Global Gene Expression Changes in Rat Retinal Ganglion Cells in Experimental Glaucoma

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Purpose. Intraocular pressure (IOP) is an important risk factor in glaucoma. Gene expression changes were studied in glaucomatous rat retinal ganglion cells (RGCs) to elucidate altered transcriptional pathways.

METHODS. RGCs were back-labeled with Fluorogold. Unilateral IOP elevation was produced by injection of hypertonic saline into the episcleral veins. Laser capture microdissection (LCM) was used to capture an equal number of RGCs from normal and glaucomatous retinal sections. RNA was extracted and amplified, labeled, and hybridized to rat genome microarrays, and data analysis was performed. After selected microarray data were confirmed by RT-qPCR and immunohistochemistry, biological pathway analyses were performed.

Results. Significant changes were found in the expression of 905 genes, with 330 genes increasing and 575 genes decreasing in glaucomatous RGCs. Multiple cellular pathways were involved. Ingenuity pathway analysis demonstrated significant changes in cardiac β -adrenergic signaling, interferon signaling, glutamate receptor signaling, cAMP-mediated signaling, chemokine signaling, 14-3-3-mediated signaling, and G-protein-coupled receptor signaling. Gene set enrichment analysis showed that the genes involved in apoptotic pathways were enriched in glaucomatous RGCs. The prosurvival gene Stat3 was upregulated in response to elevated IOP, and immunohistochemistry confirmed that Stat3 and phosphorylated-Stat3 levels were increased in RGCs in experimental glaucoma. In addition, the expression of several prosurvival genes normally expressed in RGCs was decreased.

Conclusions. There are extensive changes in gene expression in glaucomatous RGCs involving multiple molecular pathways, including prosurvival and prodeath genes. The alteration in the balance between prosurvival and prodeath may contribute to RGC death in glaucoma. (*Invest Ophthalmol Vis Sci.* 2010;51: 4084 - 4095) DOI:10.1167/iovs.09-4864

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Glaucoma is the second leading cause of blindness worldwide. It is a neurodegenerative disease that is characterized by the slow and progressive degeneration of retinal ganglion cells (RGCs) and their axons, leading to atrophy of the optic nerve and loss of vision. Elevated intraocular pressure (IOP) is the leading risk factor for loss of RGCs and development of optic nerve atrophy. It is now clear that RGCs die by apoptosis in glaucoma.²⁻⁵ However, the trigger of the apoptosis is still unknown. To successfully target the mechanisms, we must to understand the molecular signaling networks in RGC death. We used laser capture microdissection (LCM) to specifically capture RGCs, and microarray technology, an accepted and powerful tool for large-scale gene expression profiling, to elucidate a global view of gene interaction networks in RGCs.

To date, several gene expression studies have been performed on the retina as a whole, to catalog changes in transcription that accompany glaucoma. The mRNA expression patterns in whole retina of several animal models of glaucoma, including DBA/2J mice,6 cynomolgus macaque monkeys, rat,⁸⁻¹¹ and C57B16/J mice¹² have been investigated. These studies revealed up- and downregulation of many genes in response to elevated IOP. However, the outcome and interpretation of these microarray studies can be strongly affected by inherent tissue heterogenicity, as the retina is a complex tissue composed of neuronal, glial, and vascular cell types. The RGCs comprise only 1% or less of the retinal cells. It is therefore difficult to causally link changes in gene expression patterns from whole retina to specific RGC gene expression, the associated pathways and subsequent physiological effects. In addition, site-specific, pathology-related gene expression changes can be diluted by the gene expression profile of other cell types in a whole retinal homogenate and thus not be readily detectable. Previous efforts to identify the complete picture of retinal gene expression may thus have missed important RGC genes, due to the low relative frequency of RGCs and, hence, their mRNA transcripts. LCM overcomes such obstacles. 13 LCM allows morphologic preservation of tissue and the laser does not damage macromolecules such as DNA, RNA, or proteins.

In the present study, we sought to investigate the wholegenome regulation of RGCs under conditions of elevated IOP. We hypothesized that gene expression would change in glaucomatous RGCs compared with normal RGCs in a rat model of experimental glaucoma. To test our hypothesis, we applied LCM to capture an equal number of glaucomatous RGCs and normal RGCs and used microarray analysis to identify the genes and functional gene classes most altered in expression in the rat RGCs after exposure to experimentally elevated pressure. Then, we used real-time quantitative PCR (RT-qPCR) to validate selected changes in gene expression identified by microarray analysis. The JAK/STAT signaling pathway, especially the Stat3 protein, has been implicated in neurodegenerative and neuroprotective mechanisms.¹⁴ The phosphorylation of Stat3 is known to activate Stat3 and has been shown to be protective of RGCs after ischemic insult. 15-17 The increased transcript expression of the prosurvival gene, signal transducer, and activator of transcription 3 (Stat3) was confirmed at the protein level in RGCs by immunohistochemistry (IHC). In addition, Stat3 phosphorylation was seen in RGCs after IOP elevation.

Methods

Study Design

The study was performed in five adult male Brown Norway rats. RGCs were back-labeled with Fluorogold. Seven days later, unilateral elevation of IOP was produced in one eye by injecting hypertonic saline into episcleral veins. The fellow eye served as the control. After 10 days of elevated IOP, the rats were killed, and the eyes were processed for LCM. First, to confirm the enrichment of RGCs, the gene expression of glial fibrillary acidic protein (Gfap, an astrocyte marker) and thymus cell antigen 1, theta (Thy1, an RGC marker) was compared between captured RGCs and captured ganglion cell layer (GCL) from normal eyes. Next, two thousand RGCs were captured from each normal eye and each eye with elevated IOP in three rats. Gene expression in the glaucomatous RGCs was compared with that in the fellow normal RGCs by gene microarray (Rat GeneChip 230.2; Affymetrix, Santa Clara, CA). Finally, an equal number of RGCs were captured from five rats, including the three rats from the microarray experiments. RTqPCR was used to verify selected changes in gene expression initially identified by microarray analysis. Retinal tissue sections were also used to perform IHC localization of Stat3 and phosphorylated-Stat3.

Experimental Glaucoma

All animal-related procedures were performed in compliance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. Five adult male Brown Norway rats (300-450 g; Charles River, Wilmington, MA) were maintained in covered cages with a 12-hour light-dark cycle and fed with a standard rodent diet ad libitum.

RGCs were bilaterally back-labeled by stereotaxic injection of 3% Fluorogold (Fluorochrome LLC, Denver, CO) into the superior colliculus of deeply anesthetized rats, as described previously.³ The animals were allowed to recover for a week to allow for adequate back-labeling of RGCs before experimental glaucoma was induced.

Anesthesia was induced with a mixture of acepromazine maleate (1.5 mg/kg), xylazine (7.5 mg/kg), and ketamine (75 mg/kg; all from Webster Veterinary Supply, Sterling, MA). To elevate IOP, hypertonic 1.9 M saline was injected into limbal aqueous humor-collecting veins of one eye, as originally described by Morrison et al.¹⁸ The fellow eye served as the control.

IOP was measured with a handheld tonometer (TonoPen XL; Medtronic Ophthalmics, Jacksonville, FL). Before the first hypertonic saline injection, baseline IOPs for both eyes were measured on three consecutive days for each rat. During the induction of experimental glaucoma, IOP in each eye was measured every other day while the rats were in the awake state. To minimize diurnal variability in IOP, measurements were taken between 10 AM and 2 PM. To assess total IOP exposure, we calculated integrated IOP as the area under the pressure-time curve (AUC = experimental eye — control eye), beginning with the day of the first saline injection. After IOP had been elevated for 10 days, the rats were killed by asphyxiation with carbon dioxide, the eyes were rapidly removed, and the eye cups were collected.

Tissue Preparation and Assessment

After the eyes were enucleated, the posterior segments were minimally fixed in 4% paraformal dehyde for an hour and cryoprotected with serial sucrose dilutions. Eye cups were frozen in optimal cutting temperature compound (OCT; Tissue-Tek; Miles Inc., Diagnostic Division, Elkhart, IN) and sectioned in their entirety at 16 μm . These sections were used for RGC counting, LCM of RGCs, and IHC. Estimation of total RGC number in each retina was performed with statistically unbiased stereological protocols by using an optical dissector and a fractionator, as previously described.^{3,19} Fluorogold-positive cells were immunostained with an anti-Fluorogold antibody (1:200, Fluorochrome LLC) and counted (C.A.S.T. system; ver. 2.3.1.2; Olympus, Albertslund, Denmark).

Optic nerves were assessed in saline-injected eyes and fellow eyes. All corresponding optic nerve sections were taken from approximately 2 mm posterior to the globe, fixed in glutaraldehyde, embedded in plastic, stained with toluidine blue, and evaluated by light microscopy. Optic nerve damage was graded from 1 (no injury) to 5 (degeneration involving the total nerve area) by four masked observers who used a modification of the method described by Morrison.^{3,18}

LCM of Fluorogold-Labeled RGCs

For LCM, a commercial system (PixCell IIe; Arcturus, Mountain View, CA) was used, as previously described. 3,20 Briefly, the tissue sections were dehydrated by serial immersions in 75%, 95%, and 100% ethanol for 2 minutes, followed by a 2-minute immersion in xylene. Tissue sections were then air-dried for 2 to 5 minutes at room temperature. A cleanup pad (Cap-Sure; Arcturus) was used to remove any extra tissue on the slide. RGCs were identified by the presence of Fluorogold back-labeling (excitation filter, 330-380 nm; barrier filter, 400 nm). An equal number of RGCs from each eye were captured onto single caps (Cap-Sure MacroLCM Caps; Arcturus), by using the following parameters: spot size, 7.5 μ m; power, 65 mW; pulse duration, 750 μ s. For the microarray experiment, 2000 RGCs were captured from each eye. RGCs were captured from approximately 30 sections per eye. Because RGCs were captured from retinal cross sections, the sampled area included central and peripheral retina. The microdissected samples were collected in a noncontact and contamination-free manner, immediately lysed by 20 µL LCM extraction buffer (Arcturus). For RT-qPCR, 2600 RGCs were captured from each eye.

RGC Enrichment

To assess the enrichment of RGCs, LCM-captured RGCs from a normal eye were compared to LCM-captured GCL from the same normal eye. The gene expression of *Gfap* and *Tby1* was compared between LCM-captured RGCs and LCM-captured GCL.

RNA Isolation and Microarray Experiments

For each eye of the rats (n=5 eyes), the Fluorogold-labeled RGCs on each cap were lysed with 20 μ L of extraction buffer for 30 minutes at 42°C, and centrifuged (12,500g) from the caps into microfuge tubes. Isolated RGCs were processed for RNA extraction (PicoPure RNA; Arcturus) according to the manufacturer's protocol. The total cellular RNA was finally resuspended in 14 μ L elution buffer. One microliter of the isolate was used to determine the quality of the RNA by detecting 28S/18S ribosomal RNA peaks (RNA Pico 6000 LabChip with a model 2100 Expert Bioanalyzer; Agilent Technologies, Palo Alto, CA).

Isolated RNA was amplified by linear amplification (RiboAmp HS RNA amplification kit; Arcturus). One microliter of the resuspended isolate was run on the bioanalyzer, to estimate average fragment length of the aRNA molecules. The labeling and hybridization of the cRNA was performed at the NIH Neuroscience Microarray Consortium (Keck Microarray Resource, New Haven, Yale University School of Medicine, New Haven, CT). Briefly, 10 µg of total RNA was used for cRNA labeling, according to the recommended microarray protocol (Affymetrix) and hybridized to the rat gene chip (Rat 230.2 chip; Affymetrix) for 16 hours at 45°C in a rotisserie oven. After hybridization, the arrays were washed using the fluidics station and stained with streptavidin-phycoerythrin according to the technical manual (Affymetrix). Washed arrays were scanned (GeneChip Scanner 3000; Affymetrix). Scanned output files were visually inspected for hybridization artifacts and then analyzed (GeneChip Operation Software [GCOS]; Affymetrix). The software recorded intensities for perfect match (PM) and mismatch (MM) oligonucleotides, and determined whether genes were present (P), marginal (M), or absent (A).

Microarray Data Analysis

The output .CEL files obtained from the software server (GCOS; Affymetrix) were processed and normalized by dChip.²¹ The probe level analysis was performed by dChip, which normalizes arrays to the PM and MM probes before computing model-based expression levels. Briefly, invariant set normalization was performed, the PM-only model was used for generating gene signal intensities, and the significance of changes in gene expression due to elevated pressure was determined. The parameters were set with a false-discovery rate (FDR) of 5%, and a minimum change of twofold, to identify the genes with potential biological significance. Because of the small sample size, a threshold of the percentage of samples called "present" in the samples used in both groups was set as 30%, to filter the significantly changed genes. That means the significantly changed genes that meet dChip comparison criteria may be absent from some chips but not all chips. Biologically, it is reasonable that genes expressed in diseased cells may not be present in normal cells or vice versa. Therefore, only the genes absent in all samples were ignored. All microarray data have been deposited in the NIH/NLM Gene Expression Omnibus (GEO, http://www.ncbi.nlm. nih.gov/projects/geo/ provided in the public domain by the National Center for Biotechnology Information, Bethesda, MD) and are accessible through GEO Series accession number GSE12596.

Then, the genes were subjected to bioinformatics analysis. Pathway analysis (Ingenuity Systems, Redwood City, CA) was performed to determine canonical pathways with expression that is differentially affected between glaucomatous RGCs and normal RGCs (www. ingenuity.com). Gene set enrichment analysis (GSEA) was used to assess whether a set of functionally related genes would show statistically significant differences between glaucomatous RGCs and normal RGCs. ^{22,23}

Real-Time Quantitative PCR

Total RNA was isolated from 2600 RGCs from five pairs of eyes, including the same set of eyes used for microarray analysis plus an additional two pairs (Table 1). The procedure is the same as the RNA extraction for microarrays. Reverse transcription (RT) was performed, according to the manufacturer's specifications (Invitrogen Corp., Carlsbad, CA), with 13 μ L of extracted RNA from each LCM sample and 200 U of RT polymerase (Superscript; Invitrogen).

The amount of cDNA corresponding to 2600 RGCs was amplified with the primer pairs for *Gapdh*, *Nrg1*, *Cpg15*, *Ywbab*, *Ywbaz*, *Grb2*, *Map2k1*, *C3*, *C1qa*, *C1qb*, *Stat3* and *Stat5a* (Table 2). mRNA extracted from Brown Norway rat brain was reverse transcribed and used as the reference sample. RT-qPCR was performed (iCycler iQ; Bio-Rad, Hercules, CA) with SYBR green PCR master mix (Bio-Rad). A computer program (SDS2.1 software; Bio-Rad) was used to visualize the data. The standard curve method was used to determine relative changes in gene

expression levels with *Gapdb* serving as the reference, which did not change significantly in our samples. A Student's *t*-test was conducted to compare the normalized relative expression values between the control and high-IOP eyes, to determine statistical significance. To confirm the identity of real-time PCR reaction products, a DNA chip (Agilent Technologies) and a bioanalyzer (model 2100 Agilent Technologies, Palo Alto, CA) were used to verify the size of the products. The amplicons of selected genes, *Nrg1* and *Cpg15*, were sequenced at the DNA Sequencing Center for Vision Research, Massachusetts Eye and Ear Infirmary, Boston, MA. Sequence data were aligned with sequence analysis software (Chromas, ver. 2.33; Tewantin, QLD, Australia) and compared with the published gene sequence by an NCBI BLAST search (www.ncbi.nlm.nih.gov/blast/ provided in the public domain by NCBI, Bethesda, MD).

Immunohistochemistry

The avidin-biotin-peroxidase complex and 3,3-diaminobenzidine tetrahydrochloride (ABC-DAB) procedure to determine Stat3 expression was performed on retinal sections from normal and elevated-IOP eyes. The 16-µm-thick cryostat sections were thawed and air dried for 10 minutes at room temperature. The sections were incubated in 0.3% H₂O₂ for 10 minutes, washed in 0.1 M phosphate-buffered saline (PBS), blocked, and incubated with Stat3 (1:100; Cell signaling) for 48 hours. Biotinylated anti-rabbit IgG was used as the secondary antibody. After they were washed with PBS, the sections were treated in ABC (Vectastain Elite ABC kit, Vector Laboratories, Burlingame, CA) solution according to the instructions of the supplier. After ABC treatment and washing in PBS, the slides were treated with a DAB Kit (Vector Laboratories) for 2 to 3 minutes. The reaction was terminated by rinsing in water. After several washes in distilled water, the slides were dehydrated in ethanol, cleared in xylene, and coverslipped (Permount; Fisher Scientific, Pittsburgh, PA).

Immunofluorescence staining for Stat3 and phosphorylated-Stat3 was also performed on retinal sections from normal and elevated-IOP eyes. Sections were incubated with a blocking solution for 1 hour at room temperature and incubated with primary antibodies specific for Stat3 (1:400, Cell Signaling, Beverly, MA) or phosphorylated-Stat3 (1:250, cell signaling) at 4°C overnight. Anti-goat antibodies (Alexa 594; Molecular Probes, Eugene, OR) were used as secondary antibodies with 1-hour incubation at room temperature. Fluorescence staining was detected with a fluorescence microscope (Olympus, Tokyo, Japan).

RESULTS

Evaluation of the High-IOP-Induced Glaucoma Model

To quantify glaucoma progression, we calculated the integrated IOP (AUC), assessed RGC loss, and graded optic nerve

TABLE 1. IOP Characteristics of the Rats

		Ave	erage IOP*	Pe	eak IOP*
Eye Pair	Total AUC	Control	Experimental	Control	Experimental
1†‡	155	19.13	29.96	19.30	43.69
2†‡	195	17.78	34.73	18.4	44.87
3†‡	169	16.77	30.94	19.47	45.06
4‡	209	18.78	32.30	20.53	43.73
5‡	150	19.13	31.90	19.93	41.13

AUC, area under the time-pressure curve, used to assess exposure to IOP.

^{*} Both the average IOP and the peak IOP of experimental group are significantly higher than the control group (P < 0.0001).

[†] Eye pairs 1-3 used for microarray experiment.

[‡] Eye pairs 1-5 used for RT-qPCR confirmation.

TABLE 2. Primers Used for RT-qPCR

Gene Name	GenBank*				Pri	mers	Sequ	ience				Amplicon Size (bp)
C1qa: complement component 1, q subcomponent,	NM_001008515	F:	TTG		GAA			GCC				207
alpha polypeptide C1qb: complement component 1, q subcomponent, beta polypeptide	NM_019262	F:	CTT	GGG TTT		GCT	CCA	ACC TGT CCC	CTC	GGG CT	Ġ	213
C3: complement component 3	NM_016994	F:	CTC	TGG	GGA CCA	GAA	AAG		AAT	-	CCC	97
Cpg15: candidate plasticity gene 15	NM_053346	F:	TAT	CCA	AGG CTT		CTT		CG	1110	000	200
Gapdb: glyceraldehyde-3-phosphate dehydrogenase	NM_017008	F: R:		ATG	TAT AGG	CCG	TTG	TGG	AT			177
Gfap: glial fibrillary acidic protein	NM_017009	F: R:			CCA TCC	AAT TTG				AAG TTC		117
Grb2: growth factor receptor bound protein 2	NM_030846	F: R:	AAG GGT					GGA GTG		GG		180
Map2k1: mitogen activated protein kinase kinase 1	NM_031643	F: R:	TTG GTT		CAA AGG			AAG ATC		GCT GTG		135
Nrg1: neuregulin 1	NM_031588	F: R:	GGT GCT		GGA GCC					G		218
Stat3: signal transducer and activator of transcription 3	NM_012747.2	F: R:	TTA ATT	CCC	ACA	TCT	CTG	CTC	CC	GTG	С	183
Stat5a: signal transducer and activator of transcription 5a	_	F: R:	CAG ATC	TGC	ACC TGC	CGG	ATC TTC	CTG TGC	G			153
Thy1: thymus cell antigen 1, theta	NM_012673	F: R:	ACA GCT	TGT TAT					GAC	000	G	99
Ywbab: tyrosin 3-monooxygenase/tyyptophan 5- monooxygenase activation protein, beta polypeptide	NM_019377	F: R:	TCC TTC	TCT	AAA CCC	TCT	CCA	GCA	TCT	GG CC		184
Ywbaz: tyrosin 3-monooxygenase/tyyptophan 5- monooxygenase activation protein, zeta polypeptide	NM_013011	F: R:	ATC TGC		CTG GTC	GGT TCC		GCC GGT				216

^{*} http://www.ncbi.nlm.nih.gov/Genbank; provided in the public domain by the National Center for Biotechnology Information, Bethesda, MD.

damage. Table 1 shows the data for all samples processed for microarray and RT-qPCR. Animals with similar IOP histories were chosen. The average RGC loss was 33%. Two available optic nerves were graded at 3.49 and 3.50, suggesting moderate retina damage (scale: 1, no damage; 5, severe damage).

RGC Enrichment

RGCs were identified by the presence of Fluorogold backlabeling and captured for microarray analysis. Because astrocytes and amacrine cells occupy the ganglion cell layer along with RGCs, we sought to assess the efficiency of LCM in enriching each sample for RGCs. We determined the specific mRNA expression of *Gfap* (astrocyte marker) and *Tby1* (RGCs marker) between LCM-captured RGCs and LCM-captured GCL (Table 3). Compared with LCM-captured GCL, *Gfap* was 65.04fold less abundant in LCM-captured RGCs (P = 0.006). No significant differences in Thy-1 levels were found between LCM captured-RGCs and LCM-captured GCL (P = 0.07). In addition, our microarray findings showed that the transcriptional expression of macrophage markers (CD14, CD68, and epidermal growth factor module-containing mucin-like receptor 1 [EMR1]) and an amacrine cell marker (HPC-1/syntaxin 1A) were not significantly different between normal and glaucomatous RGCs. The change ratios of CD14, CD68, EMR1, and syntaxin 1A were 1.23, 1.04, -1.12, and 1.14, respectively.

These results indicate that although some astrocytes were captured along with the RGCs, Fluorogold labeling effectively guided LCM harvesting of the RGCs. LCM capture successfully captured RGCs and effectively excluded most glial cells, macrophages, and amacrine cells.

Identification of Differentially Expressed Genes

Microarrays were used to survey the change in gene expression of RGCs from normal eyes and high-IOP eyes. A total of 1532 genes were significantly changed (Supplementary Table S1, http://www.iovs.org/cgi/content/full/51/8/4084/DC1). After FDR correction for multiple testing, 905 genes remained significant with a change of twofold or more in expression, with 330 genes increasing and 575 genes decreasing in glaucomatous RGCs (Supplementary Table S2, http://www.iovs. org/cgi/content/full/51/8/4084/DC1). Analysis of the main gene ontology categories revealed that the genes involved in apoptosis and response to oxidative stress were among the most upregulated (Table 4), including Angptl4, Ednrb, Pnkp, Gpx1, Ca3, RT1-Aw2, and Nek6. On the other hand, the genes vital for neuron, axon, and dendrite development were among the most downregulated (Table 5), which include Gja1, Snap25, Cit, Nrep, Gabrg2, and Vdac3.

The expression of several prosurvival genes was altered in glaucomatous RGCs. Several prosurvival genes that are present

TABLE 3. The mRNA Expression of *Gfap* and *Tby1* among LCM-Captured RGC and GCL*

Marker	LCM RGC $(n = 3)$	LCM GCL $(n = 3)$	P	GCL:RGC Ratio
Gfap	0.49 ± 0.05 47.00 ± 1.97	31.87 ± 4.79	0.006	65.04
Thy 1		29.13 ± 4.86	NS	0.62

^{*} The relative expression level (mean \pm SD) of each gene was calculated based on the standard curve, with *Gapdb* used as a reference. *Gapdb* did not change significantly in the samples.

TABLE 4. The Thirty Most Upregulated Genes in RGCs with Elevated IOP

GenBank*	Gene Name	Change Ratio	GO Biological Process
NM_017009	Gfap: glial fibrillary acidic protein	19.9	Intermediate filament-based process
AA945643	Chi311: chitinase 3-like 1	10.4	Carbohydrate metabolic process
AI176253	Nyw1: Ischemia related factor NYW-1	8.17	Unknown
NM_021584	Ania4: activity and neurotransmitter-	6.38	Unknown
021901	induced early gene protein 4 (ania-4)	0.50	
NM_022512	Acads: acetyl-Coenzyme A dehydrogenase, short chain	5.51	Metabolic process, response to glucocorticoid stimulus, oxidation reduction
NM_013154	Cebpd: CCAAT/enhancer binding protein (C/EBP), delta	4.67	Transcription, regulation of transcription, DNA-dependent
AI170182	Man1a2_predicted: mannosidase, alpha, class 1A, member 2 (predicted)	4.07	Unknown
NM_031140	Vim: vimentin	3.86	Intermediate filament-based process
BF415939	Fos: FBJ murine osteosarcoma viral oncogene homolog	3.83	Regulation of transcription, DNA-dependent, nervous system development
BG380285	Iftm1_predicted: interferon induced transmembrane protein 1 (predicted)	3.81	Response to biotic stimulus
AA818262	Angptl4: angiopoietin-like 4	3.75	Angiogenesis, response to hypoxia, signal transduction, negative regulation of apoptosis
X57764	Ednrb: endothelin receptor type B	3.71	Neural crest cell migration, protein amino acid phosphorylation, signal transduction, apoptosis, pigmentation
NM_016994	C3: complement component 3	3.66	GPCR protein signaling pathway, complement activation, inflammatory response
AW520745	Pnkp: polynucleotide kinase 3'- phosphatase	3.69	DNA repair, response to oxidative stress, dephosphorylation, nucleotide phosphorylation
AI317842	Kif22: kinesin family member 22	3.60	Microtubule-based movement
S41066	<i>Gpx1:</i> glutathione peroxidase 1	3.54	Release of cytochrome <i>c</i> from mitochondria, regulation of inflammatory response to antigenic stimulus, metabolic process, regulation of caspase activity, regulation of neuronal apoptosis
AB030829	Ca3: carbonic anhydrase 3	3.48	Metabolic process, response to oxidative stress
AI070137	Aoc3: amine oxidase, copper containing 3	3.43	Cell adhesion, amine metabolic process
BE102350	RT1-Aw2: RT1 class Ib, locus Aw2	3.31	Apoptosis, inflammatory response, calcium-mediated signaling, regulation of synapse structure and activity, regulation of axonogenesis
NM_019144	Acp5: acid phosphatase 5, tartrate resistant	3.30	Bone resorption
BI300770	Igtp: interferon gamma induced GTPase	3.30	Unknown
NM_019358	Pdpn: podoplanin	3.30	Cell morphogenesis, prostaglandin metabolic process, transport, signal transduction, cell proliferation
NM_031832	<i>Lgals3:</i> lectin, galactose binding, soluble 3	3.29	Extracellular matrix organization and biogenesis
BF411958	Rufy1: RUN and FYVE domain containing 1	3.20	Protein transport, regulation of endocytosis
AI407092	<i>Rdx</i> : Radixin	3.18	Microvillus biogenesis, apical protein localization
BF281577	Rcn1_predicted: Reticulocalbin 1 (predicted)	3.17	Ciliary or flagellar motility
AI072892	Frzb: frizzled-related protein	3.10	Regulation of Wnt receptor signaling pathway
NM_133609	Eif2b3: eukaryotic translation initiation factor 2B, subunit 3 gamma	3.07	Regulation of Rho protein signal transductioniosynthetic process, oligodendrocyte, Regulation of translational initiation in response to stress
BE113920	Stat3: signal transducer and activator of transcription 3	3.06	initiation in response to stress Eye photoreceptor cell differentiation, transcription, acute-phase response, signal transduction, JAK-STAT cascade
BF282365	<i>Nek6:</i> NIMA (never in mitosis gene a)-related expressed kinase 6	2.90	Protein amino acid phosphorylation, apoptosis, regulation of І-кВ kinase/NF-кВ cascade

 $^{^*\} http://www.ncbi.nlm.nih.gov/Genbank; provided in the public domain by the National Center for Biotechnology Information, Bethesda, MD.$

in normal RGCs were significantly downregulated in RGCs from eyes with high IOP (Table 6), including *Cpg15*, *Nrg1*, *Map2k1*, *Ywhab*, *Ywhaz*, *Vsnl1*, *Nfl*, *Stmn2*, *Nell2*, and *Grb2* (P < 0.05). On the other hand, the message levels of two prosurvival genes, *Stat3* (3.06 \pm 0.53, P < 0.05) and *Stat5a*

(2.26 \pm 0.55, P < 0.05), were increased in RGCs in response to elevated IOP.

Two methods of biological pathway analysis were used to further analyze our data. First, pathway analysis (IPA; Ingenuity) yielded distinct canonical pathways related to the RGC

TABLE 5. The Thirty Most Downregulated Genes in RGCs with Elevated IOP

GenBank*	Gene Name	Change Ratio	GO Biological Process
AI411352	Gja1: gap junction membrane channel protein alpha 1	-10.46	Embryonic development, neuronal migration, cell communication, cell-cell signaling, positive regulation of I-κB kinase/NF-κB cascade, neurite morphogenesis, protein oligomerization, regulation of calcium ion transport
D13921	Acat1: acetyl-coenzyme A acetyltransferase 1	-10.26	Metabolic process
NM_030991	Snap25: synaptosomal-associated protein 25	-9.16	Exocytosis, neurotransmitter secretion, axonogenesis, regulation of synaptogenesis
AI411995	Ube2d3: Ubiquitin-conjugating enzyme E2D 3 (UBC4/5 homolog, yeast)	-9.00	Ubiquitin-dependent protein catabolic process
AI717472	<i>Tyrp1:</i> tyrosinase-related protein 1	-8.07	Melanin metabolic process, pigmentation
AA957183	Cit: Citron	-7.89	Mitotic sister chromatid segregation, dendrite development, generation of neurons
BE105102	Glrx2: Glutaredoxin 2 (thioltransferase)	-7.74	Transport, response to organic substance, oxidation reduction
AW529483	Slc4a5: Solute carrier family 4, sodium bicarbonate cotransporter, member 5	-7.67	Unknown
NM_057205	Ube1c: ubiquitin-activating enzyme E1C	-6.73	Mitotic cell cycle, protein modification process, metabolic process, regulation of cell cycle
AA851874	Spg3a: spastic paraplegia 3A homolog (human)	-6.64	Unknown
BG380454	Basp 1: brain abundant, membrane attached signal protein 1	-6.45	Unknown
BE104618	Dbrs1: Dehydrogenase/reductase (SDR family) member 1	-6.34	Metabolic process
BE107450	Nrep: Neuronal regeneration related protein	-5.95	Regulation of TGF β receptor signaling pathway, axon regeneration, regulation of neuronal differentiation
BG372673	Trim32: tripartite motif protein 32	-5.92	Unknown
AI408819	H3f3b: H3 histone, family 3B	-5.91	Nucleosome assembly
BE120391	Gabrg2: gamma-aminobutyric acid A receptor, gamma 2	-5.79	Transport, synaptic transmission, post-embryonic development
AI169170	Eif4a2: eukaryotic translation initiation factor 4A2	-5.68	Translation
NM_013128	Cpe: carboxypeptidase E	-5.61	Proteolysis, insulin processing
BF284830	Yipf3: Yip1 domain family, member 3	-5.53	Cell differentiation
NM_013219	Cadps: Ca2+-dependent secretion activator	-5.24	Transport, vesicle fusion, positive regulation of calcium ion-dependent exocytosis
AI029445	Ldb2_predicted: LIM domain binding 2 (predicted)	-5.17	Multicellular organismal development
AI172276	<i>Ppp1r2:</i> protein phosphatase 1, regulatory (inhibitor) subunit 2	-5.13	Metabolic process, nervous system development, regulation of signal transduction, regulation of phosphoprotein phosphatase activity
BM388861	Col9a1: procollagen, type IX, alpha 1	-5.13	Chondrocyte differentiation, phosphate transport
NM_053562	Rpe65: retinal pigment epithelium 65	-5.07	Eye development, regulation of rhodopsin gene expression, retinal metabolic process
NM_057104	Enpp2: ectonucleotide pyrophosphatase/phosphodiesterase 2	-5.05	Chemotaxis, metabolic process, lipid catabolic process
NM_013011	Ywbaz: tyrosin 3-monooxygenase/tyyptophan 5- monooxygenase activation protein, zeta polypeptide	-5.03	Anti-apoptosis, protein targeting, signal transduction
AI145413	Gabra1: gamma-aminobutyric acid A receptor, alpha 1	-4.94	Transport, gamma-aminobutyric acid signaling pathway development
NM_031825	Fbn1: fibrillin 1	-4.91	F
AF439397	Zwint: ZW10 interactor	-4.90	Exocytosis, cell cycle, mitosis, cell division

^{*} http://www.ncbi.nlm.nih.gov/Genbank; provided in the public domain by the National Center for Biotechnology Information, Bethesda, MD.

response to elevated IOP, including cardiac β -adrenergic signaling, interferon signaling (the JAK/STAT pathway exists within the interferon signaling pathway), antigen presentation pathway, lysine biosynthesis, glutamate receptor signaling, cAMP-mediated signaling, chemokine signaling, leukocyte extravasation signaling, 14-3-3-mediated signaling, synaptic long-term potentiation, and G-protein-coupled receptor signaling (Table 7).

Second, GSEA demonstrated significant enrichment in complement coagulation cascades, Fas pathway, prostaglandin, and leukotriene metabolism, JAK/STAT signaling pathway, G-protein-coupled receptor database class (GPCRDB) A rhodopsin-like, prostaglandin synthesis regulation, and caspase pathway

expression, whereas significant depletion in the citrate cycle, cholesterol biosynthesis, and G-protein-coupled receptor (GPCR) pathways was found in RGCs from high-IOP eyes (Table 8).

Validation of Selected Differentially Expressed Genes by RT-qPCR

To confirm selected microarray results, we examined expression levels by RT-qPCR after LCM in five pairs of eyes. Based on a combination of statistical analysis of microarray data and potential biological importance of the genes in question, 11 genes were chosen for RT-qPCR confirmation. These 11 genes

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TABLE 6. Prosurvival and Neurogenesis Genes with Decreased Expression in Glaucomatous RGCs

Gene	Function	Glaucomatous RGCs/Normal RGCs (Change Ratio)
Stmn2	Microtubule depolymerization and synaptic plasticity	-6.26*
Ywbaz	Mitochondrial import stimulation factor	-5.03*
Vsnl1	Calcium dependent signaling	-3.45*
Ywhab	Signal transduction	-3.33*
Ret	Neurogenesis	-2.72
Cpg15	Promotes neurite outgrowth	-2.61*
Nfl	Cytoskeleton organization	-2.45^{*}
Grb2	Signal transduction	-2.34^{*}
Nell2	Growth and differentiation of neural cells	-2.25*
Calb2	Calcium ion binding	-2.08
NCAML1	Neural and glial cell adhesion	-2.08
Map2k1	Signal transduction	-2.06*
Nrg1	Neural and organ development	-2.06*
Nrp	Neuronal development	-2.01
Rgs4	G-protein coupled receptor signaling	-2.01
Ywhag	Signal transduction	-1.74
Coch	Perception of sound	-1.68
Pik3r1	Negative regulation of apoptosis	-1.56
Plcb1	Production of the second messenger molecules	-1.56
Gap43	Neurite outgrowth and nerve regeneration	-1.47
Rgs3	G-protein coupled receptor signaling	-1.47
Nef3	Cytoskeleton organization	-1.4
Rnd1	Neuritic process formation	-1.4
Inpp4a	Phosphatidylinositol-3,4-bisphosphate 4-phosphatase activity	-1.38
Ywhah	Negative regulation of apoptosis	-1.26
Sema6b	Neurogenesis and axon guidance	-1.22
Prph1	Cytoskeleton organization	-1.17
Fbxo2	Maintaining neurons in a postmitotic state	-1.15
Ptpn5	Regulation the duration of ERK activation	1.12

^{*} P < 0.05.

represent the survival genes (Ywbaz, Ywbab, Cpg15, Grb2, Nrg1, Map2k1, Stat3, and Stat5a) and complement component genes (C3, C1qa, and C1qb). The change obtained by RT-qPCR correlated well with the direction of change obtained from the microarray data, confirming the validity of the microarray gene expression patterns (Table 9). Because RT-qPCR is more sensitive than microarray, RT-qPCR identified greater changes than microarray for these genes. In addition, several genes such as Stat5a and Ywbab were absent in some chips. RT-PCR confirmed the presence of these genes in both normal

TABLE 7. Top Canonical Pathways that Changed Significantly in RGC from Elevated IOP Eyes

Pathway	P^*	Ratio (%)†
Cardiac β-adrenergic Signaling	< 0.0001	33.60
Interferon Signaling (includes JAK/STAT)	0.001	43.30
Antigen presentation pathway	0.001	30.80
Lysine biosynthesis	0.011	7.81
Glutamate receptor signaling	0.018	29.90
cAMP-mediated signaling	0.025	29.20
Chemokine signaling	0.026	31.60
Leukocyte extravasation signaling	0.030	24.5
14-3-3-mediated signaling	0.035	28.20
Synaptic long term potentiation	0.035	29.5
G-Protein coupled receptor signaling	0.039	27.10

^{*} The P-value suggests whether there is an association between a specific pathway and uploaded dataset and whether it is due to chance.

RGCs and glaucomatous RGCs. Direct sequencing confirmed the identity of all the genes selected to validate microarray

Upregulated Stat3 and Phosphorylated-Stat3 Protein in Glaucomatous RGCs

Stat3 was detected in the ganglion cells layers and inner nuclear layer (Fig. 1A) as previously described.²⁴ Under conditions of high IOP, increased Stat3 immunoreactivity was seen in the GCL, inner nuclear layer, and inner plexiform layer, as well as in the cell processes (Fig. 1B). To further localize the increases in Stat3 and test whether Stat3 is activated in glaucomatous RGCs, the RGCs were back-labeled with Fluorogold, and the sections were fluorescently immunostained with Stat3 and phosphorylated-Stat3 antibodies. Figure 1F shows colocalization of Fluorogold and Stat3 in normal retina and Figure 1I shows increased Stat3 in an RGC under conditions of high IOP. Phosphorylated-Stat3 is expressed at low levels in normal RGCs (Fig. 1K), whereas it is increased in glaucomatous RGCs (Figs. 1N, 1O).

DISCUSSION

Apoptosis is thought to be an acute or subacute event, yet in glaucoma, it takes weeks to years of elevated IOP to trigger RGC death. In an attempt to better understand upstream IOPinduced changes that may contribute to RGC dysfunction or apoptotic degeneration, we studied RGCs in animals exposed to IOP for 10 days, at a time point at which some, but not all, RGCs were likely to have died. Since RGCs account for only a small percentage of the cells in the retina and since it is these cells that are uniquely vulnerable to the consequences of ele-

[†] The ratio is the number of molecules in a given pathway that meet cutoff criteria (twofold change), divided by total number of molecules that make up that pathway. The ratio is good for looking at which pathway has been affected the most based on the percentage of genes uploaded into IPA.

TABLE 8. Gene Sets Enriched/Depleted in Glaucomatous RGC*

Gene Set	Enrichment Score†	False-Discovery Rate q‡
Enriched in glaucomatous RGCs		
Complement coagulation cascades	0.45	< 0.0001
Fas pathway	0.50	< 0.0001
Prostaglandin and leukotriene metabolism	0.44	< 0.0001
JAK/STAT signaling	0.42	< 0.0001
G protein coupled receptor database class A rhodopsin-like	0.35	< 0.0001
Prostaglandin synthesis regulation	0.47	< 0.0001
Caspase pathway	0.47	< 0.0001
Depleted in Glaucomatous RGC		
Citrate cycle	0.55	< 0.0001
Cholesterol biosynthesis	0.81	< 0.0001
G-protein-coupled receptor pathway	0.57	< 0.0001

^{*}A total of 639 canonical pathways gene sets (http://www.broad.mit.edu/gsea/msigdb/genesets.jsp?collection=CP#; provided in the public domain by the Broad Institute, Massachusetts of Technology, Cambridge, MA) were calculated by gene set enrichment analysis.

vated IOP in glaucoma, we developed a technique using LCM to isolate RGCs for microarray analysis.

Our work had several novel outcomes. In studies of changes in transcript profiles in experimental glaucoma, whole retina was used and a relatively modest number and diversity of gene expression changes were recorded. In comparison with these studies, we found detectable changes in nearly 1000 transcripts: 330 upregulated and 575 downregulated. The pattern of these changes highlights and reinforces several broad themes, including downregulation of many neuronal prosurvival genes and upregulation of apoptosis-related pathways and markers of inflammation, including complement pathways. We describe, for example, gene expression changes in Stat3 and extend these results to include evidence of increased protein levels of Stat3. In addition, we demonstrated an increase in Stat3 phosphorylation, which leads to the activation of Stat3 and may be an important prosurvival mediator. In addition, several unexpected but robust changes in patterns of gene expression emerged, including alterations in the GPCRs. These changes suggest new therapeutic targets for neuroprotection in glaucoma distinct from direct pressure-lowering approaches. The large number of individual transcripts and pathways altered in RGCs by exposure to elevated IOP suggest that the pathophysiological processes that ultimately are manifest by RGC degeneration are proceeded by a complex, likely integrated, set of changes in gene expression. The evidence implies that elevated IOP initiates transcriptional cascades that most likely contribute to neurodegeneration.

RGCs exist in a complex milieu and receive input from multiple different cell types, any of which may be influenced by elevated IOP. It is therefore important to study RGCs in situ to truly understand gene expression changes as a result of increased IOP. Whole retinal PCR is frequently used to study retinal gene responses to increased IOP. However, this technique does not differentiate responses of RGCs from other cell types. LCM, coupled with microarray analysis, is ideally suited to this challenge. Fluorogold back-labeling can effectively guide microscopic laser capture of neurons, as has been reported. In the present study, RGCs were identified by the presence of Fluorogold back-labeling and captured by LCM. Our results indicate that some astrocytes were probably captured along with the RGCs. Similar findings have been reported

TABLE 9. Validation of Microarray Gene Expression by Real-Time Quantitative PCR

	Ratio of Change	e (Mean ± SD)
Gene	Microarray $(n = 3)$	RT-qPCR $(n = 5)$
Ywbaz: 14-3-3 family, zeta polypeptide	$-5.03 \pm 0.89^*$	$-25.77 \pm 5.42^*$
Ywhab: 14-3-3 family, beta polypeptide	$-3.33 \pm 0.55^*$	-23.14 ± 6.26 *
<i>Cpg15:</i> candidate plasticity gene 15	$-2.61 \pm 0.30^*$	$-20.93 \pm 9.79^*$
<i>Grb2:</i> growth factor receptor bound protein 2	$-2.34 \pm 0.32^*$	$-24.59 \pm 10.43^*$
Nrg1: neuregulin 1	$-2.06 \pm 0.30^*$	$-6.56 \pm 0.47^*$
Map2k1: mitogen activated protein kinase kinase 1	$-2.06 \pm 0.23^*$	$-9.73 \pm 3.43^{*}$
<i>C3:</i> complement component 3	$3.66 \pm 1.01^*$	$5.07 \pm 1.55^*$
C1qa: complement component 1, q		
subcomponent, alpha polypeptide	1.39 ± 0.32	1.89 ± 0.45
Stat3: signal transducer and activator of		
transcription 3	$3.06 \pm 0.53^*$	$7.28 \pm 1.27^*$
Stat5a: signal transducer and activator of		
transcription 5a	$2.26 \pm 0.55^*$	$2.74 \pm 0.90^*$
C1qb: complement component 1, q		
subcomponent, beta polypeptide	1.28 ± 0.11	$2.13 \pm 0.56^*$

^{*} P < 0.05.

[†] Enrichment score (ES) reflects the degree to which a gene set is overrepresented at the top or bottom of a ranked list of genes.

[‡] False-discovery rate is the estimated probability that a gene set with a given normalized ES represents a false-positive finding.

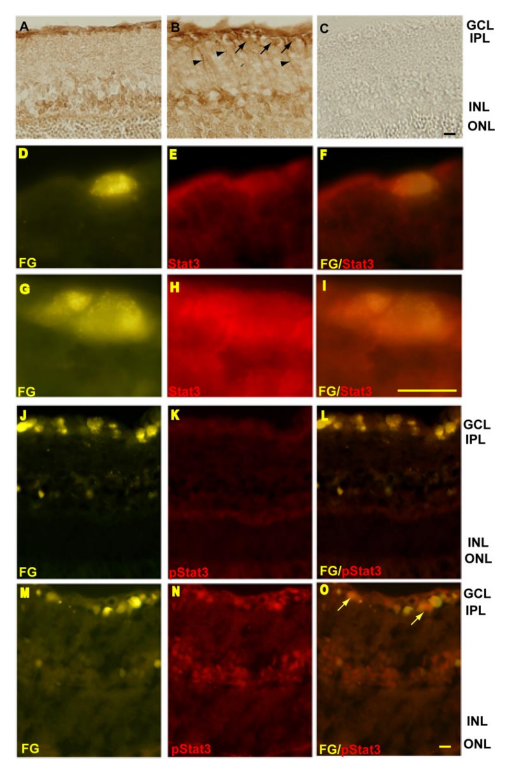


FIGURE 1. Stat3 protein increased in the glaucomatous retina and in glaucomatous RGCs, and phospho-Stat3 was also increased in glaucomatous RGCs. (A) In the normal retina, positive staining for Stat3 was seen in the inner nuclear layer (INL) and GCL. (B) The expression of Stat3 was higher in the GCL in the high-IOP retina. (C) Negative control. (E) Low levels of Stat3 immunoreactivity were seen in Fluorogold-labeled RGCs in a retina with normal IOP. (G-I) In retinas with high IOP, increased Stat3 expression was seen in Fluorogold-labeled RGCs. (K-N) Phosphorylated-Stat3 was clearly increased in the GCL, including in RGCs (O compared with L). FG, Fluorogold. Arrow: retinal ganglion cell; Arrowbead: cell processes. Scale bars: 20 μm.

for LCM of substantia nigra neurons in which oligodendrocyte genes were found in the microarray results in histologic sections of a thickness similar to that of those used in our studies. ²⁵ Yao et al. ²⁵ suggest that the co-capture of neurons along with the surrounding glial cells may underscore the close association of these two cell types. Although we cannot exclude the possibility that some of the pathways identified as changing are changing in glial cells (astrocytes and/or Müller cell processes), in our control experiments we found that expression of the glial cell gene *Gfap* was 65.04-fold less

abundant in LCM-captured RGCs than in LCM of the entire GCL. These results indicate that LCM results in a highly enriched sample of RGCs.

In this article, we report data from three pairs of eyes. Because of the small sample, we may have underestimated the actual changes, although publications of studies in which a similar number of paired samples was used have reported important data in other systems. ²⁶ qRT-PCR of independently laser-captured RGCs from five pairs of eyes confirmed the direction of change found in the microarray data in all cases.

TABLE 10. Comparison of Our Study with Previous Microarray Studies Using Whole Retina Glaucoma Model

	Our Study	Miyahara et al. ⁷	Ahmed et al. ⁸	Piri et al. ⁹	Steele et al. ⁶	Naskar et al. ¹⁰	Yang	Yang et al. ¹¹	Walsh et al. ¹²
Species Glaucoma model	Brown Norway rat Saline injection	Cynomolgus monkey Laser-induced glaucoma	Brown Norway rat Saline injection	Wistar rat Optic nerve transsection	DBA2J mouse 8 months after IOP elevation	Rat Hereditary high IOP	Wista Laser-induced glaucoma	Wistar rat Optic nerve transsection	C57 Mice Saline injection
Samples Array*	RGC Affymetrix rat 230.2	Whole retina UniGEM Human ver 2.0	Whole retina Affymetrix rat U34A	Whole retina Aligent rat G4130A	Whole retina Affymetrix mouse 430.2	Whole retina MWG Biotech 10K chip	Whole	Whole retina Affymetrix rat 230.2	Whole retina Affymetrix mouse
Statistic setting	>2-fold change	>1.7-fold change	>2-fold change	>2-fold change	>1.8fold change	>3-fold change	>2-fold	>2-fold change	>1.5-fold
Number of upregulated genes	330	45	7.4	85	32	27	347	212	50 50
Number of downregulated genes	575	17	٢	21	36	45	196	480	140
Major upregulated genes	Apoptosis genes (Table 4), Complement component genes (CIqa, CIqb, C3)	Immune response genes, gla activation genes, ceruloplasmin	Immune response genes, glia activation genes	Apoptotic genes, ceruloplasmin	Gilal activation, immune response genes, signaling, gene expression	Glial activation genes	Early phase: immune and stress response genes. Late phase: genes related to protein synthesis	Early phase: genes related to cell death	Saa1, Saa2
Major downregulated genes	Prosurvival genes (Gpg 15, Nrg1, Yurbaz, Yurbaz, Vurbaz, NJ, Stmt., Map2kt, Netl.2, Vstt.D., neurogenesis genes (Gja 1, Snap25, Gt, Nrep, Gabry2, and Varies)	Neurofilament genes	Neurofilament genes	Neurofilament genes, Neuritin (Cpg15)	Multiple crystal genes			RGC-enriched genes; neurofilament genes	

* Affymetrix, Santa Clara, CA; Agilent, Palo Alto, CA; UniGem, Incyte Genomics, Palo Alto, CA; MWG Biotech, High Point, NC.

Microarray analysis of isolated RGCs refines detection of gene responses to increased IOP. The current microarray study of RGC RNA yielded 905 genes with altered expression, many more than the number of significantly changed genes found in whole retina analyses from different glaucoma models (Table 10).6-12 Among the different microarray studies, some common themes were shared. The immune response genes were upregulated in the Morrison rat model,⁸ laser-induced monkey or rat model,^{7,11} and DBA2J mice.6 In the present study, complement component genes were enriched in glaucomatous RGCs confirming in situ hybridization observations by Stevens et al.²⁷ A decrease in neurofilament genes has been reported in the monkey and the rat model of glaucoma^{7-9,11} and in our study, mRNA levels for Nefh, Nefm, and Nefl were decreased specifically in RGCs in response to elevated IOP. In addition, other structural genes vital for the survival or development of neuronal structures (Gja1, Snap25, Cit, Nrep, Gabrg2, and Vdac3) were among the most downregulated (Table 5). Genes vital for neuronal structure are affected by increased IOP, and these changes may contribute to RGC vulnerability in glaucoma or altered cellular functions that precede RGC death. Because of the detection ability of microarrays, some genes with small changes in expression may not have been detected in our microarray experiment. For example, Thy 1 was not significantly changed in our microarray analysis of glaucomatous RGCs. We have reported that Thy1 mRNA levels were decreased approximately twofold in glaucomatous RGCs isolated by LCM followed by RT-PCR.²⁰ Because RT-PCR has been shown to be more sensitive than microarrays,²⁸ some genes with small changes in expression may not be detected by microarray experiments.

In the present study, apoptosis and oxidative stress-related genes (*Angptl4*, *Ednrb*, *Pnkp*, *Gpx1*, *Ca3*, *RT1-Aw2*, *Nek6*, and *Stat3*) were among the most upregulated (Table 4). RGC death in glaucoma has been characterized as involving the apoptotic pathway of cell death.^{2,4,5} In addition, in the GSEA analysis, the caspase cascade was significantly enriched in glaucomatous RGCs, and one of the genes that contributed significantly to this enrichment was caspase 3. Also, growing evidence supports the possibility that oxidative stress is involved om glaucomatous neurodegeneration in different subcellular compartments of RGCs.²⁹ Our findings support the view that both apoptosis and oxidative damage contribute to glaucomatous RGC death.

The GPCRs are a diverse family of transmembrane signaling proteins. They transduce a wide variety of cellular stimuli, and a large number of the drugs in current use exert their effects through GPCRs. 30 In addition, multiple endogenous ligands are thought to act through GPCR pathways. Our data show an unexpected finding in glaucomatous RGCs of significant downregulation of the GPCR pathway. In addition, we found decreased expression of several adenylate cyclase genes and cAMP-dependent protein kinase (PKA) genes in RGCs from eyes with elevated IOP. It has been shown that GPCRs modulate neuronal signaling via specific G proteins in the central nervous system, including structures of the eye.31 In the present study, the decreased message levels of GPCR and G proteins in RGCs in glaucoma may lead to a decrease in adenylate cyclase-regulated cAMP signaling, which is critical for RGC survival. 32,33

Alterations in neurotrophins and neurotrophin receptors have been implicated in RGC death in glaucoma with a decrease in retinal levels of both the levels of BDNF and in the Trkb receptors.³⁴ Our observation that the GPCR pathway is downregulated prompts consideration of another intriguing possibility. Transactivation of the Trkb receptor through a GPCR-mediated, non-neurotrophin mechanism has been described,³⁵ and one possibility is that downregulation of the

GPCR pathway has the potential to cause a further decrease in Trkb signaling in RGCs in glaucoma.

In summary, in our work that elevated IOP led to complex changes in RGC gene expression in a rat model of glaucoma. A critical finding of this study is that multiple cellular pathways were changed in RGCs in response to elevated IOP. The expression of genes involved in apoptotic pathways (Caspase pathway, Fas pathway, JNK pathway) was increased, and the genes involved in the GPCR, 14-3-3 pathway, and PI3K pathway were decreased. By contrast, gene expression of the prosurvival JAK/STAT pathway was increased, suggesting the activation of endogenous neuroprotective mechanisms. We also found decreased expression of many prosurvival genes that are normally expressed in RGCs. Thus, gene expression changes led to an alteration of the balance between prosurvival and prodeath genes and pathways in glaucomatous RGCs. We speculate that the decreased expression of prosurvival genes contributes to RGC death in glaucoma. These findings warrant further investigation of the roles of these genes in glaucomatous RGC death.

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