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Global Changes in Optic Nerve Head Gene Expression after Exposure to Elevated Intraocular Pressure in a Rat Glaucoma Model

Elaine C. Johnson, Lijun Jia, William O. Cepurna, Thomas A. Doser, and John C. Morrison From the Kenneth C. Swan Ocular Neurobiology Laboratory, Casey Eye Institute, Department of Ophthalmology, Oregon Health Sciences University, Portland, Oregon.

Abstract

Purpose— In glaucoma, the optic nerve head (ONH) is the likely site of initial injury and elevated intraocular pressure (IOP) is the best-known risk factor. This study determines global gene expression changes in the pressure-injured ONH.

Methods— Unilateral sustained IOP elevation (glaucoma, n = 46) or optic nerve transection (n = 10) was produced in rats. ONHs were removed, and the retrobulbar optic nerves were graded for degeneration. Gene expression in the glaucomatous ONH with extensive injury was compared with that in the fellow ONH (n = 6/group), by using cDNA microarrays. Data from 12 arrays were normalized, significant differences in gene expression determined, and significantly affected gene classes identified. For the remaining ONH, grouped by experimental condition and degree of injury, quantitative reverse transcriptase-PCR (qPCR) and ANOVA were used to compare selected message levels.

Results— Microarray analysis identified more than 2000 significantly regulated genes. For 225 of these genes, the changes were greater than twofold. The most significantly affected gene classes were cell proliferation, immune response, lysosome, cytoskeleton, extracellular matrix, and ribosome. A 2.7-fold increase in ONH cellularity confirmed glaucoma model cell proliferation. By qPCR, increases in levels of periostin, collagen VI, and transforming growth factor β 1 were linearly correlated to the degree of IOP-induced injury. For cyclinD1, fibulin 2, tenascin C, TIMP1, and aquaporin-4, correlations were significantly nonlinear, displaying maximum change with focal injury.

Conclusions— In the ONH, pressure-induced injury results in cell proliferation and dramatically altered gene expression. For specific genes, expression levels were most altered by focal injury, suggesting that further array studies may identify initial, and potentially injurious, altered processes.

Glaucoma is the second leading cause of blindness worldwide, affecting approximately 60 million people.¹ Although many risk factors are associated with glaucoma, IOP is the most widely recognized, and lowering IOP is the goal of current glaucoma therapies. When IOP is experimentally elevated in nonhuman primates, the pattern of optic nerve head (ONH) cupping, optic nerve axon degeneration, and selective loss of retinal ganglion cells replicates the pathologic features of human glaucoma. Clinically, human glaucomatous optic neuropathy is characterized by optic disc cupping and a pattern of visual field loss.² Cupping results from

Corresponding author: Elaine C. Johnson, Casey Eye Institute, 3375 S.W. Terwilliger Boulevard, Portland, OR 97201; johnsoel@ohsu.edu.

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the loss of optic nerve axons and posterior bowing and remodeling of the support structures of the ONH.^{3,4} Often, these changes are most pronounced in the superior and inferior parts of the nerve head. The most characteristic visual field defect is the arcuate scotoma, which arches above or below central fixation and follows the pathways of the nerve fiber bundles as they converge on the superior and inferior poles of the ONH.^{2,5,6}

Regional variation in laminar structure suggests less support and protection for axons, possibly explaining the apparent increased susceptibility of axons that pass through the superior and inferior ONH. In the face of increased or fluctuating IOP, movement of the lamina may result in preferential mechanical injury to neural tissues in these regions.⁷ In addition, because the vascular supply to the ONH tissue lies within the laminar beams, this pattern could result in regionally compromised blood flow.⁸ Current experimental evidence supporting either of these mechanisms is limited, and the cellular events that connect the regional pattern of glaucomatous optic nerve damage with the known structural anatomy of the ONH are still largely unknown. Regardless of mechanism, the variation in the structure of the ONH provides the only anatomic correlation with the characteristic pattern of glaucomatous optic nerve axon loss.

The apparent vulnerability of the ONH to pressure-induced axonal injury has led many investigators to examine changes in the composition of glaucomatous human and experimental monkey ONH tissues. These ONHs are characterized by axon loss, gliotic scaring, increased expression of matrix metalloproteinases, and abnormal deposition of extracellular matrix (ECM) materials, including collagens, tropoelastin, tenascin, and proteoglycans.^{9–20}

In the rat, glaucoma can be modeled by sclerosing aqueous outflow pathways to produce a sustained elevation of IOP.²¹ In this model, pressure causes a selective loss of retinal ganglion cells and a characteristic pattern of axon degeneration that begins in the superior quadrant of the optic nerve.^{21,22} In addition, immunohistochemical studies of ONH reveal that pressure-induced injury is accompanied by deposition of collagens and other ECM components, similar to that in human glaucoma.²³ This deposition of ECM is preceded by a loss of gap junctional connexin 43 immunolabeling and evidence of astrocytic proliferation.²⁴ By immunohistochemical analysis, the initial evidence of ECM deposition coincided with a decrease in ONH labeling for neurotrophins and astrocytic GFAP.²⁴

Although the ONH is recognized as the likely site of initial injury in human glaucoma and in experimental IOP elevation glaucoma models, very little is known about the changes in gene expression that accompany this injury. In this study, we used microarray analysis to identify the genes and functional gene classes most altered in expression in the rat ONH after exposure to experimentally elevated pressure that results in extensive and ongoing optic nerve degeneration. Then, we used real-time quantitative (q)PCR to verify selected changes in gene expression initially identified by microarray analysis and to examine several genes not included on the arrays. To expand this study, we included qPCR analyses of ONHs from eyes with focal regions of degeneration in optic nerve cross-sections. Focal injury occurs in eyes with more mild pressure elevations or those of shorter duration and probably reflects earlier responses to pressure-induced nerve injury. We compared injury in these ONHs to those in ONHs with more extensive injury due to elevated IOP and to those after optic nerve transection, to evaluate ONH responses to simple loss of axons. The expansion of the qPCR study to nerves with focal injuries allows us to explore the potential of discovering, by future microarray analyses, unique or more dramatic alterations in gene expression that occur early in the injury process. To our knowledge, this is the first genome-wide analysis of expression changes in the ONH in response to elevated IOP.

Methods

Glaucoma Model

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All animal experiments were performed in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research and were approved by the Oregon Health Sciences University (OHSU) Animal Care and Use Committee. Sustained IOP elevation was produced in Brown Norway rats by unilateral episcleral vein injection of hypertonic saline. IOP was measured 4 days per week with a handheld tonometer (Tono-PenXL; Medtronic, Minneapolis, MN) in awake animals housed in low-level, constant light to stabilize the circadian IOP oscillations that maximize during the dark phase.^{25,26} At 5 weeks after injection, the eyes were removed and rapidly chilled in cold phosphate-buffered saline, and the retinas were removed. Using a trephine, the ONH was removed from the posterior pole of the globe. The dissected ONH was then cut to 1 mm in length, a segment that included the optic disc, all the unmyelinated, anterior ONH, and the initial (approximately 0.5 mm) of the myelinated optic nerve, as illustrated in Figure 1A. The ONH was frozen on dry ice and stored at -80°C before extraction. RNA was isolated (Pico-Pure RNA Isolation Kit; Arcturus Bioscience, Inc., Mountain View, CA) by sonicating the frozen nerve heads in the kit extraction buffer using an MS 0.5 probe (UP50H Ultrasonic Processor; Hielscher USA, Inc., Ringwood, NJ) and then, after the instructions provided by the Arcturus kit, including DNase treatment to remove any DNA. Purified RNA was quantified (RiboGreen RNA Quantitation Kit; Invitrogen, Carlsbad, CA).

All corresponding optic nerves were postfixed in glutaraldehyde, embedded in plastic, crosssectioned, and evaluated by light microscopy. Optic nerve cross-sections were graded from 1 (no injury) to 5 (degeneration involving the total nerve area) by five masked observers.²⁵ Our grading scale was designed to be sensitive to early pressure-induced axonal degeneration. By correlating counts of intact axons on transmission electron micrographs to the scale, we have determined that each grade unit corresponds to an approximate 12% increase in the number of degenerating axons.²² At grade 5, at least 50% of the axons are degenerating, whereas many other axons with intact, apparently normal, axoplasm remain.

Microarray Experimental Design and Analysis

Microarray analysis was used to compare gene expression in untreated, fellow ONHs to that in ONHs with extensive optic nerve injury due to elevated IOP. For these studies, the fellow eyes (n = 6) had normal optic nerves confirmed by optic nerve grading (Fig. 1B),²⁵ whereas the glaucoma model ONHs (n = 6) were isolated from eyes with optic nerve injury grades just below 5 (4.64–4.95; Fig. 1C). Data from this group reflect extensive and ongoing elevated IOP-induced nerve injury affecting approximately 50% of the axons. We anticipated that the greatest number of genes and cellular processes within the nerve head would be affected by this injury paradigm. ONH from eyes with more advanced optic nerve gliotic scar formation, indicated by a relative reduction of axonal and myelin debris compared with the proportion of glial cell processes and extracellular matrix (Fig. 1D) were not included in the microarray study.

For each of the 12 ONHs, an aliquot of 50 ng total RNA underwent two rounds of linear amplification, was examined for integrity (Bio-Analyzer; Agilent, Palo Alto, CA), reverse transcribed and dye labeled at the OHSU Spotted Microarray Core Facility http:// www.ohsu.edu/gmsr/smc/index.html). The facility also prepared and processed the cDNA arrays and compiled the data. For each sample, two cDNA arrays, SMCmou8400A and SMCmou6600A, were used with a combined total of 15,400 mouse cDNAs probes, representing the entire initial release of the NIA (National Institute on Aging) mouse library (http://lgsun.grc.nia.nih.gov/cDNA/15k.html) and approximately half the genome. Each probe

was represented by duplicate spots on two duplicate slides, yielding four technical replicates per probe. As illustrated in the reference standard experimental design (Fig. 1E), mRNA expression patterns from each of six fellow and six grade-5 ONHs were compared separately to an ONH RNA reference standard. The standard was derived by pooling aliquots from all samples. Therefore, data represent two groups of six independent biological replicates and a total of 24 array comparisons, 12 each of the SMCmou8400A and SMCmou6600A arrays. Gene expression data are reported as the ratios of the average signal intensity (spot intensity minus local background) for each sample relative to the reference standard and were normalized by using a modified Lowess procedure.²⁷ For each of the two array sets, Significance Analysis of Microarrays, (SAM, version 2.5, www-stat.stanford. edu/~tibs/SAM/ provided in the public domain by Stanford University, Stanford, CA) was used to determine the significance of changes in gene expression due to elevated pressure injury. The parameters for SAM were set at 1000 permutations in the two-class, unpaired format, using the nearest number imputer of 10, a false-discovery rate of 2%, and a minimum change of 1.3-fold, to limit the called genes to those with potential biological significance.

Functional Classes of Significantly Altered Genes

DAVID gene ontology analysis tools (http://apps1.niaid.nih.gov/david/provided in the public domain by the National Institute of Allergies and Infectious Disease, Bethesda, MD) were used to identify and determine the number of altered genes in each significantly affected class for the categories of biological process, cellular component, and molecular function, as defined by the controlled vocabulary of the Gene Ontology consortium (http:// www.geneontology.org/). AMIGO (http://www.godatabase.org/cgi-bin/amigo/go.cgi) was used to construct gene ontology hierarchies for the classes of affected ONH genes. In addition, GenMAPP was used to visualize gene expression changes in affected KEGG pathways (http:// www.genmapp.org/).

ONH mRNA Quantitation by qPCR

Sustained IOP elevation was produced in an additional 40 Brown Norway rats, 21,22 and ONHs were collected 5 weeks later, as has been described. Optic nerve transection was produced in 10 more rats to provide a comparison group of ONHs with injury due solely to retrograde axonal degeneration.²⁸ Nerves were transected approximately 2 mm behind the globe and the ONHs collected 14 days later. ONHs from eyes exposed to elevated IOP with optic nerve injury grades between 1.5 and 4.5 demonstrated focal lesions of increasing size, from less than 15% to ~50% of axons degenerating.²² In nerves with injury grades of 5, the lesion filled the entire optic nerve cross-section and 50% or more of the axons were degenerating. In some grade-5 lesions, glial scar formation was evident. For the transection group, the average grade was 4.9.

For reverse transcription of total RNA, 30 ng of ONH mRNA from each sample, along with a pooled ONH RNA standard curve (2–480 ng) was reverse transcribed.²⁹ The cDNA was amplified using a thermocycler (LightCycler, LightCycler Software 3.5, and DNA Master SYBR Green 1 kit; Roche, Indianapolis, IN) according to the manufacturer's protocol, with a final concentration of 4 mM MgCl₂ and 0.25 μ M each of the relevant primers. The primers used for each cDNA were designed (Primer Designer 3 software for Windows; Sci-ed Software, State Line, PA) and sequences for rat mRNAs were obtained from the NIH National Center for Biotechnology Information (Bethesda, MD) databases (http://www.ncbi.nlm.nih.gov/entrez). Glyceraldehyde phosphate dehydrogenase (*GAPDH*) was used as the housekeeping gene for normalization, because its level was not significantly changed in the microarray study (*P* = 0.8) and it showed no significant correlation to either IOP or optic nerve injury grade by qPCR. *GAPDH* was measured by triplicate assay in each experimental sample and the mean value used for normalization. Table 1 lists the primers used and PCR product sizes. PCR was followed by the generation of melting curves for the amplified products to verify amplification

qPCR Data Analysis

All data are expressed as a percentage of the mean (±SEM) value obtained for uninjected, fellow-eye ONHs. Statistical analyses were performed using commercial software (Excel; Microsoft, Redmond, WA; Prism; GraphPad, San Diego, CA) statistical software packages. Elevated IOP and transection groups were compared with the fellow-eye group by using ANOVA with the Dunnett multiple comparison post test. In addition, linear and nonlinear regression analyses were used to correlate mRNA expression levels to nerve injury grade and to compare regression fits by F test. Analysis of data based on IOP history, rather than optic nerve injury grade, produced equivalent results.

ONH DNA Measurement

Total ONH cell count was determined by extracting and measuring total DNA from seven fellow and five grade-5 ONH (Picopure DNA Extraction kit; Molecular Devices, Sunnyvale, CA; and Quant-iT PicoGreen dsDNA Assay kit). Cell count was determined by dividing total ONH DNA by the DNA content per rat cell (Gregory TR (2005). Animal Genome Size Database. http://www.genomesize.com/). Significance of difference was determined by Student's *t*-test.

Immunohistochemistry

For immunohistochemistry, eight rats with unilateral IOP elevation were transcardially perfused with 4% phosphate-buffered paraformaldehyde and embedded in paraffin. Longitudinal sections through the ONH were prepared, and optic nerve cross-sections graded for degeneration.^{24,25} Immunohistochemistry was performed using the avidin-biotin technique with diaminobenzidine (DAB) chromogen with rabbit IBA-1 antibodies (Wako Chemicals USA, Richmond, VA) at 0.1 μ g/mL with purified rabbit IgG serving as a negative control.²⁴

Results

Significant Changes in Gene Expression by SAM Analysis

Using SAM analysis with a false-discovery rate of 2% and a difference of 1.3-fold, 1539 upregulated and 867 downregulated significantly altered genes were identified. All these genes have a q value (probability adjusted for multiple genes) of <0.02. Of these, 177 genes were upregulated more than twofold, 32 of which were increased greater than threefold (Table 2). Predominant among the most upregulated genes were those associated with mitotic cell division, cell adhesion, ECM, and immune responses. Forty-eight genes were downregulated by 50% or more. The most affected of these are listed in Table 3. These genes were primarily associated with synaptic and axonal mRNA, 30-32 lipid and steroid metabolism, and cell signaling.

Functional Classes of Significantly Altered Genes

The DAVID gene ontology analysis tool (http://apps1.niaid.nih.gov/david) was used to identify significantly affected gene classes using the controlled vocabulary of the Gene Ontology Consortium (http://www.geneontology.org/) from the SAM analysis lists of significantly regulated genes. This tool gives a probability (EASE score) for the significance of gene expression changes for each gene class. The significantly affected gene classes are listed for all regulated genes (Table 4), all upregulated genes (Table 5), and all downregulated genes (Table 6).

Elevated IOP Results in Significant ONH Cellular Proliferation

DAVID analysis of significantly regulated genes was dominated by altered expression of cell cycle and cell proliferation-associated messages (Table 4), implying that cellular proliferation occurs in the pressure-injured ONH. DAVID identified 113 proliferation-associated genes with altered expression. Eighty-seven of these genes were significantly upregulated (Table 5) including the neural progenitor cell marker, nestin (1.8-fold).³³ Mapping the genes to a KEGG cell cycle map revealed that every stage of cell division was affected (not shown). Among upregulated cell cycle genes was cyclin D1. Cyclin D regulates the initiation of the cell cycle. We used qPCR, to confirm the upregulation of cyclin D1 and extend it by examining its expression levels in ONH from eyes with increasing grades of optic nerve injury and in ONH after optic nerve transection (Fig. 2A). We found that the correlation between cyclin D1 and optic nerve injury due to elevated IOP was significantly nonlinear (F test for comparison of linear and third-order polynomial fit, P < 0.0001) with the highest levels of cyclin D1 mRNA occurring in focally injured ONH. These ONHs were from eyes with approximately 5-mm Hg mean pressure elevation and nerve injury grades between 1.5 and 3.5 (Fig. 2B). This, along with other evidence, ^{24,34} suggested that ONH cellular proliferation accompanies early pressure-induced nerve injury. Because cell cycle genes also can be activated during apoptosis, 35,36 we used DNA content to determine the number of cells in additional fellow and grade-5 ONHs. In fellow eye rat ONHs, we found an average content of $4,894 \pm 196$ cells, whereas the number of cells in grade-5 ONHs had increased to $13,266 \pm 2,200$ cells (271%, $P \le 0.000002$, Fig. 2C).

Effect of Elevated IOP on ONH Expression of Immune Response Genes and Evidence of Microglial Activation

Genes associated with the biological process of immune response were also significantly regulated (Table 4). Of 38 significantly regulated genes in this category, 34 of these were upregulated in expression. Predominant in this class were all five components of the complement 1 complex (range, 2- to 4.4-fold), FC receptor (3.4-fold), beta 2 microglobulin (2.5-fold), and several histocompatibility class II antigens (1.8–2.3 fold), all genes expressed by activated microglia in the nervous system. $^{37-42}$ Complement receptor related protein, Crry, was also significantly upregulated (1.55-fold). Crry inhibits the activation of the third component of complement (CR3), protects from complement-mediated autoimmune damage, and is abundant in microglia.^{43,44} Many of these messages are associated with the activation of microglia, and markers for activated microglia have been identified in the pressure-injured ONH.^{45,46} In activated microglia, the upregulation of ionized calcium-binding adaptor molecule 1 (IBA1), also known as allograft inflammatory factor 1, is associated with the formation of lamellipodia and membrane ruffles.⁴⁷ Using qPCR to measure IBA1 mRNA levels in the ONH described in Figures 2A and 2B, we found that IBA1 mRNA levels in the ONH demonstrated a positive linear correlation with the degree of optic nerve injury after 5 weeks of elevated IOP ($R^2 = 0.44$, P < 0.0001) By group, IBA1 mRNA levels were significantly elevated in ONH with injury grades greater than 3.5 (Fig. 3A). Immunostaining for IBA1 in elevated-IOP-injured nerve heads suggested that increases in IBA1 mRNA levels reflected endogenous microglia activation, without ruling out a contribution due to proliferation (Fig. 3B).

Upregulation of ONH Ribosomal Gene Classes

All three gene ontology components of DAVID analysis (Table 4) identified significant effects on ribosomal gene expression in ONH injured by pressure. For genes associated with the biological process of ribosome biogenesis and assembly, 21 of 23 significantly affected genes were upregulated (1.3- to 1.6-fold). For ribosomes and associated components (ribosonucleoprotein complex), 57 of 66 significantly regulated genes were upregulated (1.3-

to 3.5-fold). Finally, for genes that function as structural constituents of ribosomes, 52 of 56 genes were upregulated (1.3- to 3.5-fold).

Dramatic Upregulation of Lysosomal Genes

The lysosome was both the most significantly affected cell component (Table 4) and the most significantly upregulated cell component (Table 5). Among upregulated lysosomal genes were cathepsin-D, -B, -Z, -L, -8, -C, and -H (2.2- to 1.4-fold); lysosomal-associated protein transmembrane 5, a protein activated in retinal microglia in response to optic nerve transection⁴⁸ (2.4-fold); and ganglioside degrading enzymes: beta galactosidase and beta hexosaminidase-A and -B (2.2-, 2.2- and 2.0-fold, respectively).

ECM Gene Expression

Elevated IOP resulted in the upregulation of 28 ECM components including periostin (12.8-fold); fibulin 2 (3.2-fold); tenascin C (2.7-fold); isoforms of collagens I, III, IV, V, and VI (1.5-to 2.5-fold); fibronectin 1 (2.3-fold); cartilage-associated protein (2.0-fold); and decorin (1.8-fold) as well as the matrix proteases ADAMTS-1 (2.9-fold) and matrix metalloproteinase-7 (MMP7, 2.8-fold) and -2 (MMP2, 2.1-fold). In addition, increased processing of ECM and other exported proteins was suggested by the upregulation of 33 genes associated with the Golgi complex.

To confirm some of these changes, we used qPCR to measure the expression levels of ECM messages in the ONH samples from Figures 2A and 2B. Because our previous study of altered gene expression in retinas exposed to elevated IOP had shown that the gene most altered in expression was an MMP inhibitor, tissue inhibitor of metalloproteinase (TIMP)-1 (46-fold increase⁴⁹), we examined ONH TIMP-1 expression as well. This probe was not present on the cDNA arrays used in this study, whereas two other TIMPs were downregulated (described later).

For periostin, collagen IV and collagen VI mRNA, we found a significant positive linear correlation between mRNA level and optic nerve injury grade in ONHs injured by elevated IOP ($R^2 = 0.51, 0.11, and 0.37$, respectively). When grouped by injury grade, levels of periostin and collagen VI were significantly elevated in ONH with optic nerve injury grades greater than 3.5, as well as in ONH from eyes after optic nerve transection (Fig. 4A). Type 2 α -collagen IV demonstrated a similar pattern, although only the increase after ONH transection was significant. By our microarray analysis, another isoform of collagen IV (type1 α) was increased 2.3-fold.

However, the pattern expression of other ECM-related messages demonstrated a different relationship to optic nerve injury. Using the same samples, as described in Figures 2A and 2B, we found that fibulin 2, tenascin C, and TIMP-1 mRNA levels showed a significantly nonlinear response to elevated IOP (F test for comparison of linear versus third order polynomial fit, P < 0.03), similar to that demonstrated by cyclin D1. For all three ECM messages, we found that the upregulation of expression in the experimental ONH groups with focal optic nerve injury was greater than in groups from eyes with more extensive injury due either to elevated IOP or transection (Fig. 4B).

In addition, microarray analysis indicated that elevated IOP led to the significant downregulation of some ECM components. Included were osteonectin/SPARC and two other SPARC-related genes (0.53–0.68-fold): collagen XVIII, which contains anti-angiogenic endostatin (0.7-fold) and two other TIMPs, TIMP-3, and -4 (both 0.7-fold). SPARC influences collagen fibril formation⁵⁰ and has recently been shown to promote retinal ganglion cell survival.⁵¹

Differential Regulation of TGF^β Isoforms

Various studies have associated increased expression TGF β 1 and - β 2 with human glaucoma or have found responses to experimentally elevated IOP and the deposition of ECM proteins in the ONH.^{45,52–56} Only the TGF β 2 isoform was present on our microarrays. By microarray analysis, we found a 29% decrease in the TGF β 2 mRNA level, compared with a nonsignificant 19% decrease in grade-5 ONHs by qPCR, using the ONH samples from Figure 2. When samples were grouped by injury mechanism and grade, there were no significant differences between groups (Fig. 5). However, a comparison of linear and nonlinear (third-order polynomial) fits to the expression data indicated a significant nonlinear fit (P < 0.02) in the pressure-injured ONHs. In contrast, the expression of TGF β 1 demonstrated a significant positive linear correlation with optic nerve injury in the ONHs damaged by elevated IOP ($P \le P$ 0.001), and ANOVA analysis found that the increase was significant for the group with grade-5 injury (Fig. 5; $P \le 0.01$). Two-way ANOVA revealed significant differences in the level of TGF β 1 and $-\beta$ 2 isoform expression in all elevated IOP injury groups with grades greater than 1.5. Together these data indicate that elevated IOP injury to the ONH results in differential regulation of TGF β isoforms, with an upregulation of TGF β 1 and a relative downregulation of TGF β 2.

Microarray analysis also implicated significantly altered expression of TGF β receptors and signaling pathway intermediates. Upregulated were TGF β receptor 1 (2.3-fold), TGF β receptor 2 (1.5-fold), SMAD3 (1.4-fold), and SMAD-specific E3 ubiquitin protein ligase 1 (SMURF1; 1.5-fold), whereas Pura 1 and Spectrin β 2, both regulators of SMAD signaling, were downregulated 0.73- and 0.64-fold, respectively.

Differential Regulation of Cytoskeletal Messages

The cytoskeleton was also a significantly affected cell component with upregulation of 48 and downregulation of 24 genes. Upregulated cytoskeletal genes were primarily associated with mitotic cell division (kinesin family member 22, protein regulator of cytokinesis 1, lamin B1, transforming acidic coiled-coil–containing protein 3, myosin heavy chain IX, centromere protein E, nucleolar and spindle-associated protein 1, and chromosome-transforming acidic coiled-coil–containing proteins (3.6- to 2-fold). Other classes were actin cytoskeletal assembly including gene actin-related 2/3 complex proteins, capping proteins, and dynein-1 (1.3- to 1.8 fold).

Among the downregulated genes, the only statistically significant cell component was the cytoskeleton (Table 6). Down-regulated cytoskeletal genes included many retinal ganglion cell–associated genes (kinesin family members 5B and 1B, microtubule-associated protein tau, myosin 1B, adducin 3 (gamma), tropomodulin 2, neurofilament L, and spectrin $\beta 2$ (0.17- to 0.66-fold). This indicates that the loss of axonal mRNA, 30-32 due to axon degeneration, was readily detectible in this study.

Downregulation of Lipid Biosynthesis

In addition to effects on the cytoskeletal cell component, elevated IOP resulted in the downregulation of genes associated with myelin and other lipid biosynthetic processes, including 3-hydroxy-3-methylglutaryl-coenzyme A synthase 1, 24-dehydrocholesterol reductase, sterol-C4-methyl oxidase-like, isopentenyl-diphosphate delta isomerase, farnesyl diphosphate farnesyl transferase 1, low-density lipoprotein receptor, 3-hydroxy-3-methylglutaryl-coenzyme A reductase, and ELOVL family member 6 (0.12- to 0.5-fold). Elevated IOP also decreased transferases, including glutathione transferases (0.5- to 0.65-fold).

Downregulation of Aquaporin-4 mRNA

Preliminary data from our examination of ONH injured by elevated IOP, using transmission electron microscopy, found vacuoles or swellings associated with ONH astrocytic endfeet (Morrison JC et al. *IOVS*. 2002;43:ARVO E-abstract 2885). Aquaporin-4, the principle water channel in astrocytes, is associated with astrocytic endfeet and is implicated in neural edema. After traumatic brain injury, aquaporin-4 mRNA levels are altered. 57-59 Because aquaporin-4 was not on our arrays, we used qPCR to determine the aquaporin-4 mRNA levels in the ONH samples described in Figure 2. We found that mRNA for the water channel protein was significantly decreased (to approximately 50% of that in the fellow eye values in all groups with optic nerve injury; Fig. 6). In addition, there was a significant nonlinear correlation between aquaporin level and the amount of nerve injury in the pressure injured ONHs (P < 0.01), with lowest levels in the ONHs with focal injuries.

Expression Levels of Connexin 43 and GFAP in ONH are Unaffected by Elevated IOP or Transection

Probes for Connexin 43, the astrocyte gap junctional protein, and GFAP, the principal astrocyte intermediate filament protein, were not present on our microarrays. Our previous immunohistochemical study of ONHs injured by elevated IOP demonstrated early loss of gap junctional labeling and an apparent initial dilution or decrease in GFAP label intensity with injury followed by intensification of GFAP labeling as glial scars were formed.²⁴ We used qPCR and the samples from Figure 2 to measure the levels of these messages in ONH injured by either elevated IOP or transection. There was no significant correlation between optic nerve injury grade and the mRNA levels for these proteins. In addition, when grouped by injury grade, there were no significant differences between groups (Fig. 7). Whereas there appears to be a decrease in these messages in nerves with focal injuries that is similar to that seen for aquaporin-4 and TGF β 2, fit by nonlinear correlation was not significantly better than a linear fit by regression analysis (F test; P > 0.05).

Discussion

This microarray analysis revealed that the dramatic reorganization of the ONH resulting from pressure-induced injury and associated axonal degeneration is reflected in extensive alterations in gene expression. These alterations significantly affect the processes of cell proliferation, immune response, and mRNA and protein synthesis and the lysosomal, cytoskeletal, ECM, and ribonucleoprotein cellular components. The measurement of selected message levels by qPCR both confirmed and extended these microarray analysis findings. In interpreting these alterations, it is important to recognize that the dissected ONH included portions of both the unmyelinated and the initial myelinated segment of the optic nerve. The myelinated optic nerve is well known to contain progenitor cells that proliferate in response to nerve injury.^{60–62} In addition, the microarray studies were limited to ONH with substantial damage due to elevated IOP.

Global Patterns of Altered ONH Gene Expression in Glaucoma Model

The most dramatic alteration is the increased cellularity of the nerve head that occurs in response to elevated IOP and ongoing axonal degeneration. Our data indicate that in extensively damaged ONH, the number of cells increased to almost three times the initial value. The large number of regulated cell cycle genes and their association with all cycle phases suggests that the increase in cellularity occurs by proliferation of endogenous nerve head cells, most likely glia or glial precursors. Our previous analysis of time course of nerve head alterations using immunohistochemistry suggested extensive cell division within the ONH glial columns based on heavy labeling with antibodies to proliferating cell nuclear antigen.²⁴ In this study, we used qPCR to examine the message levels for the key regulator of cell cycle progression, cyclin D1,

in nerve heads with a wide range of pressure-induced injury. We found that the initiation of cell division appears to be at its maximum in nerve heads that have been exposed to mild IOP elevation and have focal axonal degeneration. In these ONH segments, astrocytes, oligodendroglia, as well as adult optic nerve progenitor cells, potentially contribute to the proliferative response resulting from elevated IOP exposure.^{60–69} In addition, moderate upregulation of the mRNA for microglial calcium-binding protein, IBA1, the increased immunostaining with IBA1 antibodies and the increased mRNA levels for complement components all suggest microglial activation and a potential for microglial proliferation as well, as has been suggested by other observations.^{37–42,45,46,70} Studies designed to identify these proliferating ONH cells are currently under way in our laboratory.

Regardless of the cell type(s) involved, the magnitude of the cell proliferation, at least in these extensively injured ONH, implies a similarly extensive disruption of cellular organization within the nerve head. The normal interposition of astrocytes between the vasculature and axons is very likely affected, disrupting the activity-dependent supply of glucose and oxygen to the axons and adversely affecting their ionic environment. Our previous immunohistochemical investigation of ONH after exposure to elevated IOP identified the loss of connexin 43 gap junctional labeling as an early event in pressure-induced nerve injury.²⁴ Disruption of gap junctions implies a reduction in the ability of astrocytes to buffer the axonal environment and supply the metabolic demands of the nerve fibers. These compromised astrocytic functions may become critical factors in axon survival in the unmyelinated anterior portion of the ONH, where the energy demands are significantly higher than those for myelinated nerve.^{71,72}

The proliferation of glial cells in response to injury is well documented in neural tissues, including the myelinated optic nerve. $^{60-68}$ However, the extent of cellular proliferation in the ONH injured by glaucoma or experimentally elevated IOP, at least in the initial stages of injury, has not been generally recognized or extensively investigated. In rhesus optic nerves after 1 to 4 years of experimental glaucoma, the total number of glia appeared constant, although an increase in the optic nerve area occupied by glial tissue was reported.⁷³ After optic nerve transection, glial cells in the optic nerve and retinal ganglion cell layer have been reported to increase 41% and 210%, respectively,^{74,75} whereas normal aging in rhesus monkeys results in a 50% increase in optic nerve glia.⁷⁶ In addition, optic nerve astrocytes and lamina cribrosa cells proliferate in vitro in various experimental conditions.^{77–80}

The formation of an astrocytic scar, labeled by GFAP, is characteristic of glaucomatous injury. ^{24,73,81,82} However, in this study, we found that ONH GFAP mRNA levels were not significantly altered. This observation is consistent with our previous observation that, coincidentally with glial cell migration and disruption of ONH glial columns, GFAP label intensity appeared decreased or diluted. ²⁴ Intensified ONH immunolabeling for GFAP occurred later and accompanied glial scar formation. This labeling pattern could suggest the proliferation of adult glial progenitor cells in the pressure injured ONH. Although glial progenitor cells may not express glial cell type markers, they do express nestin, which was significantly upregulated in this study.⁸³ Our observations suggest that glial proliferation and gliotic scar formation are sequential and distinct processes in the ONH response to pressure-induced injury. This sequence may offer windows of opportunity for therapeutic agents to affect the proliferative or scarring processes.

Previously, human glaucoma and experimental glaucoma in both monkeys and rats has been characterized by the deposition of ECM components, including collagens, elastin, tenascin, laminin, fibronectin, and proteoglycans, within the ONH. 10-13,15,17,20,23,24,84,85 It has been hypothesized that these materials increase the susceptibility of the remaining axons to elevated pressure by changing the material properties of the nerve head and may contribute to

the inhibition of axon regeneration.^{22,23,82} Our microarray analysis confirms these previous observations by identifying significantly upregulated expression for the functional classes of ECM, ECM structural constituent, Golgi apparatus, and glycosaminoglycan binding. In addition to confirming the upregulation of collagens I, III, IV, and VI; fibronectin; and tenascin C, previously unidentified ECM components were also found to be significantly affected. Predominant among these was periostin, a fasciclin homologue that induces cell attachment and spreading and binds to heparin and alpha V integrin. Periostin upregulation is also associated with expression of TGF β , vascular hypertrophy and stress overload.^{86–88} Another significantly upregulated ECM component identified by our study was fibulin 2, a protein associated with elastic fiber formation.⁸⁹

Our previous studies in the rat suggested that there is a sequence of ECM deposition and that collagen VI deposition precedes the formation of the glial scar.²⁴ In this study, using qPCR to examine changes in gene expression in relationship to the degree of optic nerve injury, we observed two patterns of gene expression response. For several ECM components, including periostin, collagen VI, and collagen IV, expression increased linearly with increasing optic nerve injury. However, for other ECM components, expression patterns were not linear, but demonstrated the highest expression in ONH with focal injuries. These components included fibulin 2, tenascin C, and the matrix metalloproteinase inhibitor TIMP-1. This nonlinear pattern was similar to that seen for cyclin D1, suggesting that the expression of these ECM components is associated with early injury and ONH cell proliferation.

TGF β isoforms play important roles in the regulation of the expression of ECM components and in glial progenitor, astrocyte, and microglial proliferation.^{90–92} Therefore, we examined the changes in TGF β isoform expression in the ONH in response to elevated IOP. Again, we found two distinct patterns of expression. TGF β 1 expression increased linearly with increasing nerve injury, whereas the nonlinear expression pattern of TGF β 2 resulted in the lowest expression values in focally injured nerves, such that the difference in expression ratios between the two TGF β isoforms was significant in focally injured nerves. Astrocytes can produce both isoforms, whereas microglia are known to produce TGF β 1, and both isoforms inhibit microglial proliferation.^{93–95} Previously, Pena et al.⁵² have reported that whereas TGF β 2 mRNA levels appeared unaffected in glaucomatous ONH, protein levels were increased by both immunohistochemistry and bioassay, leading them to suggest a posttranslational regulation of this isoform. Together, these observations suggest that the differential regulation of TGF β isoforms may play important roles in the regulation of ONH response to elevated IOP.

In a pattern similar to that of TGF β 2, the principle water channel protein in astrocytes, aquaporin-4, showed the greatest downregulation in ONH from focally injured nerves. Aquaporin-4 immunolocalizes to the astrocytic endfeet that abut brain microvessels and subarachnoidal spaces and is also expressed by activated microglia.^{96,97} Aquaporins function to maintain neural volume and water homeostasis.⁹⁸ The astrocytic endfeet probably play a key role in the transfer of metabolites and water between the axons and the vasculature and subarachnoid space. Preliminary data from our electron microscopic studies of nerve heads exposed to elevated IOP identified these endfeet as sites of early injury (Morrison JC et al. *IOVS*. 2002;43:ARVO E-Abstract 2885). The downregulation of aquaporin-4 may be associated with the retraction of these endfeet accompanying the glial migration or proliferation that is associated with the disruption of the normal columnar arrangement of glial nuclei in the unmyelinated ONH. This morphologic alteration is a characteristic of exposure to elevated IOP in experimental monkey and rat glaucoma models.^{24,82} Our data confirm these observations and further point to astrocytic responses as early events in axonal injury.

For our qPCR studies, we included a group of ONH at 2 weeks after transection. Axon degeneration in the distal optic nerves from these eyes was similar to that in the highest pressure-induced injury group and gene expression levels were similar, so that no pressure-specific changes in gene expression were identified. However, this study was limited both by the number of genes measured and by the limitation of the transection ONH analysis to a single time point.

Our ONH microarray study also demonstrated downregulation of genes primarily associated with retinal ganglion cells. This most likely reflects a loss of optic nerve axons and the mRNA they contain. $^{30-32}$ Because axonal mRNA levels are low, we feel this gives a strong indication of the sensitivity of the cDNA microarrays used in the study.

Comparison of This Study with Other Microarray Studies of Cultured ONH Cells

In other microarray studies gene expression has been examined in cells cultured from the normal and glaucomatous ONH or normal ONH cells exposed to conditions hypothesized to be important in glaucoma pathogenesis. Although differences in microarray platform, array genes, species, experimental design, and determination of significance of expression changes complicate comparisons of these studies to ours, some commonalities can be found in all cases. In microarray studies of astrocyte cultures from normal and glaucomatous human ONH, 150 genes were identified as changed in expression.⁹⁹ These genes represented functional classes of signal transduction, cell adhesion, and proliferation, ECM synthesis, and degradation. In further studies of normal human ONH astrocytes exposed to hydrostatic pressure, approximately 600 genes were found to be altered in expression. 100 In common with this study, we also found increased expression of Fos, Ki-67 antigen, AXL receptor tyrosine kinase, Smad3, and Kcnn4 potassium channel. GFAP-negative cells isolated from human ONH have been exposed to mechanical strain or TGF β 1, to model proposed glaucomatous processes. With mechanical strain, the functional classes of ECM, cell proliferation, growth factor activity, and signal transduction were affected in these cells.¹⁰¹ Exposure to TGF β 1 resulted in expression of genes for ECM components, cell proliferation, angiogenesis, and growth factor binding. ¹⁰² To explore a potential role for epidermal growth factor in glaucoma, rat ONH astrocyte cultures were stimulated with EGF, and the upregulation of EGF receptor and other messages were evaluated.¹⁰³ In common, we found an upregulation of leukemia inhibitory factor, versican, Cd44, TGF β 1, and endothelin converting enzyme 1 in the pressure-damaged ONH.

Comparison of This Study with Other Glial Cell Microarray Studies

Most microarray studies of neurodegenerative diseases focus on the gene expression changes in affected neuronal populations, rather than those in the glial cells associated with degenerating nerves. However, there are several microarray analyses of glial cell responses that may be compared with our study.

Demyelinating disease, such as multiple sclerosis, is the focus of many of these investigations. 104-107 Genes upregulated in demyelinating disease that were also upregulated in our study included complement component 1q, $\beta 2$ microglobulin, FC receptor, histocompatibility (HLA) antigens, and polyA-binding protein.

Other studies have been conducted to investigate the difference between glial cell environments permissive and restrictive to axon outgrowth and regeneration. Kubo et al.¹⁰⁸ examined the changes in gene expression in the distal segment of peripheral nerves after transection, to identify genes associated with an environment supportive of nerve regeneration. In common with this peripheral nerve study, 22 of the same genes were similarly upregulated in the pressure-damaged ONH. These included tenascin C, CDC28 protein kinase regulatory subunit

2, complement component 1qb, prostaglandin E receptor 4, Ms4a6d, SKI-like, cathepsin D, tubulin β 2b, and β 4 thymosin.

For the study of characteristics of a central nervous system environment conducive to axonal regeneration, cultures of olfactory ensheathing cells (OECs) that promote axon outgrowth were compared to similar OEC cultures that do not.¹⁰⁹ Genes associated with a permissive environment that were also upregulated in our study included matrix metalloproteinase 2, versican, brain acidic membrane protein, and calretinin, whereas those upregulated in our study that were associated with an inhibitory environment were insulin-like growth factor 2 receptor, benzodiazepine receptor (translocator protein), and annexin.

Although in all these other studies, the expression of specific genes or gene classes seemed to respond similarly to the changes we observed in the ONH damaged by elevated pressure, it is difficult to draw any overall conclusions. However, several studies of cultured ONH cells did identify cell proliferation, the most significantly affected gene category in our study, as among the affected functional classes of genes.^{99,101,102}

Altered Expression of Specific ONH Genes in Focal Optic Nerve Injury

One unique feature of our study is the use of qPCR to compare selected gene expression changes with the degree of IOP damage. Using this powerful tool, we were able not only to confirm selected microarray findings but to extend them to discover two patterns of gene expression response in ONH injured by elevated IOP. One of these showed a positive correlation with the extent of nerve damage. The other pattern was distinctly nonlinear, demonstrating the greatest change in the eyes with focal injury and thus providing an opportunity to identify early gene changes that may play an important role in the pathogenesis of axon damage.

In ongoing work in our laboratory, we are extending the present study to isolate genes significantly regulated by early, focal injury in the unmyelinated portion of the optic nerve. By identifying these genes, we expect to identify specific processes by which the stress resulting from elevated IOP is translated to axonal injury within the ONH. The identification of these processes and determining their molecular pathways is likely to suggest therapeutic interventions that have the potential of limiting axonal injury in glaucoma.

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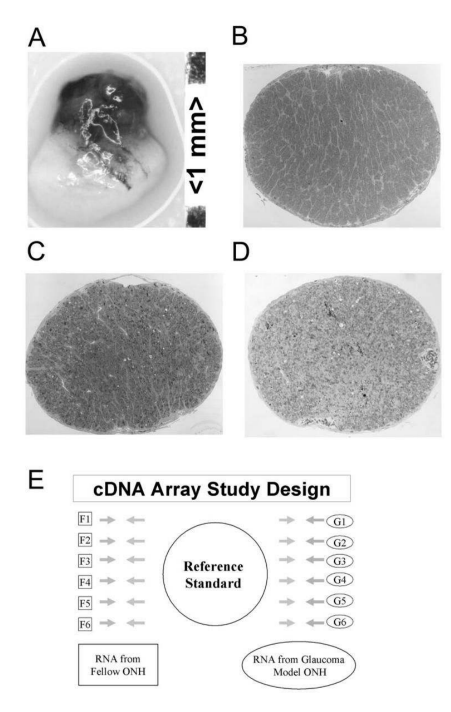


Figure 1.

(A) Example of the dissected ONH used in this study. Each 1-mm-long ONH consisted of the optic disc and the unmyelinated and the initial myelinated portions of the nerve. A representative optic nerve cross-section from a normal, fellow eye (control) is shown in (**B**). All fellow nerves were graded 1.0. For the microarray study, glaucoma model nerve cross-sections exhibited extensive axonal degeneration (**C**). This nerve was graded 4.7. In the microarray study, glaucoma model ONH had nerves with grades between 4.6 and 4.95, indicating ongoing and extensive axonal degeneration affecting approximately 50% of the axons. ONH with grade-5 optic nerves that demonstrated more extensive gliosis (**D**) were excluded from the microarray study. The experimental design of the microarray study is

illustrated in (E). For each gene, RNA from each of six fellow and six glaucomatous model eyes were independently compared by using 12 separate arrays with an ONH RNA reference standard.

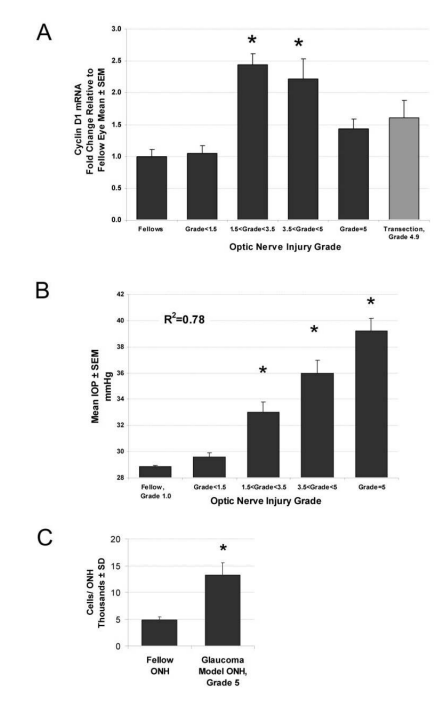


Figure 2.

Elevated IOP results in ONH cellular proliferation. mRNA levels of cyclin D, the key regulator for cell-cycle progression, are shown in (**A**) for ONH with graded optic nerve injury due to elevated IOP. ONHs were grouped according to injury grade (\blacksquare ; n = 7-11/group). (\blacksquare) For comparison, ONH from eyes with transected optic nerves (n = 7) are shown. Each bar represents the mean \pm SEM for that group relative to the mean for untreated fellow eyes (n = 14). *Significant increases compared with fellow eye levels (ANOVA with the Dunnett posttest, P < 0.01). (**B**) In a pressure-injured ONH, the relationship of mean IOP to optic nerve injury grade is shown over the 5-week experimental period, by group. Mean IOP differences between

groups are significant for all injury groups (*P < 0.05), except that between fellow and injury grade <1.5. For two separate groups of fellow and grade-5 ONH, total DNA measurements are shown in (C), illustrating the approximately threefold increase in cellularity in ONH with grade-5 optic nerve damage due to elevated IOP (*P < 0.000002).

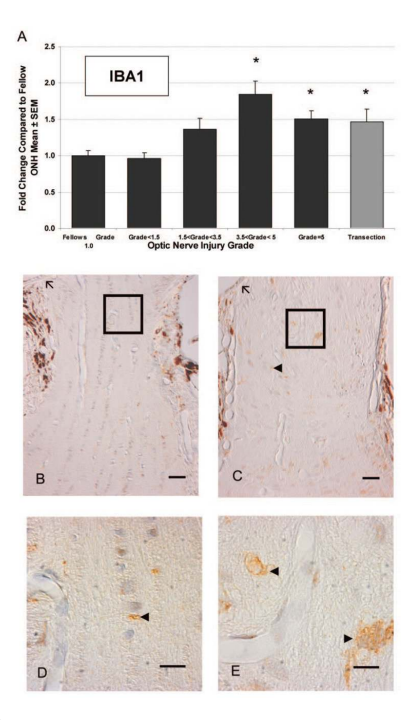


Figure 3.

Microglial activation in response to elevated IOP. In the same nerve injury groups as described in Figure 2, mRNA levels for the microglial marker, IBA1, were measured (**A**). *Significantly different from fellow eye (P < 0.05). IBA1 mRNA levels were significantly elevated in ONHs damaged by elevated IOP (**•**) with optic nerve injury grades greater than 3.5, as well as in ONHs at 2 weeks after optic nerve transection (**•**). Immunohistochemical labeling indicated activation and potential proliferation of ONH microglia. Representative fellow ONH (**B**) and glaucoma model ONH with an optic nerve injury grade of 3.6 (**C**). (**D**, **E**) Higher magnifications of the *boxed regions* in (**B**) and (**C**), respectively. *Arrowheads*: labeled cells; *arrows*: Bruch's

membrane. *Brown*: chromogen labeling; *pale blue*: counterstained nuclei. Scale bars: (**B**, **C**) 10 μ m; (**D**, **E**) 3 μ m.

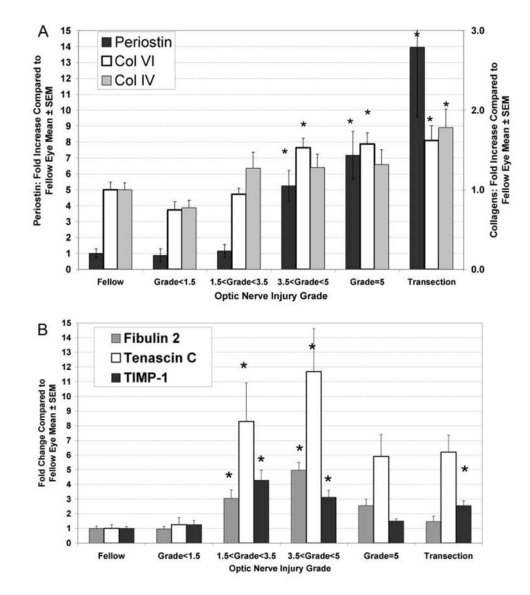


Figure 4.

Two response patterns for altered ONH ECM message expression accompanying pressureinduced injury. For the ECM components, periostin, collagen IV, and collagen VI, increasing optic nerve injury was accompanied by linear increases in ONH message level (**A**). For periostin and collagen VI, ONH groups with injury grades greater than 3.5 demonstrated significant increases compared with the fellow eye group (*P < 0.05). Note that the magnitude of increase for periostin is much greater, as indicated by the fivefold difference in *y*-axis scale labels. Collagen IV demonstrated a similar pattern, although the increase was only statistically significant in the transection group. In contrast, message levels for fibulin2, tenascin c, and TIMP-1 were highest in ONHs from eyes with focal optic nerve injury (**B**). *Significant elevation relative to untreated, fellow ONH (P < 0.05). In addition, for fibulin 2, the grade-3.5 to less than grade-5 group, had significantly greater responses than all other groups. For TIMP-1, the grade-1.5 to less than grade-3.5 group, had significantly greater values than all groups, except the grade-3.5 to less than grade-5 group. ANOVA with the Tukey multiple comparison posttest.

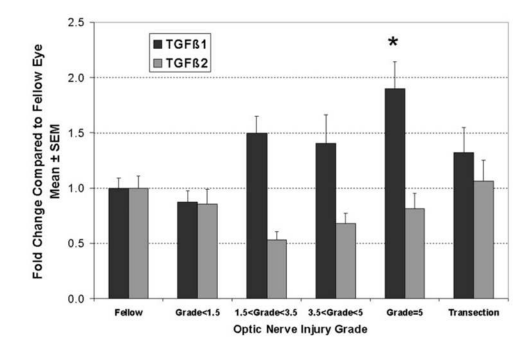


Figure 5.

TGF β isoform mRNAs were differentially regulated in pressure-injured ONH. For TGF β 1, there was a significant increase in mRNA level that correlated with the degree of optic nerve injury, so that levels were significantly elevated in the group of pressure-injured ONHs with grade 5 (**P* < 0.01). In contrast, expression levels for TGF β 2 demonstrated a significantly nonlinear correlation to optic nerve injury grade, with lowest levels in ONH with focal optic nerve injury. This resulted in a significant (approximately twofold) difference in the ratio of TGF β 1 to TGF β 2 in all ONHs with pressure-induced optic nerve injury grades greater than 1.5.

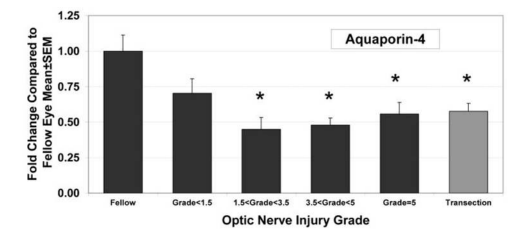


Figure 6.

ONH Aquaporin-4 mRNA levels decreased in a pattern similar to that of TGF β 2. For all injury groups, except that with injury grades <1.5, there was a significant decrease in aquaporin-4 mRNA expression (**P* < 0.05) compared with levels in the fellow eye. Correlation of expression levels with optic nerve injury grade demonstrated a significant, nonlinear fit, with the lowest expression levels in ONH from eyes with focal optic nerve injury (F test; *P* < 0.01).

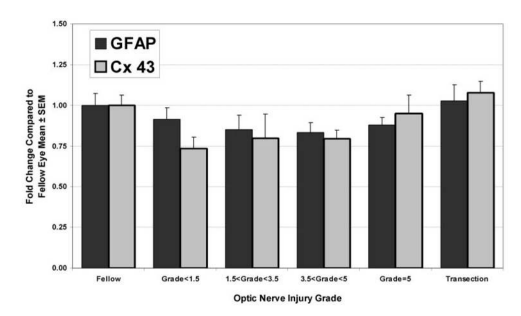


Figure 7.

Expression levels of two other proteins associated with differentiated astrocytes, GFAP and connexin43, were not altered in the ONH by either elevated IOP or transection. Although the pattern of expression in the various optic nerve injury groups may appear similar to that seen for TGF β 2 and aquaporin-4, levels in the optic nerve injury groups were not significantly different from fellow eye levels. Regression analysis also indicated no significant correlation.

Primers Sequences for qPCR

Target	Forward Primer	Reverse Primer	bp
Aquaporin-4	CATGAATCCAGCTCGATCCT	TTCTCGTCTCCACGGTCAAT	297
Collagen IV ($\alpha 2$)	AGGGATACCCGGCATAAC	GGCCTCTGCTTCCTTTCT	119
Collagen VI $(\alpha 1)$	CCAGGTCGAGCGTCTCTTCA	ACGTCGTACTCGGCGGTCTT	332
Connexin 43	CGTGCCGCAATTACAACAAG	TTGGTCCACGATGGCTAATG	194
Cyclin D1	CTGCGATGCAAGGCCTGAAC	GCGCAGGCTTGACTCCAGAA	152
Fibulin-2	CGCTGTCTGCGCTTCGATTG	GGCCGATGCGGAAGATATGG	175
GAPDH	TGCCACTCAGAAGACTGTGGATG	GCCTGCTTCACCACCTTCTTGAT	249
GFAP	GCGGAGAACAACCTTGGCTGTGTA	GCAGTTGGCGGCGATAGTCATTA	376
Microglia (IBA1)	TCCGAGGAGACGTTCAGTTA	CTGGTCTCACAACTGCTTCTT	237
Periostin	GGAAGAACGCATCATCACAG	CATCGCCACCTTCAATGAAC	137
Tenascin C	GAGTCTGTGGATGGTACAGT	ATGTATAGAGGCCAGAGGT	236
TGFβ1	TTGGAGCCTGGACACACA	TTGCGACCCACGTAGTAGAC	132
TGFβ2	CATGCCCTTATCTGTGGAGTTC	TGGGCGTATTGCCAATGT	150
TIMP1	GTTCCCTGGCATAATCTGAG	GGGATGAGGATCTGATCTGT	155

Table 1

Table 2 Genes Upregulated More Than Threefold in ONHs with Elevated-IOP-Induced Injury

Change (x-Fold)	GenBank Accession	Unigene	Name	Gene Ontology Summary
12.8	AW553287	Mm.	Periostin, (fasciclin I related)	Integrin-mediated cell adhesion (beta-3),
6.1	AW550270	236067 Mm.980	Tenascin C	ECM, enhances motility Extracellular space, localized in tendons, expressed by astrocytes, disrupts adhesion supports migration
5.9	BG083139	Mm.6856	Securin, pituitary tumor-	Cell cycle, chromosome segregation,
5.3	BG083522	Mm.	transforming 1 CDC28 protein kinase	cysteine protease inhibitor activity Cell cycle, cyclin-dependent protein kinas
4.7	AU018403	222228 Mm.6856	regulatory subunit 2 Nuclear receptor subfamily 5, group A, member 2	activity; cytokinesis Regulates cholesterol degradation, regulation of transcription, DNA dependent
4.6	BG085568	Mm.29798	CD34 antigen	Cell adhesion, external side of plasma membrane
4.4	AW547306	Mm.3453	Complement component 1, q subcomponent, gamma polypeptide	Complement activation
4.4	BG079311	mm.113278	Potassium voltage-gated channel, delayed-rectifier, subfamily S, member 3	Potassium ion transport, voltage-gated io channel activity
4.3	BG080700	Mm.28518	Tumor necrosis factor receptor superfamily, member 12a, TWEAK receptor	Migration of endothelial cells, positive regulation of axon extension, adhesion
4.3	BG069098	Mm. 274086	Kif18a: kinesin family member 18A	ATP binding, microtubule-based movement, motor activity
4.3	BG079289	Mm. 373656	Riken cDNA 2700099C18 gene	Cell division and chromosome segregation
4.2	BG063624	Mm.24337	PDZ binding kinase	Phosphorylates MAP kinase p38. Seems be active only in mitosis.
4.1	BG076632	Mm. 288206	Adenylate cyclase 7	Adenylate cyclase activation, guanylate cyclase activity, integral to membrane
4.0	BG080803	288200 Mm.9749	B-cell linker	B-cell differentiation humoral immune
4.0	BG079781	Mm. 181237	Ribonuclease T2	response Mitochondrial RNAse activity
4.0	BG079350	Mm. 225956	Kinetochore associated 2	Actin binding, chromosome, pericentric region, cytoskeleton, mitotic metaphase
3.8	BG087868	Mm.2570	Complement component 1, q subcomponent, beta polypeptide	Complement activation
3.7 3.7	BG087390 BG074248	Mm.4237 Mm. 286488	Topoisomerase (DNA) II alpha Kinesin family member 22	Chromosome segregation Microtubule-based process, anaphase, nucleus
3.5	BG067419	Mm. 281298	Growth arrest and DNA- damage-inducible 45 gamma	Activation of MAPKK, negative regulation of protein kinase activity, regulation of co cycle
3.5	BG083088	Mm. 273049	Cyclin D1	Cyclin-dependent protein kinase holoenzyme complex, protein kinase activity
3.5 3.4	BG063496 BG074814	Mm.9916 Mm.370	Centrosomal protein 55 Complement component 1, q subcomponent, alpha polypeptide	Cell division and chromosome partitionin Complement activation
3.5	BG085131	Mm.22673	Fc receptor, IgE, high affinity 1, gamma polypeptide	Antigen presentation, neutrophil chemotaxis, positive regulation of immu response, positive regulation of interleukin-10 biosynthesis, transmembrane receptor activity
3.4 3.3	BG070808 BG073145	NA Mm.22448	Immunoglobulin Cytoskeleton-	Unknown
3.3	BG064593	Mm.12508	associated protein 2 Karyopherin (importin) alpha 2	Nuclear protein import as an adapter
3.2	BG085966	Mm. 249146	Fibulin 2	protein for nuclear receptor KPNB1 Calcium-dependent binding to ECM components, elastic fiber assembly, elastosis
3.2	BG069532	Mm. 282556	Niemann Pick type C2	Egress of lipids from the lysosome
3.2	BG076991	282556 Mm.3468	Suppressor of cytokine signaling 3	Negative regulation of cytokine signalin, via the JAK/STAT pathway, activated by insulin-like growth factor-1 receptor, blocks IL-6 signaling.

Change (x-Fold)	GenBank Accession	Unigene	Name	Gene Ontology Summary
3.1	BG078426	Mm. 380027	Homologous to G2/mitotic- specific cyclin B1	Control of the cell cycle at the G2/M (mitosis) transition
3.1	BG064098	Mm. 171335	Ĝolgi phosphoprotein 2	Transport of protein cargo through the Golgi

Table 3 The 30 Genes Most Downregulated in Expression in ONHs with Elevated-IOP-Induced Injury

Change (x-Fold)	GenBank Accession	Unigene	Name	Gene Ontology Summary
0.04	BG069400	Mm.41035	Unc-13 homolog C (C. elegans)	Synaptic and impulse transmission neurotransmitter release
0.12	BG069739	Mm.61526	3-hydroxy-3-methylglutaryl- Coenzyme A synthase 1	Cholesterol biosynthesis
0.14	BG076743	Mm. 200203	RUN and TBC1 domain containing 2	GTPase activation, vesicle-mediate transport
0.17	BG076766	Mm.2108	Transthyretin	Retinol and thyroxine transport from the bloodstream to the brain
0.26	BG086656	Mm. 250866	Aldehyde dehydrogenase (NAD)	Aldehyde dehydrogenase (NAD) retinoic acid metabolism
0.27	BG074458	Mm. 133370	24-dehydrocholesterol reductase	Cholesterol biosynthesis, inactivation of MAPK
0.27	BG065170	Mm. 173337	Mitogen-activated protein kinase 8 interacting protein 2	JNK cascade, kinase activity, prote binding
0.30	BG083949	Mm.30119	Sterol-C4-methyl oxidase-like	Sterol biosynthesis
0.30	BG076999	Mm. 223744	Kinesin family member 5B, Kinesin 1	Microtubule-based movement, heavy chain, axonal transport
0.32	BG068358	Mm. 181166	Rabphilin 3A	Rab GTPase binding, recruitment e synaptic vesicles for exocytosis
0.33	BG079889	Mm.29847	Isopentenyl-	Cholesterol biosynthesis
			diphosphate delta isomerase	
0.35	BG073378	Mm. 360538	Acetyl-coenzyme A acetyltransferase 2	Acyltransferase activity, transferas activity, cholesterol biosynthesis
0.35	BG084178	Mm.45372	Protein phosphatase 1, regulatory (inhibitor) subunit 1B	Protein phosphatase inhibitor activity, dopaminergic signal transduction
0.36	BG069211	Mm. 371560	Farnesyl diphosphate farnesyl transferase 1	Cholesterol biosynthesis
0.38	BG063992	Mm.3213	Low-density lipoprotein receptor	Cholesterol metabolism
0.38	BG080229	Mm.7729	Aldolase 3, C isoform	Glycolysis
0.39	BG068310	Mm. 229532	Endothelin receptor type B	G-protein-coupled receptor activit nonspecific endothelin receptor, regulation of blood pressure
0.42	BG070284	Mm. 292102	RAB30	Small GTP-binding protein of the RAB family
0.42	BG071079	Mm. 228798	Patched homolog 1	Hedgehog receptor activity, negati regulation of mitosis, Wnt and TGFβsignaling
0.42	BG076141	Mm. 257073	Chimerin (chimaerin) 1	Rho GTPase activating protein, is transcriptionally upregulated after exposure to neurotrophins, enhance neurite outgrowth
0.43	BG079550	Mm. 272361	Protease (rosome, macropain) 26S subunit, ATPase 5	Protein catabolism, thyroid hormo receptor interactor 1
0.43	BG082257	Mm. 273108	Abhydrolase domain containing 3	Biological process unknown, catalytic activity, hydrolase activit integral to membrane
0.43	BG088107	Mm.55143	Dickkopf homolog 3 (Xenopus laevis)	Negative regulation of Wnt recept signaling pathway
0.43	BG087831	Mm. 320691	Potassium voltage-gated channel, Shal- related family, member 2	Repolarization phase of the action potential, outward K+ currents
0.44	BG071519	Mm.70950	SRY-box containing gene 21	Repression of NGF-induced neuri outgrowth in PC12 cells
0.44	BG071963	Mm.54126	IPLA2-2	Membrane-associated, calcium- independent phospholipase A2 gamma, fatty acid metabolism
0.44	BG078816	Mm. 316652	3-Hydroxy-3-methylglutaryl- coenzyme A reductase	Cholesterol biosynthesis
0.45	BG073755	Mm. 130063	Riken cDNA 9330182L06 gene	Receptor activity
0.45	BG066372	Mm.1287	Microtubule-associated protein tau	Negative regulation of microtubul depolymerization, axonal microtubules
0.45	BG068973	Mm.13148	Riken cDNA 1110064P04 gene, mRNA	Homology to AHA1, activator of heat shock 90-kDa protein ATPas homolog 2

Table 4

Significantly Regulated Gene Classes by DAVID Analysis of All Genes Identified by SAM as Significantly Altered in Expression

Biological Process GO Term: Number of Genes (P)	Cell Component GO Term: Number of Genes (P)	Molecular Function GO Term: Number of Genes (P)
Cell proliferation: 113 (<0.0002)	Lysosome: 26 (<0.000009)	Structural constituent of ribosome: 56 (<0.00001)
Immune response: 38 (<0.0004)	Cytoskeleton: 72 (<0.0006)	ECM structural constituent: 13 (<0.002)
Ribosome biogenesis: 23 (<0.02)	ECM: 34 (<0.002)	Cytoskeletal protein binding: 39 (<0.002)
Sterol synthesis: 7 (<0.02)	Ribonucleoprotein complex: 66 (<0.002) Spindle: 9 (<0.02) Cell surface: 7 (<0.04)	Transmembrane receptor activity: 36 (<0.01) Peroxidase activity: 7 (<0.02) Glycosaminoglycan binding: 9 (<0.03) Rho small monomeric GTPase activity: 6 (<0.03) Metal ion binding: 97 (<0.04) Cytokine binding: 6 (<0.05) Transferase activity, transferring hexosyl groups: 12 (<0.05) Long-chain-fatty-acid-CoA ligase activity: 4 (<0.05)

Listed in order of ascending probabilities, P < 0.05 for all listed classes. For nested gene ontology (GO) terms, the most descriptive parent term is listed.

Table 5

Significantly Upregulated Gene Classes by DAVID Analysis of Significantly Upregulated Genes Identified by SAM

Biological Process GO Term: Number of Genes (P)	Cell Component GO Term: Number of Genes (P)	Molecular Function GO Term: Number of Genes (P)
Immune response: 34 (<0.00001)	Lysosome: 25 (<0.0000001)	Structural constituent of ribosome: 52 (<0.00000001)
Cell proliferation: 87 (<0.0001)	Ribonucleoprotein: complex: 57 (<0.00002)	ECM structural constituent: 12 (<0.0003)
Ribosome biogenesis and assembly: 21 (<0.0003)	ECM: 28 (<0.0006)	Glycosaminoglycan binding: 9 (<0.003)
Protein metabolism: 182 (<0.001)	Cytoskeleton: 48 (<0.02)	Transferase activity, transferring glycosyl groups: 19 (<0.003)
Cell adhesion: 35 (<0.03)	Cell surface: 6 (<0.04)	Kinase regulator activity: 10 (<0.02)
	Spindle: 7 (<0.04)	Cytoskeletal protein binding: 27 (<0.02)
	Golgi apparatus: 33 (<0.05)	Transmembrane receptor activity: 26 (<0.04)
	5 1 1	Hydrolase activity, acting on glycosyl bonds: 11 (<0.04)
		Rho small monomeric GTPase activity: 5 (<0.04)
		Peptidase activity: 39 (<0.05)

Data are as described in Table 4.

Table 6 Significantly Downregulated Gene Classes by DAVID Analysis of Downregulated Genes Identified by SAM

Biological Process GO Term: Number of Genes (P)	Cell Component GO Term: Number of Genes (P)	Molecular Function GO Term: Number of Genes (P)
Lipid biosynthesis: 11 (<0.01)	Cytoskeleton: 23 (<0.04)	Transferase activity, transferring alkyl or aryl (other than methyl) groups: 8 (<0.002)
Main pathways of carbohydrate metabolism: 9 (<0.04)		Ligase activity, forming carbon-sulfur bonds: 4 (<0.01)
Amino acid derivative metabolism: 6 (<0.05)		Metal ion binding: 38 (<0.02)

Data are as described in Table 4.