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Citation

Published Version
doi:10.1167/iovs.16-20571

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Astrocytes in the Optic Nerve Head of Glaucomatous Mice Display a Characteristic Reactive Phenotype

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Submitted: August 19, 2016
Accepted: January 11, 2017
Citation: Wang R, Seifert P, Jakobs TC. Astrocytes in the optic nerve head of glaucomatous mice display a characteristic reactive phenotype. Invest Ophthalmol Vis Sci. 2017;58:924–932. DOI:10.1167/iovs.16-20571

Purpose. Optic nerve head astrocytes, a subtype of white-matter astrocytes, become reactive early in the course of glaucoma. It was shown recently that in the DBA/2J mouse model of inherited glaucoma optic nerve astrocytes extend new longitudinal processes into the axon bundles before ganglion cell loss becomes apparent. The present study aims at testing whether this behavior of astrocytes is typical of early glaucomatous damage.

Methods. Mice expressing green fluorescent protein in individual astrocytes were used to evaluate the early response of astrocytes in the glial lamina of the optic nerve head after increasing the IOP using the microbead occlusion method. Tissue sections from the glial lamina were imaged consecutively by confocal and electron microscopy.

Results. Confocal and electron microscope images show that astrocytes close to the myelination transition zone in the hypertensive nerve heads extend new processes that follow the longitudinal axis of the optic nerve and invade axon bundles in the nerve head. Ultrastructurally, the longitudinal processes were largely devoid of subcellular organelles except for degenerating mitochondria.

Conclusions. The longitudinal processes are a common feature of glaucomatous optic nerve astrocytes, whereas they are not observed after traumatic nerve injury. Thus, astrocytes appear to fine-tune their responses to the nature and/or timing of the injury to the neurons that they surround.

Keywords: astrocytes, glial lamina, glaucoma, electron microscopy, confocal microscopy

A large body of evidence points to the optic nerve head (ONH) as a site of active tissue remodeling and retinal ganglion cell (RGC) axon damage in glaucoma. In humans and other primates the opening in the sclera that allows the ganglion cells’ axons to exit the eye, is fortified by plates of collagenous tissue forming the lamina cribrosa, and pathologic changes of this structure have been reported in glaucoma.1–6 In this region of the nerve the first signs of blockage of retrograde axonal transport occur.7,8

However, rodents do not have a true collagenous lamina cribrosa.9–12 Yet, several rodent models of glaucoma, spontaneous and induced, exist and share many features with the human disease.13–18 In humans, glaucomatous degeneration of retinal ganglion cells typically follows a sectorial pattern, and this feature is replicated in rodents.19–22 Furthermore, in rats and mice, an elevation of IOP leads to blockage of axonal transport at the level of the ONH where ganglion cell axons run as discrete bundles with a topographic relationship to the retina.12,25–26

As apparently a collagenous lamina cribrosa is not necessary to develop glaucoma, the glial cells of the optic nerve have attracted attention as possible players in the pathophysiology. In the nerve head, the RGC axons are unmyelinated, but they are ensheathed by a dense meshwork of GFAP-positive astrocytes, the glial lamina.10,12,27 Individual cells making up the glial lamina are relatively big and usually span at least half the diameter of the nerve, they overlap extensively, and their processes are oriented perpendicularly to the long axis of the nerve.12,28 In response to traumatic injury or glaucoma, the astrocytes become reactive and lose their typical arrangement, reorient or retract many of their processes, and become irregular in shape.29–36 Using the DBA/2J model of inherited glaucoma, we recently demonstrated that astrocytes in the glial lamina grow out new longitudinal processes that invade the axon bundles rather than participating in forming the glial tubes around them.38 These abnormal longitudinal processes appeared before there was detectable damage to the RGCs (at 6–7 months of age), but it was not clear what their function might be. We report an extension of our earlier study to the level of electron microscopy. We also have used the microbead occlusion model of ocular hypertension38,40,41 rather than the DBA/2J strain. In the microbead model, the onset of the injury is defined more clearly in time and allows the study of very early changes in astrocyte morphology.

Materials and Methods

Animal Strains and Husbandry

All animal experiments were done according to the guidelines of the Association for Research in Vision and Ophthalmology (ARVO) Statement for the Use of Animals in Ophthalmic and Vision Research and were approved by the Institutional Animal
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Care and Use Committee at the Schepens Eye Research Institute. We used hGFAPpr-GFP mice\(^1\) that were backcrossed onto the C57bl/6 background. These mice express green fluorescent protein (GFP) in individual astrocytes in the optic nerve and other parts of the central nervous system (CNS).\(^1\) The strain was maintained in heterozygous state and was backcrossed in each generation to C57bl/6 (obtained from the Jackson Laboratories, Bar Harbor, ME, USA) to avoid genetic drift. Individual animals were tested for GFP expression by inspecting ear snips under a fluorescent microscope.\(^3\) Male and female mice of 2 to 8 months were used for all experiments.

**Microbead Injection**

Ocular hypertension was induced by the injection of microbeads (15 μm diameter; Invitrogen, Carlsbad, CA, USA) into the anterior chamber of the right eye, essentially as described by the developers of the method.\(^4\) Another group of mice was injected unilaterally with 2 to 3 μl sterile saline solution as an additional control. Intraocular pressure (IOP) was measured with a Goldmann applanation tonometer (TonoLab; iCare, Espoo, Finland). All measurements were taken at the same time in the morning to minimize circadian variation in IOP. The cumulative IOP (cIOP) was calculated for each eye as the area under the curve of IOP over time with a user-written Matlab (Mathworks, Natick, MA, USA) routine.\(^4\)

**Tissue Preparation and Embedding**

The head with the eyes and optic nerves in situ were fixed for 2 hours in 4% paraformaldehyde. Retinas and optic nerves were dissected from the surrounding tissue as described previously.\(^1\) Optic nerves were detached from the retina, embedded in 6% agarose, and sectioned at 100 μm, or 200 μm (when electron microscopy was performed after confocal imaging) in transverse orientation using a vibratome (Leica VT1000 S; Leica Microsystems, Buffalo Grove, IL, USA) and mounted with Vectashield (Vector Laboratories, Burlingame, CA, USA). Retinas were counterstained with the nuclear dye DAPI (Life Technologies, Grand Island, NY, USA), and mounted in Vectashield.

**Confocal Microscopy and Image Reconstruction**

Images of astrocytes in the unmyelinated segment of the optic nerve were obtained on a Leica TCS SP8 confocal microscope using a ×63 glycerol immersion objective. Individual agarose sections were kept in order starting from 1 (at the level of the sclera to 100 μm behind the sclera) to 4 (300–400 μm behind the sclera). We restricted the analysis to the glial lamina, and astrocytes in the optic nerve proper (myelinated portion) were not imaged. Image stacks were taken through the whole extent of the cell. Image size was 1024 × 1024 pixels in x/y, corresponding to a resolution of 0.1805 μm per pixel. The z step size was 0.33 μm. The 3D reconstruction of astrocytes was accomplished using a volume visualization program (Amira; Visage Imaging, San Diego, CA, USA).

**Transmission Electron Microscopy**

After confocal microscopy the optic nerve sections with the surrounding agarose were immediately immersion fixed with half strength Karnovsky’s fixative for 2 hours at room temperature. The agarose section was sandwiched between two filters to help stabilizing and handling the delicate tissue. The samples were processed for electron microscopy and imaged using a FEI Tecnai G2 Spirit transmission electron microscope (FEI, Hillsboro, OR, USA) at 80 kV equipped with an AMT XR41 digital CCD camera (Advanced Microscopy Techniques, Woburn, MA, USA). A complete description of the tissue preparation procedure is given in the Supplementary Methods.

**Immunohistochemistry**

Whole-mounted retinas were incubated in primary antibodies for 3 to 4 days at 4°C. The primary antibodies used were: rabbit anti-β-III tubulin (1:200; Cell Signaling Technology, Danvers, MA, USA), mouse anti-Brn3a (1:200, Nippon Chemi-Con, Tokyo, Japan). Secondary antibodies were FITC-conjugated donkey anti-rabbit IgG, and rhodamine-conjugated donkey anti-mouse IgG (both from Jackson Immunoresearch Laboratories, Inc., West Grove, PA, USA). Retinas were counterstained with the nuclear dye DAPI (Life Technologies, Grand Island, NY, USA), and mounted in Vectashield.

**Ganglion Cell Counting**

Retinal ganglion cells were counted on whole-mounted retinas after immunostaining for Brn3a and β-III tubulin. The images were acquired with a ×65 glycerol immersion objective on a Leica TCS SP8 confocal microscope. The retina was divided into quadrants and two mid-peripheral regions (defined as equidistant to the ONH and the border of the retina) were imaged in every quadrant (8 regions of 184.52 μm × 184.52 μm per retina). To include all cells in the ganglion cell layers, z-stacks were taken through the ganglion cell layer at a step size of 0.5 μm, and maximum-intensity projections of these stacks were made in ImageJ. All cells in the ganglion cell layer that colocalized with anti-β-III tubulin and anti-Brn 3a were counted and the ganglion cell density per retina was calculated. Cells positive for β-III tubulin were counted manually by individuals blinded to different groups of the animal using ImageJ (National Institute of Health [NIH], Bethesda, MD, USA), Brn3a-positive cells were counted semiautomatically in a user-written Matlab routine (version R2013b). Differences in RGC counts were tested for significance using Student’s t-test.

**Cell Tracing**

Every animal received a number code after the experiment and the investigator performing morphometric measurements and statistical analysis was ignorant of the IOP history of the eye. Unprocessed confocal image stacks through the glial lamina were imported to ImageJ. To quantify longitudinal processes individual astrocytes were traced using the Simple Neurite Tracer plugin.\(^4\) Image stacks containing more than 10 labeled cells were not analyzed because in these cases it was impossible to discern individual cells. For examples of digitized astrocytes, see Supplementary Figure S1. The length of all processes (transverse and longitudinal) was measured for each cell and the ratio of longitudinal process length to total process length (L/T) was calculated. Differences between groups were tested for significance using Kruskal-Wallis and Wilcoxon rank sum tests. All statistical tests were done in Matlab.

**Data Sharing**

All original unprocessed images and image stacks were deposited in the Harvard Dataverse and are accessible in the
RESULTS

Elevation of IOP Induced by Microbead Injection Into Anterior Chamber

We injected 27 GFP-positive hGFAPpr-GFP mice unilaterally with microbeads. The mice were 6 to 8 months of age at the beginning of the experiment because we first observed longitudinal processes in the astrocytes of DBA/2J mice at that age.38 Four mice were excluded from the study because they showed corneal opacities that did not resolve within days after the injection. The contralateral eye was left untreated. Recently, it was demonstrated that elevation of the IOP in one eye leads to glial cell activation in the contralateral eye, and in some mouse strains, even to RGC loss.47–50 Therefore, the contralateral eye cannot be considered truly “normal.” Therefore, we included 8 naïve mice age matched to the young animals (5 months) to determine whether age alone has an influence on astrocyte morphology. We also injected another group of 8 young adult mice unilaterally with sterile saline solution to exclude the possibility that any manipulation of the eye may cause glial activation or ganglion cell loss. The IOP was measured every 3 or 4 days in the following 4 weeks after surgery (Supplementary Figs. S2A, S2B).

Due to the variability of the level and duration of IOP changing between individual animals, we used the cumulative IOP, defined as the area under the curve of IOP over time, measured in mm Hg days.8,4 However, as we did not use the contralateral eye as the “normal” control, we calculated the cIOP for each eye rather than the difference in cIOP between the injected and contralateral eyes. The cIOP was 479.6 ± 44.58 mm Hg days (mean ± SD, n = 23) for the microbead-injected eyes, which was higher than for the contralateral eyes (410.8 ± 46.82, P < 0.001, t-test, Fig. 1A). For the sham operated group (saline injection), there was no significant difference in the cIOP comparison between saline injected eyes (300.4 ± 12.26, n = 8) and contralateral eyes (312.6 ± 23.5, n = 8, P = 0.21, Fig. 1B). The maximum IOP for the microbead group was 23.7 ± 3.16 mm Hg (mean ± SD, n = 23), which was higher than in the contralateral group (19.5 ± 3.07, n = 23, P < 0.001, Fig. 1C). We also noticed that the IOP in the contralateral eyes of the old mice was higher than baseline (15.16 ± 0.96 mm Hg in the contralateral eyes versus the baseline of 12.08 ± 1.6 mm Hg, P < 0.01). There was no significant difference in maximum IOP between saline injected eyes (15.6 ± 0.9) and contralateral eyes (14.5 ± 1.8, n = 8, P = 0.12, Fig. 1D).

Ganglion Cell Loss After IOP Elevation

Brn3a and β-III tubulin were used to assess RGC loss (Figs. 2A–I). We compared the RGC densities obtained from these two antibodies in 10 retinas. The RGC number was 8.58% ± 2.8% lower estimated by Brn3a staining rather than by β-III tubulin staining, but the differences between these two counting methods were consistent (Fig. 2J).

We first determined the RGC density (by staining for Brn3a) in a group of age-matched naïve mice that had no procedures performed on them as a baseline value. The RGC density in this group was 4135 ± 333 cells/mm² (n = 8). The RGCs densities in the eyes that received microbead injections (n = 23) were significantly lower than those of the contralateral eyes (3835 ± 363 vs. 4170 ± 242 cells/mm², P < 0.001, Fig. 2K). There was no significant difference between the naïve eyes and the contralateral eyes, suggesting that unilateral microbead injection did not cause RGC loss in the contralateral eye in this strain. RGC loss was only slightly correlated with cIOP (R² < 0.1).

We also injected a group of young (2 months) mice unilaterally with sterile saline solution. There was no RGC loss in the injected eye in this group (4650 ± 181 vs. 4726 ± 298 cells/mm², not significant, Fig. 2L). The overall higher RGC density in the younger mice is consistent with the observation that mice spontaneously lose cells with age, even without the presence of glaucoma.59

Longitudinal Astrocytic Processes in the Glial Lamina

Though the shape of the astrocytes in the glial lamina is fairly stereotypical, the cells are quite complex. To quantify cell morphologies, we digitally traced all astrocytes in the sample that were well-isolated enough from neighboring labeled cells to unambiguously identify their processes. Two sets of strain-matched naïve mice served as controls. First, we used 3-month-old animals and traced 30 individual astrocytes from 11 nerves to establish the baseline morphology of normal ONH astrocytes. However, morphologic changes might occur with normal aging, so we additionally traced 24 astrocytes from 6
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Figure 2. Ganglion cell loss after microbead injection. (A, D, G) Mouse anti-Brn3a antibody labeled ganglion cells in microbead-injected eyes, their contralateral eyes, and aged-matched naive eyes. (B, E, H) Rabbit anti β-III tubulin antibody labeled ganglion cells in microbead-injected eyes, their contralateral eyes and aged-matched naive eyes. (C, F, I) Merged images show double labeled ganglion cells in all three groups. Scale bar: 20 μm. (J) Comparison of ganglion cell densities obtained from mouse anti-Brn3a antibody labeling and rabbit anti β-III tubulin antibody labeling in 10 whole mount retinas. The ganglion cell densities obtained from Brn3a staining were relatively lower estimated than β-III tubulin staining, but the differences between these two counting methods were consistent. All images represent maximum-intensity projections of image stacks through the ganglion cell layer, taken at a z-step size of 0.5 μm. (K) Ganglion cell densities in microbead-injected eyes were significant lower than those in the contralateral eyes. N = 25 (B), n = 8 (C). **P < 0.001 (t-test). Data are presented as mean values ± SD.

nerves at 8 to 9 months to serve as strain- and age-matched controls for the microbead injected eyes. In all cases, the unmyelinated region was subdivided into 100 μm sections beginning at the sclera (region 1) to the myelination transition zone (region 4). Sections at more than 400 μm behind the sclera contained myelinated optic nerve and were not imaged. Figure 3 shows typical ONH astrocytes from the naïve B6.hGFAPpr-GFP mice at 3 (Figs. 3A–D) and 8 to 9 (Figs. 3E–G) months by regions 1 to 4. The processes of astrocytes from naïve optic nerves are long and smooth, spanning over half the diameter of the optic nerve. Three-dimensional reconstruction shows that the whole astrocyte is slightly concave, with most of the processes oriented transversely to the long axis of optic nerve—the typical morphology of astrocytes in the glial lamina.12 There are numerous processes emanating mostly from the cell body and the proximal segments of the main processes (arrows in Fig. 3), but as shown in 3D reconstructions, very few longitudinal processes are present in naïve astrocytes (Fig. 3). There was no difference between the astrocytes from the young and old animals. The total process length was 1140.34 ± 459.44 μm for the 3-month group and 994.73 ± 345.52 μm for the 8- to 9-month group (not significant). The L/T ratio (length of all longitudinal processes divided by total process length) was 0.0316 (range, 0–0.0874) for young and 0.0248 (range, 0–0.0944) for old animals (not significant). This indicates that the growth of longitudinal processes is not driven by age alone.

The astrocytes from the microbead-injected group showed obvious morphologic changes (Figs. 4C–E) and delicate longitudinal processes were found to grow out from the new formed processes and ganglion cell axon bundles. We imaged individual astrocytes in 200 μm agarose sections of the ONH 1 month after microbead injection with confocal microscopy to verify that longitudinal processes were present. Afterwards, we used the same tissue to generate thin sections for electron microscopy to inspect the relationship between newly formed processes and ganglion cell axon bundles. Longitudinal processes should be identifiable because their intermediate filaments will be sectioned transversely (and, was localized. As shown in Figure 5A, astrocytes in the initial 200 μm behind the sclera, rarely had longitudinal processes. Closer to the myelination transition zone (200–400 μm) these processes were far more common. We next tested whether the presence of longitudinal processes correlates with RGC loss. For this, each of the 14 nerves was treated as one biologic replicate since all cells in the same nerve were presumably subject to the same pressure insult and ganglion cell loss. As longitudinal processes rarely occurred in the initial 200 μm, only cells from 200–400 μm were considered. In those cases we had more than one traced astrocyte per nerve their L/T ratios were averaged. The L/T ratios correlated with RGC loss (R² = 0.41, Fig. 5B). There was no obvious difference between nerves from male and female mice (Fig. 5B).

Finally, we asked whether longitudinal processes only occurred in old eyes with ocular hypertension. Therefore, as an additional control, we injected 3 young B6.hGFAPpr-GFP mice with microbeads and digitized 8 astrocytes from their glial laminae 200 to 400 μm behind the sclera. Total process length in the young injected cells was similar to the naïve controls (1029 ± 179.75 μm, not significant), but the L/T ratio was higher (0.0868; range, 0.0097–0.2518, P < 0.05). Taken together, these findings indicate that the growth of longitudinal processes is likely driven by the increase in IOP or ganglion cell distress and are a common feature of reactive astrocytes in the optic nerve regardless of age.

Transmission Electron Microscopy

We imaged individual astrocytes in 200 μm agarose sections of the ONH 1 month after microbead injection with confocal microscopy to verify that longitudinal processes were present. Afterwards, we used the same tissue to generate thin sections for electron microscopy to inspect the relationship between newly formed processes and ganglion cell axon bundles. Longitudinal processes should be identifiable because their intermediate filaments will be sectioned transversely (and,
thus, appear as puncta) rather than longitudinally, as in the regular astrocyte processes that form the glial tubes. Furthermore, profiles of longitudinal processes should occur in the middle of axon bundles. Following these rules, we detected numerous profiles of longitudinal processes in the ONH from microbead-injected eyes (Figs. 6A, 6B). The longitudinal processes did not contain subcellular organelles, but often electron-dense inclusion bodies were present that were surrounded by a membrane. Higher magnification images showed that these were degenerating mitochondria (Figs. 6A’, 6B’). In ONHs after microbead injection, we also observed vacuoles of different shapes and sizes (Fig. 7A). Some of these vacuoles seemed to be adjacent to a longitudinal process (Fig. 7B). Though most of the ganglion cell axons appeared morphologically normal, the space between single axons was loosened, and the astrocytes grew out numerous small processes between the axons (Figs. 7C-E). In some axons we observed abnormally large mitochondria or mitochondria with very few cristae (Fig. 7F). Neither longitudinal processes nor vacuoles were found in naïve optic nerves from young mice (Figs. 6C, 6D). The axons within each bundle were tightly packed with very few astrocyte processes that invaded the bundles (Fig. 6C). Axonal mitochondria always had visible cristae (Fig. 6D).

Comparison of Glaucomatous Optic Nerves With Aged Optic Nerves

In electron microscopy, our primary control for a normal ONH were young (3 months old), nonglaucomatous C57bl/6 mice. However, mice lose RGCs spontaneously with age, and some of the pathologic changes we observed in our experimental animals may be related to age rather than to an elevation in IOP. Therefore, we used the optic nerves from a naïve 6-month-old C57bl/6 mouse, prepared in exactly the same way as described above, as an additional control. In these older nerves, we did not observe either longitudinal processes or vacuoles (Fig. 6E).

In Figure 3, typical ONH astrocytes from naïve B6.hGFAPpr-GFP mice. (A-D) Astrocytes obtained from ONHs of 3 months naive mice. (A-D) Three-dimensional reconstructions of the astrocytes in (A-D). (E-H) Astrocytes obtained from ONH of 9 month naive mice. (E-H) Three-dimensional reconstructions of astrocytes in (E-H). The processes are long and smooth (arrowbeads in [B–E, G]). There are numerous fine processes emanating mostly from the cell body and the proximal segments of the main processes (arrows in [C, D, E, G]). Only occasionally there are processes in the longitudinal direction in naïve ONH astrocytes (arrow in [G]). R1, region 1, 0 to 100 μm behind the sclera; R2, region 2, 100 to 200 μm behind the sclera; R3, 200 to 300 μm behind the sclera; R4, 300 to 400 μm behind the sclera. Scale bar: 50 μm.

In Figure 4, green fluorescent protein–positive astrocytes in the ONH from naïve, contralateral, and microbead-injected eyes. (A, B) Typical ONH astrocytes from the naïve (A) and the contralateral (B) eyes. The processes of astrocytes from naïve optic nerves are long and smooth (arrowbeads), spanning over half the diameter of the optic nerve. There are numerous very short processes emanating mostly from the cell body and the proximal segments of the main processes (arrows). (C-E) Optic nerve head astrocytes from the microbead-injected eyes. Some processes became thicker (arrowbeads). (A) Three-dimensional reconstruction of astrocyte in (A). The whole astrocyte is fairly flat, slightly concave, with most of the processes oriented transversely to the long axis of the optic nerve. Very few longitudinal processes are present in naïve astrocyte. (D) Three-dimensional reconstruction of astrocyte in (D). A large number of longitudinal processes were found to grow out from the processes of ONH astrocytes. The schematics in (A) and (A) indicate the plane of section. Scale bar: 50 μm.
In an earlier study, we reported that astrocytes from the ONH extend new longitudinal processes into the axon bundles in 6- to 8-month-old DBA/2J mice. The changes in the optic nerve preceded overt RGC loss, and were the first obvious signs of reactivity we observed in astrocytes. However, there are two caveats about our earlier study. First, the observations were made in DBA/2J mice. DBA/2J is a valuable model of glaucoma, but it differs from other rodent models of glaucoma in terms of gene expression, and its reaction to radiation treatment. A possible reason may be the pronounced inflammatory component of DBA/2J glaucoma. Given the differences between rodent glaucoma models, generalizations from one of them to glaucoma in general must be treated with caution. Second, in our earlier study, we did not use electron microscopy and, therefore, were unable to obtain more details about the relationship between these newly formed longitudinal processes and injured optic nerve axons.

Using unilateral microbead injection in B6.hGFAPpr-GFP mice, we asked whether longitudinal processes are a common feature of glaucomatous optic nerve rather than a peculiarity of the DBA/2J strain. In our cohort of 7- to 8-month-old mice, microbead injection led to a significant increase in cIOP. However, we noted that the IOP in the contralateral eye in these older mice is higher than in younger animals, agreeing with previous reports. Older mice on the C57Bl/6 background have been shown to be less vulnerable to ganglion cell loss in the microbead model than their younger counterparts or mice of other strains. We did observe a significant, albeit moderate, loss of RGCs in the microbead injected eyes; however, the loss was poorly correlated to the pressure insult the eye had sustained. A possible explanation for this observation might be that taking IOP measurements twice per week does not truly reflect the IOP history of the eye during the observation period and misses events of elevated IOP or pressure spikes that may be detectable only by continuous telemetry. In addition, even in inbred strains individual differences in the vulnerability of RGCs to IOP may exist.

### DISCUSSION

**FIGURE 5.** Longitudinal processes in ONH astrocytes. (A) Longitudinal processes were fairly rare in the initial 200 µm behind the sclera. The L/T ratios are given for every 100 µm segment. Filled dark blue, 0 to 100 µm, n = 3 cells; red, 100 to 200 µm, n = 8; green, 200 to 300 µm, n = 22; violet, 300 to 400 µm, n = 12. Black bars represent group means. Open symbols represent L/T ratios of age- and strain-matched naïve control cells (n = 3, n = 4, n = 11, n = 6 for the four regions, respectively). Gray bars represent group means for the naïve cells. *P < 0.05, Wilcoxon rank sum test. (B) Correlation of L/T ratios versus retinal ganglion cell loss (in %). Blue symbols represent nerves from male mice; red symbols represent nerves from female mice. Data are presented as mean values ± SD.

**FIGURE 6.** Electron microscopy of the ONH from microbead-injected eyes and naïve eyes. (A, B) Optic nerve head after microbead injection. Regular astrocyte processes (asterisks) with intermediate filaments. The longitudinal processes can be identified by the orientation of their intermediate filaments and the lack of continuity with the other processes. There are two longitudinal processes in the middle of the axon bundle in (A) and two longitudinal processes (a big one in the lower left and a small one in upper right) in the middle of the axon bundle in (B). Arrows point to materials in the longitudinal processes. (A’, B’) The insert shows a high-power image of the region corresponding to the black box in (A) and (B). The materials inside the longitudinal processes are degenerating mitochondria, with some cristae still visible. (C-F) Optic nerve head of a 3-month-old naïve mouse (C, D) and of a 6-month-old naïve mouse (E, F). Asterisks indicate regular astrocyte processes that form the glia tube. No longitudinal processes were found. The axons within each bundle were tightly packed with very few astrocyte processes that invaded the bundles (C, E). Axonal mitochondria (arrows in [D]) in 3-months-old ONH always had visible cristae. Some axons in 6-months-old ONH contained abnormal mitochondria (arrows in [F]). Scale bars: 2 µm (A, C, E), 500 nm (B, D, F), 200 nm (A’, B’).
We observed longitudinal processes similar to those that are present in DBA2/J nerves after microbead injection, indicating that they are a common feature of glaucomatous optic nerve astrocytes. This behavior is not found in protoplasmic astrocytes after a stab wound to the gray matter, where the astrocytes occupy nonoverlapping spatial domains which are maintained even after injury. Nor is it observed after severe traumatic injury in the astrocytes of the glial lamina and the optic nerve proper. Glaucoma leads to slow, but progressive, RGC damage, and the astrocytes may react to this type of injury in a unique way.

The presence of the longitudinal processes depended on the localization of the cell within the unmyelinated segment of the nerve: closer to the myelination transition zone more longitudinal processes were encountered. This may be an indication of phagocytic activity in these cells, as astrocytes in the myelination transition zone are active phagocytes. Ultrastructurally, the longitudinal processes contain sparse intermediate filaments and no subcellular organelles other than apparently degenerating mitochondria. Davis et al. recently suggested that optic nerve astrocytes aid in the transcellular degradation of axonal mitochondria, and the mitochondria in the longitudinal processes may be of axonal origin.

Compared to the normal ONH, the space between single axons appears loose after IOP elevation, and fine astrocytic processes intermingle with the unmyelinated axons, sometimes encircling single axons. Recent studies identified decreased axon packing and an increase in astrocyte processes for the myelinated portion of the optic nerve, also. One of these studies also reported a loss of axonal mitochondria in glaucomatous nerves before overt degeneration and abnormal mitochondria accumulate in the axons of glaucomatous DBA/2J mice. Thus, an imbalance of energy production and metabolic demand of the RGCs, particularly in the aging nerve, may be an important factor in ganglion cell degeneration.

We hypothesized that newly formed astrocyte processes are attracted to distressed axons by signals that are as yet unknown. As in the spinal cord, one of their functions may be to surround and limit the spread of axonal damage. The processes also may be phagocytic and possibly relieve the axons of degenerating mitochondria. Finally, astrocytes may be involved in clearing the debris of axons that did eventually degenerate and die.

Acknowledgments

The authors thank Elio Raviola and Frederic Jakobiec for advice on electron microscopy; and Richard Masland, Daniel Sun, and Anthony Pappas for critically reading the manuscript. Supported by NIH Grants 2R01EY019703 and R01EY022092, the NIH Core Grant for vision research P30EY003790, and grants from the Chinese Scholarship Council and Research to Prevent Blindness.

Disclosure: R. Wang, None; P. Seifert, None; T.C. Jakobs, Santen, Inc. (R)

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