Optic Nerve Head Astrocytes Display Axon-Dependent and -Independent Reactivity in Response to Acutely Elevated Intraocular Pressure

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PURPOSE. Optic nerve head (ONH) astrocytes provide support for axons, but exhibit structural and functional changes (termed reactivity) in a number of glaucoma models. The purpose of this study was to determine if ONH astrocyte structural reactivity is axon-dependent.

METHODS. Using rats, we combine retrobulbar optic nerve transection (ONT) with acute controlled elevation of intraocular pressure (CEI), to induce total optic nerve axon loss and ONH astrocyte reactivity, respectively. Animals were euthanized immediately or 1 day post CEI, in the presence or absence of ONT. ONH sections were labeled with fluorescent-tagged phalloidin and antibodies against β3 tubulin, phosphorylated cortactin, phosphorylated paxillin, or complement C3. ONH label intensities were quantified after confocal microscopy. Retrobulbar nerves were assessed for axon injury by light microscopy.

RESULTS. While ONT alone had no effect on ONH astrocyte structural orientation, astrocytes demonstrated significant reorganization of cellular extensions within hours after CEI, even when combined with ONT. However, ONH astrocytes displayed differential intensities of actin (phosphorylated cortactin) and focal adhesion (phosphorylated paxillin) mediators in response to CEI alone, ONT alone, or the combination of CEI and ONT. Lastly, label intensities of complement C3 within the ONH were unchanged in eyes subjected to CEI alone, ONT alone, or the combination of CEI and ONT, relative to controls.

CONCLUSIONS. Early ONH astrocyte structural reactivity to elevated IOP is multifaceted, displaying both axon dependent and independent responses. These findings have important implications for pursuing astrocytes as diagnostic and therapeutic targets in neurodegenerative disorders with fluctuating levels of axon injury.

Keywords: optic nerve head, astrocyte reactivity, axon, elevated intraocular pressure, glaucoma
through signaling and crosstalk between astrocytes and the axonal/extracellular environment at the site of injury. While astrocyte-axon communication and crosstalk has been well documented in several models, the role of astrocyte-axon communication in initiating astrocyte reactivity remains an open question. Local elimination of axons prior to inducing astrocyte reactivity in a model is one way to address this question.

Here, using an adult rat model, we induced complete optic nerve axon loss by retrobulbar optic nerve transaction (ONT). In addition, we used acute controlled elevation of IOP (CEI) to induce local ONH astrocyte reactivity. We have shown previously that optic nerve axons exhibit minimal observable injury immediately after CEI (day 0 and day 1 post-IOP elevation), while optic nerve axons exhibit essentially total degeneration 2 weeks post ONT as assessed by light microscopy. Features of ONH astrocyte reactivity, including structural reorientation of astrocyte extensions and changes in label intensities of actin, focal adhesion-, and complement-based mediators were studied in eyes after ONT alone, CEI alone, and combined ONT and CEI, and were compared with control eyes.

**Materials and Methods**

**Animals**

All animals were treated in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research and all experimental methods were approved by the Oregon Health & Science University Institutional Animal Care and Use Committee. Animals underwent unilateral procedures only. Thirty-nine 8- to 9-month-old male Brown Norway rats (350–400 g) were randomized and underwent either unilateral retrobulbar ONT (n = 8), CEI (n = 16), or ONT followed by CEI 2 weeks post ONT (n = 15). Only male rats were used in this study. In prior experiments it proved difficult to consistently achieve a stable plane of anesthesia for the prolonged periods necessary in female rats, and mortality rates were unacceptable (data not shown).

**Optic Nerve Transection (ONT)**

For animals undergoing ONT (n = 23), rats received systemic anesthesia with intraperitoneal injection of ketamine (37.5 mg/kg; JHP Pharmaceuticals, Rochester, MI, USA), xylazine (7.5 mg/kg; RXV, Greeley, CO, USA), and acepromazine maleate (1.5 mg/kg; VET ONE, Boise, ID, USA). Topical 0.5% proparacaine hydrochloride ophthalmic solution (Akorn, Lake Forest, IL, USA) was applied to the ocular and conjunctival surface unilaterally. During anesthesia, animals remained on a temperature-controlled blanket at 37°C to maintain body temperature. The lateral canthus was clamped with a curved hemostatic mosquito forceps, followed by a lateral canthotomy with Wescott scissors. A conjunctival peritomy was performed at the superior six clock hours of the limbus. The conjunctiva was retracted by an assistant to expose the superior rectus muscle insertion near the limbus. To gain access to the retrobulbar optic nerve space, the superior rectus muscle was cauterized with low-temperature cautery (Bovie Medical, Clearwater, FL, USA) and transected with Vannas scissors. Care was taken to avoid injury to the superotemporal and suponasal vortex veins exiting the sclera. The eye was then infero-oriented by grasping the remaining superior rectus muscle insertion using 0.2-mm toothed forceps. The junction of the superior optic nerve and the globe was exposed using a two-pronged instrument modified from a Lewis Lens Loop, with care to avoid any trauma or excess stretch to the optic nerve. Next, a 25-G microvitreoretinal blade was used to carefully puncture the superior optic nerve sheath, as far away from the globe as possible. This was followed by dissection of the superior optic nerve sheath, using Vannas scissors. After adequate optic nerve sheath opening was confirmed, one arm of the Vannas scissors (with sharp edge oriented superiorly) was carefully placed between the optic nerve and the inferior optic nerve sheath, followed by total axotomy as far posterior in the retrobulbar space as possible. The inferior optic nerve sheath and its vasculature was preserved without manipulation. The orbital contents and conjunctiva were carefully replaced in their original position, and the conjunctiva was re-approximated at the limbus using two interrupted 10-0 nylon sutures (Medtronic, Minneapolis, MN, USA). The periocular surface was cleaned and erythromycin ointment (Perrigo, Minneapolis, MN, USA) was applied to the ocular surface. Animals received postsurgical subcutaneous injections of 50 μL of 0.3 mg/mL buprenorphine (Buprenex; Reckitt Benckiser Pharmaceuticals, Richmond, VA, USA) for analgesia and were allowed to awaken from anesthesia on a warm platform. Animals were allowed to survive for 2 weeks prior to euthanization (ONT alone, n = 8) or CEI (combined ONT and CEI, see below).

**Controlled Elevation of Intraocular Pressure (CEI)**

Naive animals (n = 16) or a portion of the animals which had undergone ONT 2 weeks prior (n = 15), underwent unilateral CEI as previously described. Briefly, for IOP elevation, animals were maintained under general anesthesia using 2.25% isoflurane mixed with 100% oxygen, and kept at a temperature of 37°C using a water bath-heated temperature therapy pad. After topical 0.5% proparacaine application to the eye, a corneal incision was made using a 31-G needle with care to avoid iris or lens trauma. Next, the anterior chamber was cannulated through the corneal incision with a 2 cm length of 0.010” OD polyurethane tubing (Instech Labs, Plymouth Meeting, PA, USA) linked via larger tubing to a reservoir filled with pH-balanced saline solution (BSS Plus; Alcon Laboratories, Fort Worth, TX, USA). The saline reservoir was raised to produce a pressure of 60 mm Hg, as confirmed by a pressure sensor coupled with the infusion line (Harvard Apparatus, Holliston, MA, USA) and independent measurements every 30 minutes using a handheld tonometer (TonoLab; Icare Finland Oy, Espoo, Finland). Both eyes received topical proparacaine and topical BSS, alternating every 15 minutes to keep the ocular surface anesthetized and moist, respectively. Animals received a 1-mL subcutaneous injection of 0.9% saline every hour to maintain systemic hydration while under general anesthesia. After 8 hours of elevated IOP exposure, the saline reservoir was lowered to 20 mm Hg for 5 minutes, after which the cannula was removed and animals were sacrificed either immediately after IOP normalization (CEI day 0, n = 8 for CEI day 0 only, n = 8 for CEI day 0 + ONT), or allowed to survive for 1 day (CEI day 1, n = 8 for CEI day 1 only, n = 7 for CEI day 1 + ONT) prior to euthanization. For euthanization, animals were placed under deep sedation using 5% isoflurane anesthesia and underwent systemic perfusion-fixation via cardiac access with 1 L of buffered 4% paraformaldehyde over 20 minutes, as previously described.

**Axonal Injury Assessment**

For axonal injury assessment, animals were euthanized under isoflurane anesthesia and eyes were perfusion-fixed with buffered 4% paraformaldehyde as described above. Eyes were enucleated and retrobulbar optic nerves 1 mm distal to the
Axon-Dependent Astrocyte Reactivity

Immunofluorescence Labeling of Optic Nerve Tissue

Perfusion-fixed eyes (including the ONH) were cryopreserved in 15% sucrose/PBS, followed by 30% sucrose/PBS. The eyes were positioned for vertical longitudinal sectioning centered on the optic nerve in 1:1 solution of optimal cutting temperature support medium (Sakura Finetek, Torrance, CA, USA) and 30% sucrose/PBS. Eyes were then frozen in liquid isopentane cooled by liquid nitrogen, and cryosectioned (5-μm thickness) onto glass slides. The superior and inferior orientation of the ONH was determined based on the anatomic location of the central retinal vein, located just inferior to the ONH. Tissue sections were blocked with 1% bovine serum albumin (BSA) in PBS for 1 hour at room temperature. Actin filaments were labeled with 1 μg/mL tetramethylrhodamine (TRITC)-labeled phalloidin (Sigma, Sigma-Aldrich Corp., St. Louis, MO, USA) in 1% BSA/PBS at room temperature for 1 hour, followed by three cycles of wash with PBS at room temperature for 5 minutes. Tissue sections were co-labeled with primary antibodies in 1% BSA/PBS at 4°C overnight, using antibodies against axonal tubulin (TuJ1; mouse monoclonal against βIII tubulin, 1:500 dilution; Covance, Seattle, WA, USA), phosphorylated paxillin (p-paxillin; rabbit polyclonal against phosphorylated Tyr 118, 1:100 dilution; Abcam, Cambridge, MA, USA), phosphorylated cortactin (p-cortactin; rabbit polyclonal against phosphorylated Tyr 421, 1:500 dilution; EMD Millipore, Billerica, MA, USA), or complement C3 (B-9 mouse monoclonal against amino acids 541-640 of human C3, 1:100 dilution; Santa Cruz Biotechnology, Dallas, TX, USA). Nonspecific mouse or rabbit IgG antibodies (Sigma) were used at similar concentrations to the respective antibodies above for negative control sections. Sections were washed as above, followed by incubation with secondary Alexa 488-labeled goat anti-mouse/rabbit monoclonal antibodies (ThermoFisher Scientific, Waltham, MA, USA) in 1% BSA/PBS for 1 hour at room temperature. After washing as above, cell nuclei were stained with the mounting media (Prolong Gold with DAPI; ThermoFisher Scientific). All immunofluorescent study n’s represent individual animals as biological replicates (i.e., 1 ONH section from each animal was labeled and analyzed for each n). A single section was chosen as close as possible to the center of the ONH from each animal, to improve anatomic consistency between samples. To reduce any potential labeling discrepancies within each antibody condition, each immunolabeling set (including the same laser settings, magnification, exposure time, intensity range, and image capture size) was completed in a single run. Confocal images were imported directly into FIJI image analysis software (http://fiji.sc/Fiji; an open source image processing package based on the National Institute of Health, Bethesda, Maryland, USA software Imagej: http://imagej.nih.gov/ij/), and were analyzed as Z-stacks in a semiautomated fashion (including analyzing the entire Z-stack image at once without user manipulation, and using the same analysis settings for all images). ONH actin bundle orientation was determined using the Directionality plug-in in FIJI as previously described. Mean fluorescence pixel intensities for phosphorylated cortactin, phosphorylated paxillin, and complement C3 were calculated using FIJI software. Upon completion of image acquisition and analysis, the key for identifying which sample/data belonged to which treatment group was then used to group the samples. Statistical analysis was performed by 1- or 2-way ANOVA (GraphPad Prism software, La Jolla, CA, USA), followed by Sidak’s and/or Dunnett’s multiple comparisons testing. Data were log transformed when necessary to meet the assumption of normal variance prior to ANOVA analysis. All data generated or analyzed during this study are included in this published article.

RESULTS

Axon Degeneration Following IOP Elevation and Optic Nerve Transection

We confirmed that optic nerves of animals undergoing CEI exhibited no observable axon injury immediately following CEI relative to controls (Figs. 1A, 1B, 1D, 1E). In contrast, 2 weeks after ONT, optic nerves exhibited essentially total axon degeneration (Figs. 1C, 1F).

IOP Elevation Induces Similar Astrocyte Structural Remodeling in the Presence or Absence of Viable Optic Nerve Axons

In order to assess the structural response of ONH astrocytes to IOP elevation in the presence or absence of viable axons, we quantified the orientation of astrocyte extensions within the ONH. The ONH is rich in astrocytic actin filaments (Fig. 2A) that delineate astrocyte extensions, with high concordance with astrocyte-specific markers such as glial fibrillary acidic protein (GFAP) and aquaporin 4. As previously reported, control ONH astrocytes exhibit highly arranged, under confocal microscopy and camera settings (FV1000 microscope; Olympus, Center Valley, PA, USA; and an UplanFLN ×40/NA1.30 oil objective; Olympus) to be within the linear range of the camera and avoid over/undersaturation. These settings were then applied to all samples going forward to maintain consistent imaging. We have previously shown that ONH astrocyte extension rearrangement in response to CEI was more significant in the inferior ONH, relative to the superior ONH. In this study, we confirmed that the inferior ONH indeed showed more astrocyte extension rearrangement in response to CEI, relative to the superior ONH (data not shown). Thus, we present data from the inferior ONH in this study.

Confocal images of the inferior ONH (the anterior and posterior borders defined as 0 and 100 μm posterior to the termination of Bruch’s membrane, respectively) were obtained. Images were captured using FV10-ASW version 4.0 software (Olympus) with laser wavelength settings of 405, 488, and 559 nm at 1 μm/slice. Confocal microscopy image acquisition was performed in the same manner on all slides within an antibody condition (including the same laser settings, magnification, exposure time, intensity range, and image capture size). Captured images were analyzed as Z-stacks in a semiautomated fashion (including analyzing the entire Z-stack image at once without user manipulation, and using the same analysis settings for all images). ONH actin bundle orientation was determined using the Directionality plugin in FIJI as previously described. Mean fluorescence pixel intensities for phosphorylated cortactin, phosphorylated paxillin, and complement C3 were calculated using FIJI software. Upon completion of image acquisition and analysis, the key for identifying which sample/data belonged to which treatment group was then used to group the samples. Statistical analysis was performed by 1- or 2-way ANOVA (GraphPad Prism software, La Jolla, CA, USA), followed by Sidak’s and/or Dunnett’s multiple comparisons testing. Data were log transformed when necessary to meet the assumption of normal variance prior to ANOVA analysis. All data generated or analyzed during this study are included in this published article.
Astrocytes Display Differential Phosphorylation of Cytoskeletal Mediators in Response to Elevated IOP in the Presence or Absence of Viable Optic Nerve Axons

We have previously shown that the phosphorylation status of actin cytoskeletal and focal adhesion mediators within ONH astrocytes are significantly altered immediately after IOP elevation. Here, we asked if the phosphorylation status of ONH cortactin (involved in dynamic actin turnover) and paxillin (involved in focal adhesion dynamics) in response to IOP elevation, was affected by the presence or absence of viable axons.

Control ONHs displayed low label intensity (62.5 ± 5.3 arbitrary units [AU]) for phosphorylated cortactin, which remained low (38.5 ± 4.1 AU) immediately after IOP elevation (CEI day 0), but became significantly elevated (87.0 ± 12.3 AU, P < 0.005) 1 day after IOP elevation (CEI day 1 versus CEI day 0; Figs. 3A, 3B). However, IOP elevation in the setting of ONT (CEI day 0 + ONT, and CEI day 1 + ONT) resulted in significantly lower phosphorylated cortactin label intensity (60.7 ± 8.8 AU and 47.2 ± 6.6 AU, respectively) relative to ONT alone (95.1 ± 6.6 AU, P < 0.05; Figs. 3A, 3B). By 1 day post CEI, phosphorylated cortactin label intensities within the ONH with intact axons were significantly higher compared with values in the absence of viable axons (87.0 ± 12.3 AU vs. 47.2 ± 6.6 AU, P < 0.05; Figs. 3A, 3B).

Control ONHs displayed low phosphorylated paxillin label intensity (23.4 ± 1.8 AU), which significantly increased 1 day after CEI alone (32.9 ± 2.4 AU, P < 0.05), relative to controls (Figs. 4A, 4B). ONT alone significantly increased ONH phosphorylated paxillin label intensity (34.2 ± 2.3 AU) relative to control (24.1 ± 1.6 AU) and CEI day 0 alone (P < 0.05; Figs. 4A, 4B). However, when CEI and ONT were combined, ONH phosphorylated paxillin label intensities (27.6 ± 1.6 AU for CEI day 0 + ONT; 30.1 ± 3.5 AU for CEI day 1 + ONT) were not significantly different than controls (Figs. 4A, 4B). Both immediately after CEI (CEI day 0) and 1 day post CEI (CEI day 1), phosphorylated paxillin label intensities within the ONH were similar in the presence or absence of viable axons (Figs. 4A, 4B).

Astrocytic Complement C3 Label Intensities are Unaffected by Acute IOP Elevation or Optic Nerve Transection

The complement cascade has been implicated in astrocyte reactivity and elevated levels of complement C3 mRNA and
protein have been observed in an optic nerve crush model and a genetic glaucoma model, respectively. We asked if complement C3 label intensities were altered in response to acute IOP elevation, in the presence or absence of viable axons. Control ONHs displayed moderate label intensity of complement C3 (37.4 ± 3.3 AU), which were not significantly altered immediately (33.2 ± 2.3 AU) or 1 day (43.5 ± 2.9 AU) after CEI alone (Figs. 5A, 5B). Similarly, ONH complement C3 label intensity was unaltered after ONT alone (35.8 ± 4.1 AU), relative to controls (Figs. 5A, 5B). The combination of CEI and ONT did not show any significant change in complement C3 label intensity within the ONH relative to controls (36.7 ± 2.2 AU for CEI day 0 + ONT; 31.4 ± 4.2 AU for CEI day 1 + ONT; Figs. 5A, 5B). Thus, complement C3 label intensities within the ONH appear to be unchanged after acute IOP elevation, ONT, or combined IOP elevation and ONT.

**DISCUSSION**

Astrocyte reactivity is observed in multiple neurodegenerative disorders and blocking astrocyte reactivity reduces axon injury. In glaucoma models with elevated IOP, ONH astrocyte reactivity occurs prior to observable axon injury. However, it remains unclear if in vivo ONH astrocyte reactivity is initiated de novo in direct response to the glaucomatous insult (including biomechanical injury to the ONH), or if astrocyte reactivity is initiated in response to ONH axonal, extracellular, or microglial cues. Here, using a well-established model of acute IOP elevation (which leads to ONH astrocyte reactivity), in combination with ONT (which leads to nearly total loss of axon viability), we show that ONH astrocyte reactivity displays both axon-dependent and -independent features.
Our results show that structural ONH astrocyte responses to elevated IOP (namely the re-orientation of astrocyte extensions) occurs despite the total loss of axons, in a similar fashion to ONHs with intact axons. Thus, some ONH astrocyte structural responses to elevated IOP appear to be independent of viable axons, and do not depend on yet unidentified axon signaling. This is supported by three-dimensional astrocyte culture models in which deformations of the culture matrix cause re-orientation of astrocyte, despite the absence of neurons/axons in the culture model. In addition, cultured ONH astrocytes in the absence of neurons display significant actin-based motility within 1 day after initiation of a wound healing assay, and become four to six times more motile 12 hours after exposure to elevated hydrostatic pressure in vitro. ONH astrocyte reactivity to IOP elevation may lead to local remodeling of the extracellular matrix and glial architecture that results in glaucomatous optic disc features seen clinically. However, additional studies in models of ONH remodeling that include a collagenous lamina cribrosa and lamina cribrosa cells (which are not present in the rodent ONH), as well as consideration of other mechanistic factors, such as vascular insufficiency, reduced metabolic support for axons, neuroinflammation, and the contribution of microglia may shed further light on the pathophysiology of ONH remodeling and axon injury in human glaucoma.

**Figure 3.** ONH cortactin phosphorylation label intensities respond differentially after IOP elevation, in the presence or absence of viable axons. (A) ONH sections labeled with antiphosphorylated cortactin (p-cortactin) antibodies in control eyes and eyes exposed to CEI, ONT, or a combination of CEI + ONT. ONH p-cortactin label intensity significantly increased 1 day post CEI (A, B, CEI day 1 versus CEI day 0). However, when CEI and ONT were combined, ONH p-cortactin label intensity significantly decreased after CEI (A, B). (B) Mean p-cortactin label intensities within the ONH of control eyes and eyes exposed to ONT only, CEI only, or combined CEI + ONT. Error bars indicate SEM; *P < 0.05, **P < 0.005, and ***P < 0.0001 and indicates statistical significance; n = 5, 8, 6, 6, 6, and 7 for each column from left to right, and represents individual animals.

**Figure 4.** Paxillin phosphorylation label intensities within the ONH increase after ONT, but not when combined with IOP elevation. (A) ONH sections labeled with antiphosphorylated paxillin (p-paxillin) antibodies in control eyes and eyes exposed to CEI, ONT, or a combination of CEI and ONT. ONH p-paxillin label intensity significantly increased 1 day after CEI alone, relative to controls (A, B). However, when CEI and ONT were combined, ONH p-paxillin label intensity did not significantly change relative to controls or CEI alone (A, B). (B) Mean p-paxillin label intensities within the ONH of control eyes and eyes exposed to ONT only, CEI only, or combined CEI + ONT. Error bars indicate SEM; *P < 0.05 and indicates statistical significance; n = 6, 8, 6, 7, 7, and 4 for each column from left to right, and represents individual animals.
Despite our findings that ONT does not affect the structural response of astrocytes to CEI, we show evidence that certain molecular ONH astrocyte responses to elevated IOP display axon-dependent features. For example, phosphorylation of ONH cortactin (which is a substrate for Src kinase and activates Arp2/3 complex leading to increased actin filament formation) is significantly increased between day 0 and day 1 after IOP elevation. Cortactin phosphorylation enhances actin polymerization through Arp 2/3 activation and is necessary for rapid actin cytoskeletal turnover, particularly in the leading edge of motile cells and protruding cellular extensions. The significantly increased label intensity of phosphorylated cortactin within the ONH 1 day after IOP normalization is likely involved in astrocyte extension re-formation and re-orientation back to baseline levels. This is consistent with astrocyte structural changes observed in models with randomly labeled astrocytes within the ONH, which show that acute IOP elevation leads to extension retraction, followed extension reformation once IOP is normalized. In this study, when IOP elevation leads to extension retraction, followed extension formation within the ONH after IOP normalization, this may be consistent with enhanced astrocyte cellular matrix interactions as seen in a two-dimensional culture model of astrocytes without axons. However, the increase in phosphorylated paxillin label intensity post IOP relatively to controls was not seen when ONT was combined with CEI. This may imply that astrocyte-axon interactions may be necessary for optimal focal adhesion formation within the ONH post-IOP normalization. Thus, there are likely intimate links between astrocyte actin/focal adhesion mediators and axons, which may be facilitated through axonal crosstalk and regulation. This may be expected given the intimate physical relationship of astrocyte extensions, axon bundles, and the complex extracellular matrix of the ONH (Johnson EC, et al. IOVS 1994:ARVO Abstract 1850).

While we note that structural, actin-based responses of ONH astrocytes to elevated IOP (namely extension orientation) are unchanged in the absence of axons, we also noted that some of the actin- and focal adhesion-based phosphorylation events in response to elevated IOP are dependent on the presence of viable axons. In addition to large structural astrocyte orientation changes, there are likely ultrastructural actin-based changes occurring within ONH astrocytes in the setting of IOP elevation (including changes in actin bundle thickness, length, and/or turnover rate), that we simply cannot quantify in a tissue section of densely interdigitated astrocytes. In a mouse model expressing green fluorescent protein in random individual astrocytes, Wang et al. have shown that after IOP elevation ONH astrocytes display new, relatively short cellular extensions along the longitudinal axis of the optic nerve that invade axon bundles of the ONH. Many of these finer ultrastructural changes may involve changes in phosphorylation levels of cortactin and paxillin, which we do see in our model. Thus, astrocyte-axon crosstalk may be an important component of overall astrocytic response to elevated IOP within the ONH.

Complement C3 is central to both classical and alternative pathways of complement activation. Within the ONH, complement C3 gene expression increases in various models of chronic IOP elevation, including the genetic DBA/2J mouse model and a laser model of IOP elevation. However, in a
chronic model of IOP elevation in rats using episcleral hypertonic saline injection, Johnson et al. found that ONH complement C3 gene expression was not significantly elevated with either early or advanced injury. Within the optic nerve, Ohlsson et al. observed significantly elevated message levels with either early or advanced injury. Within the optic nerve, complement C3 gene expression was not significantly elevated in mice that underwent laser-induced IOP elevation. In our study, we found no significant changes in complement C3 label intensity within the ONH after IOP elevation, ONT, or their combination. The lack of changes in complement C3 label intensity within the ONH in our model further highlights the specificity of the other changes noted (namely the structural and actin/focal adhesion–mediator reactivity described above).

Our results support astrocyte reactivity as a possible link between the initial biomechanical insult to the ONH after IOP elevation, and eventual axon degeneration. Early axon-dependent or -independent structural and molecular astrocyte reactivity may reduce the ability of astrocytes to provide mechanical and metabolic support for axons. Indeed, reactive astrocytes lose the ability to promote neuronal survival in culture. Taken together, these results favor a model in which astrocytes are the primary and earliest sensors of ONH injury in glaucoma, and their response to injury precedes and may lead to eventual axon degeneration in glaucoma. Future studies in other model paradigms will help overcome some of the limitations of this study, including the relatively short time course of IOP elevation and follow-up time period, as well as the lack of a true lamina cribrosa in the rodent ONH. Additionally, gene expression studies and proteomic analysis within the ONH may shed further light on the additional cellular and molecular pathways involved in astrocyte reactivity to elevation IOP.

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