Laser-Induced Mouse Model of Chronic Ocular Hypertension

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PURPOSE. To develop an inducible mouse model of glaucoma. METHODS. An obstruction of aqueous humor outflow in adult C57BL/6J mice was induced by combined injection of indocyanine green (ICG) dye into the anterior chamber and diode laser treatment. To evaluate intraocular pressure (IOP), tonometry was performed with a modified Goldmann tonometer. The function of the retina was evaluated with electroretinography (ERG).

RESULTS. IOP was significantly elevated in surgical eyes compared with control eyes: before surgery, 15.2 ± 0.6 mm Hg; 10 days after surgery, 33.6 ± 1.5 mm Hg (P < 0.001); and 30 days after surgery, 27.4 ± 1.2 mm Hg (P < 0.001). However, 60 days after surgery, IOP in the surgical eyes decreased to 19.5 ± 0.9 mm Hg and was not significantly different compared with control eyes (control, 17.3 ± 0.7 mm Hg; P = 0.053). ERG amplitudes, expressed as a ratio (surgical/control), were decreased in surgical eyes. The amplitudes for b-wave were: before surgery, 107.6% ± 4.6%; 28 days after surgery, 61% ± 4% (P < 0.001); and 56 days after surgery, 62% ± 5.6% (P < 0.001). Oscillatory potentials were the most dramatically affected: before surgery, 108.6% ± 6.7%; 28 days after surgery, 57.5% ± 5% (P < 0.01); and 56 days after surgery, 57% ± 8.5% (P < 0.001). Amplitudes of the a-waves had relatively smaller but still significant deficits: before surgery, 105.8% ± 6.9%; 28 days after surgery, 72.2% ± 5.4% (P < 0.01); and 56 days after surgery, 79.8% ± 11.0% (P < 0.01). Histologic analysis of the surgical eyes revealed development of anterior synechia, loss of retinal ganglion cells (RGCs), and thinning of all retinal layers. Electron microscopy of optic nerve cross sections revealed swelling and degeneration of the large diameter axons and gliosis.


Glaucoma is a progressive optic neuropathy with characteristic optic disc changes and progressive visual field defects.1 The elevated intraocular pressure (IOP) is considered a primary risk factor for the initiation and progression of glaucomatous neuropathy.2 However, in many patients, despite adequate control of IOP, the loss of vision continues to progress, which dictates the necessity for the further identification of molecular mechanisms responsible for glaucomatous neurodegeneration.3

Despite new discoveries, the genetic and environmental factors causing and modifying glaucoma have not been clearly defined.4 Different molecular mediators have been proposed as potential suspects in glaucomatous neurodegeneration (inducible nitric oxide synthase [iNOS],5–7 TNF-α,8,9 lack of growth factors or their receptors,10 and alteration of optic nerve head matrix molecules11). A disruption of adequate blood supply and energy metabolism12–14 or dysfunctional protein secretion15 have also been suggested to be causes of the selective death of retinal ganglion cells (RGCs) in glaucoma.

A potentially powerful model for the investigation of these factors is the mouse, in which the genome can readily be altered by adding, changing, or deleting genes.16–17 However, the small size of the mouse eye has precluded noninvasive monitoring of IOP and optic nerve function, greatly restricting the use of a mouse model in glaucoma studies.16

A procedure for the reliable induction of chronic ocular hypertension in genetically manipulated mice may provide a unique advantage in the investigation of the link between other neurodegenerative diseases and glaucoma.18–20 Furthermore, the availability of transgenic mice that mimic different human diseases could provide an opportunity to identify possible risk factors that may predispose to and/or are responsible for RGC death due to chronic elevation of IOP. The induction of chronic ocular hypertension in transgenic or knockout mice in which signaling molecules and/or their receptors are altered may offer a powerful tool for the investigation of signal transduction mechanisms involved in the RGC death that occurs in glaucoma.17

We recently adapted and validated noninvasive techniques currently used to study aqueous humor outflow properties (fluorophotometry),21 tonometry, and retinal function22 in small laboratory animals. The goal of this study was to investigate whether it is possible to induce reliably a chronic elevation of IOP in mice, evaluate potential functional deficits of the retina using electroretinography (ERG), and determine whether morphologic changes resemble changes in glaucomatous human eyes.

MATERIALS AND METHODS

Laser Induction of Chronic Ocular Hypertension

All animal studies were conducted in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research, 2000, and with guidelines of the Animal Care and Use Committee of the University of Iowa. The studies were approved by the University of Iowa Animal Care and Use Committee. All animal studies were conducted in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.
and procedures were approved by the Iowa State University Committee on Animal Care. C57/Bl6 mice (6–8 months of age, n = 54) were used in the experiments. Mice were kept in a 10:14-hour light–dark regimen.

Laser treatment of the trabecular meshwork in monkeys and rats produces chronic elevation of IOP.23–24 We hypothesized that this would also be true of the mouse eye. We injected the photosensitive dye (indocyanine green) in the anterior chamber of one eye to increase the local photocoagulative effect of the laser treatment.

Briefly, mice were anesthetized with 2.5% isoflurane+100% oxygen, and body temperature was maintained with a heating pad. Indocyanine green (3 mL, 10 mg/mL; Sigma-Aldrich, St. Louis, MO) was injected into the anterior chamber of the eye at a very slow rate with a syringe (Hamilton, Reno, NV) attached to a microinjector pump, to avoid an abrupt elevation of IOP during the procedure. Animals were pretreated with 4% pilocarpine hydrochloride eye drops, which increased outflow of the dye into the trabecular meshwork and episcleral veins. The pilocarpine treatment also caused miosis, which served to protect the posterior pigmented structures of the eye from the diode laser energy (the pigmented iris served as a barrier for any potential stray energy). Twenty minutes after injection, a diode laser (DioVet; Iridex Corp., Mountain View, CA) was used to deliver 810-nm energy pulses through a 50-μm fiberoptic probe to the region of the trabecular meshwork and episcleral veins in proximity to the limbal region. Careful positioning of the fiberoptic probe insured that the orientation of the laser was away from the pigmented structures of the retina. We demonstrated that the aqueous fluorescein clearance are decreased in monkeys with laser-induced chronic ocular hypertension.27 Thus, we were interested in determining whether that is the case in laser-treated mouse eyes as well.

Fluorophotometric Evaluation of Aqueous Humor Flow

Fluorophotometry was pioneered by Goldmann26 as an objective method to measure aqueous humor flow by measuring the rate of clearance of fluorescein from the anterior chamber. It has been demonstrated that the aqueous flow and fluorescein clearance are decreased in monkeys with laser-induced chronic ocular hypertension.27 Thus, we were interested in determining whether that is the case in laser-treated mouse eyes as well.

Iontophoretic Application of Fluorescein. Fluorescein was applied to the anterior chamber of the mouse eye by iontophoresis, as we have described.21 Briefly, uniform (0.465 g) blocks of 2% agarose gel containing 0.5% fluorescein (AK-Fluor; Akorn, Inc., Decatur, IL) were placed on the cornea. The gels were held in position by a syringe filled with 0.9% saline, containing the negative electrode of an iontophoretic device (Phoresor II, Motion Control Inc., Salt Lake City, UT). The positive electrode of this device was placed on the ocular}

**IOP Monitoring**

IOP was measured with a modified Goldmann tonometer (Haag-Streit USA, Inc., Mason, OH), as previously described (Fig. 2).22,25 Tonometry in mice required a modified applanation prism with a biprism angle of 36° and applied weight of 25 mg per scale division (2 g full scale), as previously described by Cohan and Bohr.25 A calibration of the modified Goldmann tonometer was performed by comparing IOP results measured by invasive manometry and modified tonometry ($y = 1.74 \pm 0.97x, r^2 = 0.89$).22

IOPs were measured in anesthetized mice, under 1% halothane and 30% NO-70% O2 anesthesia. The Goldmann tonometer tip was applied toward the center of the cornea, and 5 to 10 readings were obtained for the calculation of mean pressures in each eye.

**FIGURE 2.** Goldmann tonometer recording procedure in a mouse. A 25-gauge needle connected to a reservoir containing 0.9% NaCl was carefully inserted in the posterior chamber, and IOP was then raised to 30 mm Hg by elevating the reservoir. (A) The Goldmann tonometer scale was set at 50 mm Hg. (B) Reading from the Goldmann tonometer was obtained when fluorescent rings reached a matching position (reading was 50 mm Hg, which corresponded well to the preset IOP value). (C) The Goldmann tonometer scale was set on 20 mm Hg (fluorescent rings do not overlap). Images were obtained with a video camera attached to the slit lamp biomicroscope in which the optical axis was aligned with the optic prism.
region of the anesthetized animal, and a 2.5-mA current applied for 5 minutes. Fluorescein is a negatively charged molecule and can diffuse through the electrical gradient toward the positive electrode. The Phoresor II has the capability of automatically stopping the current and time countdown if corneal contact with the gel is lost, ensuring that a constant amount of fluorescein is delivered during iontophoresis.

Fluorophotometry. Fluorophotometry was performed with a slit lamp biomicroscope equipped with a high integration charge coupled device (Cohu, San Diego, CA) attached to a bioimage analysis system (model 500; Bioimage Co., Chicago, IL) that can detect low-level fluorescence. The images were digitized and the optic density quantified with image-analysis software (Scion Corp., Frederick, MD). Excitation of fluorescein was achieved with a cobalt blue filter. A barrier filter (Wratten filter no. 15; Kodak film tablet; Eastman Kodak Co., Rochester, NY) was used for the selective detection of fluorescence from the anterior chamber. Baseline fluorescence was measured, and system calibration was performed in each animal by using the calibrated gray scale (Kodak film tablet; Eastman Kodak Co.) as a standard before application of fluorescein. Fluorescence was measured at 15-minute intervals for 160 minutes after application of fluorescein. Fluorescein clearance was calculated using the published formula: $C = \frac{(A - B)/(A + B)}{2} \cdot t_{m}$, where $C$ is clearance, $A$ is fluorescence at time point A, $B$ is fluorescence at time point B; and $t_{m}$ is the time interval between points A and B.

Electroretinography

Electroretinography was used to determine physiological amplitudes and latencies of the retinal responses in 22 mice. Animals were dark adapted for at least 6 hours before each experiment. The animals were anesthetized as previously described, and the pupils dilated with 1% tropicamide and 2.5% phenylephrine. Mice were placed in a specially designed dome with an interior that was completely covered with aluminum foil to obtain a Ganzfeld effect. Body temperature was maintained with a microwave-heated thermal pad. A light stimulus was delivered through the ceiling of the dome (PS-22 stimulator; Grass-Telefactor, West Warwick, RI). Measured luminance in all quadrants was $1600 \pm 200$ cd/m$^2$. Two electrodes made of a DTL fiber (Chieldex Trading GmbH, Palmyra, NY) and Ag/AgCl cells were used to simultaneously record electroretinograms from both eyes. The reference electrode was positioned in the ear, and the ground electrode was placed subcutaneously on the back. An evoked potential measuring system (Neuropack-MEB 7102; Nihon-Kohden America, Foothill Ranch, CA) was used to deliver a triggered output to the flash stimulator and collect signals from both eyes. A flash ERG routine was delivered at a 0.2 Hz frequency (10 averaged signals per recording session, sensitivity 100 $\mu V$/division, low-cut frequency 0.5 Hz, high-cut frequency 10 kHz, analysis time 500 ms). Oscillatory potentials (OPs) were recorded by delivering light stimuli at a 0.2 Hz frequency (10 averaged signals per recording session, sensitivity 100 $\mu V$/division, low-cut frequency 50 Hz, high-cut frequency 500 Hz, analysis time 100 ms). Isolated cone responses were recorded from light-adapted eyes by delivering stimuli at 20 Hz (50 averaged signals per recording session, sensitivity 50 $\mu V$/division, low-cut frequency 0.5 Hz, high-cut frequency 10 kHz, analysis time 500 ms). To avoid potential bias due to electrode differences, recordings were repeated with electrodes switched to the opposite eye.

Histologic Examination

Sixty-five days after surgery, mice were deeply anesthetized with a high dose of phenobarbital (100 mg/kg) and perfused intracardially with ice-cold heparinized saline followed by 4% paraformaldehyde in 0.1 M phosphate buffer. Optic nerves (dissected 1 mm posterior to the sclera) were postfixed in a solution containing 4% paraformaldehyde-2% glutaraldehyde in 0.1 M phosphate buffer (pH 7.4) and then rinsed in cacodylate buffer, postfixed in 2% osmium tetroxide in cacodylate buffer, dehydrated in alcohol, and embedded in epoxy resin. Cross sections (1 $\mu m$ thick) were cut with an ultramicrotome, mounted on glass slides, and stained with 1% toluidine blue. Ultrathin sections of the optic nerves were examined by electron microscopy. Eye globes were paraffin embedded, and 7-10 $\mu m$-thick sections of the retina were collected onto poly-l-lysine-coated glass slides, and stained with hematoxylin and eosin. The eye globe tissue sections were examined with a photomicroscope (Microphot FXA; Nikon Corp., New York, NY). Images were captured with a digital camera (Megaplan model 1.4; Eastman Kodak Corp.) connected to a framegrabber (MegaGrabber; Eastman Kodak Corp.) in a computer (Macintosh 8100/80 AV; Apple Computer, Cupertino, CA) using NIH Image 1.58 VDM software (available by ftp at zippy.nimh.nih.gov/ or at http://rsb.info.nih.gov/nih-image; developed by Wayne Rasband, National Institutes of Health, Bethesda, MD).

The morphometric analysis of the retina was performed as we have described. Briefly, sections of the retina at the level of the optic nerve head were prepared, and each section was divided into 12 optic fields (6 fields of the central retina and 6 fields of the peripheral retina). Within each field we measured the thickness of the inner plexiform and RGC layer (IPL+RGC), inner nuclear (INL), and outer nuclear layer (ONL) by measuring each layer in four places and calculating the average thickness per section. Measurements of retinal layer thickness were performed using a calibrated scale of the objective of the light microscope.

Statistical Analysis

Statistical analysis was performed by Student’s t-test, a paired t-test, the Kruskal-Wallis nonparametric test, and correlation analysis performed on computer (GraphPad, San Diego, CA).

RESULTS

Tonometry and Fluorophotometry

IOP was significantly elevated in eyes of 22 (88%) of 25 surgically altered mice after laser-induced surgery compared with control (nonsurgical) eyes (Fig. 3). Preoperative IOPs ($n = 22$) were $15.2 \pm 0.6$ mm Hg (left eyes) and $15.3 \pm 0.6$ mm Hg (right eyes); 12 hours after surgery, $33.9 \pm 4.7$ mm Hg (left eyes-surgical) and $14.5 \pm 1.2$ mm Hg (right eyes-control; $P = 0.006$, paired $t$-test; $n = 6$); 10 days after surgery, $33.6 \pm 1.5$ mm Hg (surgical eyes) and $17 \pm 0.7$ mm Hg (control eyes; $P < 0.001$, paired $t$-test; $n = 22$); and 30 days after surgery, $27.4 \pm 1.2$ mm Hg (surgical eyes) and $15.8 \pm 0.8$ (control; $P < 0.001$, paired $t$-test, $n = 22$). However, 60 days after surgery IOP in the surgical eyes decreased to $19.5 \pm 0.8$ (control; $P < 0.001$, paired $t$-test, $n = 22$).
0.9 mm Hg and was not significantly different than in control eyes (control, 17.3 ± 0.7 mm Hg; P = 0.053, paired t-test, n = 21). To evaluate whether laser treatment induced an immediate elevation of IOP and to rule out the possibility that we had an immediate spike in IOP that might cause absolute retinal ischemia, six mice were operated on, and IOP was evaluated 12 hours after surgery. IOP was significantly elevated in the surgical eyes (33.9 ± 4.7 mm Hg, P = 0.006, paired t-test) when compared with the control eyes (14.5 ± 1.2 mm Hg). Only one animal had IOP above 40 mm Hg (45 mm Hg). Immediate recording of IOP was not possible, because of the aqueous humor leak from the needle track that was created during the injection of indocyanine green dye.

An analysis of the fluorophotometry data (Fig. 4) revealed significantly decreased fluorescein clearance in surgical eyes at 15 (P = 0.02, Student’s t-test) and 30 (P = 0.01, Student’s t-test) days after surgery.

**Electroretinography**

Electroretinography was used as an objective method to evaluate the functional status of the inner and outer retina. To evaluate the effect of the chronic elevation of IOP on different populations of photoreceptors, we used full-field scotopic flash ERG to evaluate combined rod and cone activity and photopic flicker ERG (fERG) for the isolated cone activity.

ERG amplitudes, expressed as a ratio (surgical/control), were decreased in surgical eyes, and retinal inner function was predominantly affected (Fig. 5).

The amplitudes of a-waves (Fig. 6A) had relatively smaller but still significant deficits compared with b-waves and OPs: control, 105.8% ± 6.9%; 28 days after surgery, 72.2% ± 5.4% (P < 0.01, Kruskal-Wallis nonparametric test with the Dunn post test), and 56 days after surgery, 79.8% ± 11% (P < 0.01, Kruskal-Wallis nonparametric test with the Dunn post test). fERG analysis (Fig. 6B) revealed a mild but nonsignificant decrease of the function in surgical eyes: control, 99% ± 9%; 28 days after surgery, 78.5% ± 13.3% (P > 0.1, Kruskal-Wallis nonparametric test with the Dunn post test), and 56 days after surgery, 88.3% ± 13.3% (P > 0.1, Kruskal-Wallis nonparametric test with the Dunn post test). However, when absolute amplitudes were compared between surgical and control (nonsurgical) eyes there was a significant decrease in surgical eyes at 28 (P = 0.001, paired t-test) but not at 56 (P = 0.5, paired t-test) days after surgery. The amplitudes for b-wave (Fig. 6C) were control, 107.6% ± 4.6%, and 28 days after surgery, 61% ± 4% (P < 0.001, Kruskal-Wallis nonparametric test with the Dunn post test; n = 22); 56 days after surgery, 62% ± 5.6% (P < 0.001, Kruskal-Wallis nonparametric test with the Dunn post test). OPs were the most dramatically affected (Fig. 6D): control, 108.6% ± 6.7%; 28 days after surgery, 57.5% ± 5.5% (P < 0.01, Kruskal-Wallis nonparametric test with the Dunn post test); and 56 days after surgery, 57% ± 8.5% (P < 0.001, Kruskal-Wallis nonparametric test with the Dunn post test).

The latency of fERG responses, b-waves, and OPs was significantly increased in surgical eyes 28 days after surgery (Fig. 7): a-wave latency, control, 22.2 ± 0.7 ms, and latency_surgical, 23 ± 0.7 ms (P = 0.06, paired t-test); b-wave latency, control, 19.6 ± 0.4, and latency_surgical, 22 ± 0.9 ms (P = 0.03, paired t-test); OPs latency_control, 25.2 ± 0.9 ms, and latency_surgical, 26.5 ± 0.8 ms (P = 0.04, paired t-test); and fERG latency control, 50.6 ± 1.5 ms, and latency_surgical, 53.9 ± 1.5 ms (P = 0.03, paired t-test).

However, when latencies were analyzed 56 days after surgery, only b-wave latencies of surgical eyes were significantly increased compared with control (nonsurgical) eyes: a-wave

**Figure 4.** Fluorophotometry analysis revealed decreased fluorescein clearance in surgical eyes compared with healthy, nonsurgical control eyes. Data are the mean ± SEM.

**Figure 5.** Original ERG traces from one of the surgically treated mice 56 days after surgery. (A) Scotopic flash ERG. (B) Scotopic OPs. (C) Photopic fERG. Laser-induced surgery was characterized by relatively nonaffected a-wave amplitudes (scotopic flash ERG) and photopic flicker ERG amplitudes; however, oscillatory potentials had dramatic deficits, which imply primary damage to the inner retina.
latency_{control} \pm \text{SD} = 23.5 \pm 0.9 \text{ ms}, \text{ and } latency_{surgical} \pm \text{SD} = 24.2 \pm 1 \text{ ms} (P = 0.5, \text{ paired } t\text{-test}); b\text{-wave latency}_{control} \pm \text{SD} = 15.7 \pm 0.9, \text{ and } latency_{surgical} \pm \text{SD} = 20 \pm 1.7 \text{ ms} (P = 0.02, \text{ paired } t\text{-test}); OPs latency_{control} \pm \text{SD} = 28.2 \pm 0.7 \text{ ms}, \text{ and } latency_{surgical} \pm \text{SD} = 29.4 \pm 1.3 \text{ ms} (P = 0.3, \text{ paired } t\text{-test}); \text{ and } \text{flERG latency}_{control} \pm \text{SD} = 49.7 \pm 1.4 \text{ ms}, \text{ and } latency_{surgical} \pm \text{SD} = 51.4 \pm 1.9 \text{ ms} (P = 0.14, \text{ paired } t\text{-test}).

Because we were concerned that laser energy by itself or injection of the indocyanine green might cause retinal damage,

Figure 6. ERG amplitudes represented as a ratio (surgical/control). (A) Scotopic flash ERG a-wave, (B) photopic flicker ERG, (C) scotopic flash ERG b-wave, (D) scotopic OPs.

Figure 7. Laser cauterization of ICG-filled episcleral veins caused almost complete obstruction of the venous lumen. (A) Control. Arrows: borderlines of an episcleral vein that is positioned in proximity to the conjunctiva of the limbal region. (B) Treated eye. Open arrows: borderlines of an episcleral vein. A fibrous proliferation was evident that almost completely obstructed the venous lumen. C, conjunctival epithelium; CB, ciliary body; S, sclera; EV, episcleral vein lumen. Bar, 50 µm.
we analyzed ERG amplitudes (a- and b-waves) from mice that received indocyanine green injection but not laser treatment \((n = 5)\) and mice that received laser treatment only \((n = 4)\). An indocyanine green injection did not cause decrease of a-wave \((P = 0.6, \text{paired } t\text{-test})\) or b-wave \((P = 0.2, \text{paired } t\text{-test})\) amplitudes 3 weeks after surgery. Mice that received only laser treatment did not show ERG deficits in surgical eyes compared with control (nonsurgical) eyes 3 weeks after surgery (a-wave amplitudes, \(P = 0.34\); b-wave amplitudes, \(P = 0.16\), paired \(t\text{-test})\).

**Correlation Analysis of Tonometry and ERG Data**

Detailed analysis of tonometry and electroretinography data revealed significant correlation between ocular hypertension expressed as a coefficient of IOP elevation and ERG amplitudes: \((\text{IOP}_{\text{surgical eyes}} - \text{IOP}_{\text{control eyes}}) \times \text{duration of hypertension (mm Hg } \times \text{days) versus ERG amplitudes expressed as a ratio (surgical/control, %).}

Correlation analysis data were IOP versus a-wave \((r^2 = 0.2; P = 0.0002)\), IOP versus b-wave \((r^2 = 0.4; P < 0.0001)\), and IOP versus OPs \((r^2 = 0.2; P = 0.0004)\). Analysis did not reveal significant correlation between IOP and fERG amplitudes \((r^2 = 0.02; P = 0.18)\).

**Histologic Analysis**

Our electrophysiological studies revealed a reduction of the function in all retinal layers. Histologic analysis was performed to determine whether functional defects were in agreement with the morphologic appearance of the surgical eyes.

Light microscopy analysis revealed an obstruction of the episcleral veins (Fig. 7), trabecular meshwork destruction with partial or complete obstruction of Schlemm’s canal (Fig. 8), and development of anterior synchia in nine surgical eyes (Fig. 10A). The thickness of all retinal layers appeared decreased in surgical eyes compared with the control (Fig. 10E).

Electron microscopy analysis of the optic nerves (Fig. 12) revealed loss and degeneration (electron-dense appearance, Fig. 12B) of the large diameter axons, swelling of myelin sheaths, condensation and degeneration of the cytoskeleton (Fig. 12C), and activation of the glial cells in proximity to degenerating axons (Fig. 12C–F).

For the analysis, we compared measurement of the thickness of the IPL and RGC layer (IPL+RGC), INL, and ONL, and total retinal thickness between control and surgical eyes (Table 1). Morphometric analysis revealed significant reduction of the total retinal thickness. However, a separate analysis of the ONL, INL, and IPL+RGC layers in surgical eyes did not reveal statistically significant thinning compared with control (nonsurgical) eyes.

**Correlation Analysis of Histology and ERG Data**

Linear regression analysis revealed poor correlation between OP amplitudes and combined retinal thickness of the RGC and IPL layers, expressed as a ratio between surgical and control eyes \((r^2 = 0.09, P = 0.6)\). Calculated correlation coefficients did not reveal significant correlations between morphologic and functional analysis for other retinal layers: a-wave ratio versus ONL thickness \((r^2 = 0.15, P = 0.09)\), b-wave ratio versus INL thickness \((r^2 = 0.34, P < 0.05)\), a-wave amplitudes versus ONL thickness \((r^2 = 0.34, P < 0.05)\), and b-wave amplitudes versus INL thickness \((r^2 = 0.34, P < 0.05)\).

**Figure 8.** (A) Histology of the anterior segment in a control eye. Arrow: epithelial lining of the trabecular meshwork; (•) Schlemm’s canal lumen. (B) Histology of the anterior segment of the surgical eye (60 days after surgery). Partial obstruction of Schlemm’s canal was detectable (•), and the trabecular meshwork structure was completely destroyed and obstructed (arrow). Outer segments and retinal pigment epithelium of the peripheral retina did not have signs of damage induced by direct application of the diode laser. AC, anterior chamber; CB, ciliary body; I, iris; ICA, iridocorneal angle; RET, peripheral retina. Bar, 50 μm.

**Figure 9.** Latency analysis for different ERG components. a, a-wave latency; b, b-wave latency; OPs, oscillatory potentials latency; fl, flicker ERG latency. *\(P < 0.05\).
versus INL thickness ($r^2 = 0.13, P = 0.1$) and b-wave ratio versus IPL+RGC layer thickness ($r^2 = 0.03, P = 0.42$).

**DISCUSSION**

The experimental approach used in this study allowed the precise monitoring of the dynamics of retinal damage during chronic elevation of IOP. Laser cauterization of the indocyanine green–pretreated eyes was a successful procedure for the induction of chronic elevation of IOP in mouse eyes. Laser-induced cauterization caused rapid elevation of IOP during the first 12 hours of the postoperative period (34 mm Hg), which was sustained for at least 4 weeks.

Analysis of the ERG parameters (amplitudes and latency of scotopic flash ERG, OPs, and fERG) was an effective strategy for the functional monitoring of different cell populations in the retina. Previous reports have indicated that recording the OPs may be a relatively sensitive method to detect retinal damage in glaucomatous eyes. Our results show that, indeed, OPs were the most dramatically affected in chronically hypertensive mouse eyes. Chronic hypertension in the mice also affected a- and b-wave components of the scotopic flash ERG. Similar type of changes have been reported in a study that observed ERG changes in DBA/NNi mice, which undergo development of spontaneous glaucoma. Detailed analysis of the pattern ERG responses in DBA/2NNi mice revealed development of significant functional deficits as early as 5 months of age. An ERG analysis of scotopic flash ERG amplitudes revealed development of significant a-wave deficits by 7 months of age, whereas b-wave deficits were detected at 8 months of age. It has been reported that DBA mice have the maximum elevation of IOP at the age of 9 months (20.3 mm Hg), which corresponds to the development of significant ERG deficits, and thus it was not unexpected to observe similar ERG deficits in our model, because elevation of IOP was present immediately after surgery in ranges that were much higher than in DBA mice. Because mice do not have a developed lamina cribrosa, there is a possibility that damage induced by elevation of IOP is distributed not only to the optic nerve but also to the different retinal layers. Functional analysis of the sham-operated mice that received only laser treatment or indocyanine green injection did not reveal functional deficits com-

**FIGURE 10.** Light microscopy of surgical (A, C, E) and control (B, D, F) eyes. Surgical eyes were characterized by development of trabecular meshwork destruction with partial or complete obstruction of Schlemm’s canal, whereas a minority of eyes ($n = 9$) had detectable development of the anterior synechia. Optic nerve head distortion, without evidence of prominent cupping and decreased thickness of the retina with a loss of RGCs were major features of surgical eyes. Laser-induced damage to the peripheral retina (uveitis, retinal detachment, RPE, and photoreceptor destruction) was not detected by light microscopy. AC, anterior chamber; C, cornea; CB, ciliary body; I, iris; OPL, outer plexiform layer; OS, outer segments. Arrow: corneal edema.
pared with the control eyes, which rules out the possibility that invasive manipulation of the eyes and/or laser energy are responsible for the detected functional deficits. Furthermore, detailed histologic analysis of peripheral retinas in surgical eyes, which received combined treatment (indocyanine green injection and laser application) did not reveal uveitis, retinal detachment, destruction of the retinal pigment epithelium, or photoreceptors in proximity to the trabecular meshwork, where the effect of direct laser energy should be the most extensive.

**Figure 11.** Light microscopy analysis of optic nerves revealed different levels of damage. (A) Control; (B–D) surgical eyes. Massive degeneration of the optic nerves was evident. Bar, 100 μm.

**Figure 12.** Electron microscopy analysis of optic nerves of the control (A) and surgical (B–F) eyes. Degeneration of large diameter axons and gliosis were the most predominant features in the optic nerves of the surgical eyes. (E) Gliosis of the optic nerve in the surgical eye. Scale bars: (A, C, E) 500 nm; (B, D, F) 1 μm.
Histologic and electron microscopy analysis data revealed damage of the optic nerve and thinning of all retina layers. Detailed examination of the anterior chambers in surgical eyes revealed complete destruction of the trabecular meshwork, complete or partial obstruction of Schlemm’s canal, and development of anterior synechia that were probably responsible for the decreased aqueous humor outflow and elevated IOP that were the final result. However, we cannot exclude the possibility that photocoagulation of episcleral veins did not participate at least partially in the development of chronic ocular hypertension.

DBA/2 inbred mouse strains have been found to exhibit spontaneous development of a form of a secondary glaucoma phenotype: pigment-dispersion glaucoma. The exact genetic defect responsible for the DBA/2J (D2) phenotype was recently identified. Using mapping techniques, sequencing, and functional genetic tests, Anderson et al. showed that iris pigment dispersion (IPD) and iris stromal atrophy (ISA) result from mutations in related genes encoding melanosomal proteins (Gpumnb and TYRP1). They hypothesized that IPD and ISA alter melanosomes, allowing toxic intermediates of pigment production to leak from melanosomes, causing iris disease and subsequent pigmentary glaucoma. However, a recent study did not identify mutation of the TYRP1 gene in humans with pigmentary glaucoma.

Although models of spontaneous development of glaucoma have considerable value for glaucoma research, inducible models have the unique advantage of providing control parameters from the same animal (one eye surgically treated and the fellow eye as the control) with the dramatically increased speed of data generation from different transgenic lines.

Table 1. Morphometric Analysis of the Retina

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<th>IPL + RGC</th>
<th>INL</th>
<th>ONL</th>
<th>Total Thickness</th>
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<td>Control</td>
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<td>25.5 ± 0.9</td>
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<td>23.9 ± 1</td>
<td>45.3 ± 1.2</td>
<td>103.7 ± 3.5*</td>
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Data are mean micrometers ± SEM.

*p < 0.05; paired t-test.

A mouse model of laser-induced chronic ocular hypertension for studying glaucoma may have the potential to become a powerful experimental tool in the development of neuroprotective drugs and molecular determination of the pathogenesis of glaucoma.

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References


