. This study is the first to investigate the effect of these structural changes on the mechanical compliance of the lamina cribrosa as a function of age.

Methods

SOURCE OF TISSUE

Ten human eyes aged between 7 and 86 years were collected within 48 hours of death from the Bristol Corneal Transplant Service Eye Bank, after their corneas had been removed for transplantation. The eyes had no known pathology and diabetic eyes were excluded. Each eye was processed and analysed on the day of collection.

EXPERIMENTAL DESIGN

The posterior section of the eye, encompassing the back third of the sclera including the optic nerve, was isolated from each eye. The vitreous, retina, and choroid were removed and the interior rinsed with 0.9% saline. The glial tissue overlying the lamina cribrosa was dissected away, and the lamina cribrosa was stained using 0.5 mg/ml fluorescein isothiocyanate (Sigma) in 50 mM bicarbonate buffer, pH 9.2 for 1.5 hours. Following rinsing, and a 30 minute incubation in bicarbonate buffer, the tissue was then sealed to a glass slide using cyanoacrylate glue (Loctite UK, Welwyn Garden City, Herts). Air in the cavity was drawn out using a 23 gauge hypodermic needle and replaced with saline via another 23 gauge nee-
dle connected to a graduated reservoir (Fig 1). The pressure within the cavity was controlled by raising or lowering the graduated reservoir, which was used as a simple manometer. As pressure increases the lamina cribrosa distends into a cup-like shape. The shape and volume bounded by the distorted lamina cribrosa was measured using a Bio-Rad MRC-500 scanning confocal microscope. The optical sectioning facility of this instrument permits images in the plane of the lamina cribrosa (xy scans) and in a plane perpendicular to the lamina cribrosa (xz scans or yz scans) to be obtained. In each case, one xz and one yz scan, together with an xy scan were taken in order that the three dimensional shape of the cup could be calculated. The preparation was oriented such that a line joining the two central retinal vessels was parallel to the x axis of the scan. After an xz and xy scan had been performed, the preparation was rotated by 90°, and the xz and xy scans were repeated.

The pressure was raised in steps from atmospheric pressure to 43 mm Hg above atmospheric (that is, 0, 11, 22, 33, and 43 mm Hg) and then lowered back down stepwise to the original pressure. At each step the lamina cribrosa was allowed to equilibrate to the new pressure level for 15 minutes before xy and xz scans were performed as described above. Any eye, which leaked at any stage during the performance of the experiment, was discarded and a new experiment started using a fresh eye.

MEASUREMENT OF VOLUME
The photographic negatives of the xz scans were enlarged and the surface of the lamina cribrosa, identified by the fluorescent marker fluorescein isothiocyanate, was traced for each pressure induced deformation. The centre of gravity (c) and the area (A) bounded by each distorted cup was quantitated by a computer linked visual imaging display system (VIDS) linked to a Hipad digitiser (Houston Instrument, TX, USA).

Thus the volume (V) of the resulting cup was calculated using these parameters in the theorem of Pappus14:

\[ V = 2\pi Ac \]

The calculated volume was multiplied by 1.3346 (refractive index of saline) in consideration of the foreshortening of depth measurements produced by imaging the lamina cribrosa across an air saline interface.

CALCULATION OF REVERSIBILITY
The ability of the lamina cribrosa to recover its shape was measured by the reduction of the volume of the distended cup formed by the lamina cribrosa on removal of applied pressure.

Reversibility was calculated as: volume recovered on removal of pressure/total volume change

\[ \% \text{ Reversibility} = \frac{(100\% - \text{residual volume})}{(V_{43} - V_0)} \]

where:

- \( V_{43} \) = volume at 43 mm Hg above atmospheric pressure
- \( V_0 \) = volume at atmospheric pressure following reduction of pressure

Residual volume = (volume at atmospheric pressure following reduction of pressure) − (volume at atmospheric pressure before pressure has been applied).

MEASUREMENT OF STRAIN
In order to check for anisotropy in the deformation of the lamina cribrosa, the strain in the x direction and the y direction were measured separately at each applied pressure. Each enlarged xz and yz scan was converted into a digital image using a Scanjet Plus (Hewlett Packard) into a computer drawing package (Deskscan). The z coordinates of each scan were multiplied by 1.3346 to correct for depth measurement and the xz and yz scans were reconstructed. The perimeter of the depression bound by the deformed lamina cribrosa was measured using VIDS.

The extent of the deformation of the lamina cribrosa was calculated as perimeter strain (change in length/original length at atmospheric pressure) for each pressure.

\[ \text{Strain} = \frac{(S_x - S_0)}{S_0} \]

where \( S_0 \) = strain at atmospheric pressure before pressure increase and \( S_x \) = strain induced by an applied pressure (for example, 11, 22, 33, or 44 mm Hg) above atmospheric pressure

DETERMINATION OF MECHANICAL COMPLIANCE
Differential equations of pressure-strain plots were calculated for each eye to determine the
incremental gradients (change in pressure/change in strain = dP/dS) of these curves at known pressure values. The incremental gradients were then plotted as a function of the pressure applied and age in order to assess the mechanical compliance of the lamina cribrosa.

**STATISTICAL ANALYSIS**

Spearman’s rank correlation was used to test the tendency of values of Y (that is, reversibility, incremental gradients) to increase (or decrease) as values of X (age) increase.

**Results**

**PRESSURE INDUCED DEFORMATION OF THE LAMINA CRIBROSAL**

A typical xy scan is shown in Figure 2, illustrating the fluorescently labelled surface of the lamina cribrosa. Figure 3 demonstrates a typical xz scan, illustrating the backward bowing of the lamina cribrosa after the pressure had been increased to 43 mm Hg above atmospheric pressure. These scans were used calculate the volume and strain of the deformation created at each pressure in each lamina cribrosa.

**Volume**

Figure 4 shows the applied pressure versus the volume bounded by the cupped lamina cribrosa for two extremes of the age range investigated. Initially a large volume change at low pressure levels was seen, but as the stress applied increased with increasing pressure, the lamina cribrosa became progressively stiffer. Stiffness is therefore defined as the ability of the lamina cribrosa to be deformed as pressure was applied.

The volume of the lamina cribrosa increased and decreased in all ages as the pressure applied was raised and lowered respectively. The gradient of the loading curve increased with age, as shown by comparison of the two curves in Figure 4. Gradients of curves from other ages fell between those demonstrated by the 9 and 86 year old in Figure 4. This illustrates that pressure induced deformations of the lamina cribrosa were greatest in the younger eyes and progressively decrease with age.

**Reversibility**

Reversibility of the lamina cribrosa after loading, as measured by the recovery towards original volume on reduction of pressure, decreased with age from 80% recovery in the 7 year old to less than 30% in the 86 year old (Fig 5, \(r^2=0.776\), line shown is a fitted second order polynomial regression \(y=81.089 - 0.89614x + 4.153e^{-3x^2}\)). The Spearman’s rank correlation coefficient \(r_s=0.745\), which is significant at 0.013. Therefore the ability of the ageing lamina cribrosa to recover its original
shape and volume after a pressure induced deformation was progressively reduced with age.

**Strain**
Pressure strain curves were similar in shape to the pressure-volume in Figure 4. Their shape illustrates the varying degrees of non-linearity in the lamina cribrosa over the range of applied stress and strain. To quantify this, gradients at several points on the pressure-strain curves were calculated from the differential of the equations from polynomial regression of these curves. The incremental gradients of each curve, plotted as a function of age and pressure (Fig 6), increased with age and pressure, indicating a stiffening of the lamina cribrosa. Second order polynomial regression curves of incremental gradients versus age and pressure demonstrated varying correlation coefficients with r values between 0.65 and 0.75 for all curves shown.

Figure 7 Strains in two directions across the lamina cribrosa S1 and S2: the strain in one of the two directions was always greater in one dimension.

Incremental gradients (dP/dS, that is to say the change in pressure relative to the change in strain) were calculated for each pressure point from a polynomial regression curve through the loading curve of each lamina cribrosa. The incremental gradient defined the change in the mechanical compliance of the lamina cribrosa with age, as pressure was increased. (A) Incremental gradients of the pressure-strain curves for three of the 10 eyes analysed: 7 (squares), 44 (triangles), and 86 (circles) year old, versus the pressure applied to produce each strain. The incremental gradient increased with age and pressure, indicating a stiffening of the lamina cribrosa. (B) Incremental gradients dP/dS versus age for given pressure increments, indicating a rapid increase in stiffness of the lamina cribrosa after 40–50 years of age. Lines show second order polynomial regression. The correlation values (r) vary between 0.65 and 0.75 for all curves shown.

Some individuals demonstrated very clear differences in the strains measured in the two directions (perpendicular to each other) at all pressures, as demonstrated in Figure 7, suggesting evidence of an anisotropy in the deformation of the lamina cribrosa.
Discussion
The mechanical properties of biological soft tissues are dependent upon the interaction of its individual components. The extracellular matrix of the lamina cribrosa is composed of elastin, collagen, and proteoglycans which define in Figure 4 the non-linear properties of a composite under stress. This mechanical behaviour can be interpreted in terms of the highly extensible elastin components and relatively stiff and inextensible collagen fibres, which have an elasticity modulus 1000 times greater than elastin. At low stress, it is likely that the elastin component allows the initially large volume change formed by the distending lamina cribrosa and as pressure increases and the extended collagen fibres limit further deformation, the lamina cribrosa becomes progressively stiffer. Various in vivo techniques have been designed to quantitate optic disc topography in glaucoma patients and optic disc displacement in animal models of experimental glaucoma. Pressure induced displacement of the optic nerve head in human cadaver eyes has been monitored by scanning laser tomography, and radiographic evaluation of platinum wires placed in the lamina cribrosa. In contrast with these in vitro methods, the technique described in this study identifies the three dimensional parameters of the pressure induced deformation at the level of the human lamina cribrosa, and uses this information to determine the biomechanical properties of the tissue as a function of age.

Previously described alterations in the extracellular matrix of the ageing human lamina cribrosa include an age related increase in the non-enzymatic collagen cross link pentosidine, total collagen and elastin content, and decreased type III to type I collagen ratio and total sulphated glycosaminoglycans content. Consistent with our findings in this study identifies the three dimensional parameters of the pressure induced deformation at the level of the human lamina cribrosa, and uses this information to determine the biomechanical properties of the tissue as a function of age.

Reversibility of the lamina cribrosa towards original volume on reduction of pressure decreased with age, indicative of decreased resilience. When force is applied and then removed cyclically, many materials (including biomaterials) demonstrate fully recoverable (that is, elastic) deformations up to a certain point (the elastic limit). However, after the elastic limit, plastic flow occurs and the internal structure of the tissue may be permanently altered. In such cases strain does not return to zero on removal of stress. Plastic flow can continue in a tissue until it reaches its failure strength. In this study, the difference between the volume after complete removal of pressure and the original volume is known as the residual volume (see Fig 4) and is representative of permanent deformation (plastic flow) which has occurred within the lamina cribrosa. The age related decrease in reversibility following pressure induced deformation implies that permanent deformation in structure as a result of plastic flow increases with age. In vivo studies in monkeys have demonstrated chronic posterior deformation of the optic disc surface, indicative of plastic flow, 3–4 weeks following the onset of experimentally induced glaucoma. Consistent with our findings of higher resilience in younger laminae cribrosae is the reversibility of cupping seen in infant glaucoma and young adults, perhaps due to higher yield points or their ability to undergo larger deformations up to their elastic limit. The degree of damage and mechanical compliance of the cribiform plates differs depending on age and the level of pressure applied; it is likely that age related stiffening of the lamina cribrosa limits its ability to distend before permanent deformation occurs.

These alterations in mechanical properties of the ageing lamina cribrosa are likely to influence the susceptibility of the retinal ganglion cell axons to damage due to the close anatomical arrangement of the cribiform plates and the nerve axons, which pass through them. In a young eye the stress exerted on the nerve axons during the compression and rearrangement of the cribiform plates may be effectively much less than in the more rigid structure of the elderly lamina cribrosa. The change in mechanical compliance appears to be most marked after 40–50 years, the age at which the incidence of primary open angle glaucoma (POAG) is increased.

In primary open angle glaucoma, the intraocular pressure is often raised and larger fluctuations in the diurnal variation of intraocular pressure occur. Each time the pressure fluctuates above the yield point, the extracellular matrix of the lamina cribrosa may undergo plastic flow with resultant permanent damage. Rigidity of the lamina cribrosa increases with progression of the disease, perhaps indicative of accumulative damage within the connective tissue plates, compatible with total misalignment of the cribiform plates seen in advanced glaucoma. The alteration in mechanical compliance of the lamina cribrosa may be significant in the
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