Susceptibility to Neurodegeneration in a Glaucoma Is Modified by Bax Gene Dosage

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In glaucoma, harmful intraocular pressure often contributes to retinal ganglion cell death. It is not clear, however, if intraocular pressure directly insults the retinal ganglion cell axon, the soma, or both. The pathways that mediate pressure-induced retinal ganglion cell death are poorly defined, and no molecules are known to be required. DBA/2J mice deficient in the proapoptotic molecule BCL2-associated X protein (BAX) were used to investigate the roles of BAX-mediated cell death pathways in glaucoma. Both Bax+/+ and Bax−/− mice were protected from retinal ganglion cell death. In contrast, axonal degeneration was not prevented in either Bax+/+ or Bax−/− mice. While BAX deficiency did not prevent axonal degeneration, it did slow axonal loss. Additionally, we compared the effects of BAX deficiency on the glaucoma to its effects on retinal ganglion cell death due to two insults that are proposed to participate in glaucoma. As in the glaucoma, BAX deficiency protected retinal ganglion cells after axon injury by optic nerve crush. However, it did not protect retinal ganglion cells from N-methyl-D-aspartate (NMDA)-induced excitotoxicity. BAX is required for retinal ganglion cell death in an inherited glaucoma; however, it is not required for retinal ganglion cell axon degeneration. This indicates that distinct somal and axonal degeneration pathways are active in this glaucoma. Finally, our data support a role for optic nerve injury but not for NMDA receptor-mediated excitotoxicity in this glaucoma. These findings indicate a need to understand axon-specific degeneration pathways in glaucoma, and they suggest that distinct somal and axonal degeneration pathways may need to be targeted to save vision.

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Introduction

Glaucoma is a common blinding disease affecting approximately 70 million people worldwide [1]. Glaucoma is often associated with elevated intraocular pressure (IOP). IOP elevation and glaucoma are typically spontaneous, progressive, idiopathic processes and are most common in the elderly [2]. Although IOP-lowering treatments slow the development and progression of glaucoma in many patients [3,4], it is not always possible to reduce IOP to a “safe” level [5]. Vision loss in glaucoma is the result of retinal ganglion cell (RGC) death with accompanying optic nerve atrophy, so glaucoma is a neuropathy. IOP elevation is not detected in a significant subset of glaucomas [6,7]. Thus, the unifying characteristic of glaucoma is RGC death. While there are several hypotheses as to why elevated IOP kills RGCs, both the precise biochemical cascades that are triggered within RGCs and the nature of the proximal insult(s) that trigger these cascades remain superficially defined [8]. No treatments that directly protect the neurons are in routine clinical use.

The complex nature of glaucoma makes studies of its pathogenesis difficult [9]. Consequently, no specific molecules have been shown to be essential for RGC death in glaucoma. Standard glaucoma-relevant models include direct RGC trauma, direct optic nerve trauma, and suddenly induced IOP elevation [10–18]. Although these induced models have provided valuable information, the relevance of specific damaging mechanisms may differ significantly between spontaneous and experimentally induced glaucomas. Thus, studies using inherited glaucoma models are also necessary.

Apoptosis is known to contribute to RGC death following experimentally induced insults including axotomy and IOP elevation (e.g., [19,20]), and there is also some evidence that apoptosis is involved in human glaucoma [21,22]. A number of molecules that are known to affect apoptosis are reported to be important regulators of RGC death after various induced insults. These include X-linked inhibitor of apoptosis protein (XIAP) [23–25], p38 [26], several caspases [27–30], the B-cell lymphoma/leukemia 2 (BCL2) family of apoptotic regulators [20,31–34], and members of the c-Jun N-terminal kinase (JNK) [35,36] and tumor necrosis factor (TNF) [35,37] signaling pathways. One of these molecules, BCL2-associated X protein (BAX; a proapoptotic member of the BCL2 family), has a major role in mitochondrial-mediated apoptosis in different neuronal cell types [38,39]. In mice, BAX deficiency increases the number of RGCs in the adult retina by 220% by allowing more RGCs to survive during development [39]. Genetic or induced BAX deficiency is also known to prevent RGC apoptosis after optic nerve crush and axotomy [13,18,40]. Thus, BAX-mediated apoptosis is clearly an important mechanism of stress-induced RGC death. Whether
Synopsis

Glaucoma is a group of diseases whose unifying characteristic is death of nerve cells (retinal ganglion cells) that connect the eye to the brain. Glaucoma is often associated with a harmfully high pressure inside the eye (intraocular pressure) contributing to nerve cell death. Various treatments are used to lower eye pressure, but currently no commonly used treatments directly protect the nerve cells. DBA/2J mice develop elevated eye pressure with age, and this pressure kills retinal nerve cells. The authors use this mouse model to investigate how these nerve cells die in glaucoma. They show that there are distinct degeneration pathways activated in different parts of the retinal nerve cells. They found that the biochemical pathway in the nerve cell body, which resides in the retina, requires a molecule called BAX (BCL2-associated X protein). In contrast, pathways in the part of the cell (axon) that connects the cell body to the brain do not require BAX. Because degeneration pathways in the cell body and of the axon also may be molecularly different in human glaucoma, it will be important to consider them all when designing therapies. Their data also suggest that the BAX gene is a candidate to modulate glaucoma susceptibility.

or not this pathway has a role in IOP-induced RGC death in either experimentally induced or inherited glaucomas is not known.

Understanding the pathophysiologic mechanisms of RGC death in glaucoma and the genetic susceptibility factors contributing to this process is important for the development of effective and individualized treatments. Here, we use the genetically uniform DBA/2J mouse model of glaucoma [41–43] to assess the importance of mitochondrially mediated apoptosis in an inherited glaucoma. Importantly, we show that in this model of inherited glaucoma there are distinct RGC death and axonal degeneration pathways. The RGC death pathway is BAX dependent and, therefore, apoptotic. The axonal degeneration pathway is BAX independent. Finally, our data suggest that reducing BAX levels in the retina may retard the rate of vision loss in glaucoma.

Results

Apoptosis is Physiologically Relevant for RGC Death in an Inherited Glaucoma

To determine if RGC apoptosis has a significant role in an inherited glaucoma, we assessed the effects of BAX deficiency on RGC survival and on RGC axonal degeneration (see Materials and Methods). Our results show that BAX is not required for RGC axon degeneration. Bax+/− mice developed severe optic nerve damage, including essentially complete loss of axons (Figure 2). In contrast, our experiments show that BAX is required for RGC death in glaucoma (Figure 3). Despite severe axonal degeneration, the numbers of cell bodies in the RGC layer of Bax+/− mice were normal. As a stringent test of this observation, we counted RGC-layer cell bodies in the retinas of mice with severe (≥ 95% loss) axon degeneration. The number of RGC cell bodies was normal in Bax+/− mice with more than 95% axon loss (Figure 3). Importantly, Bax+/− mice were also protected against glaucomatous RGC death. Bax+/− mice with an axon loss of 95% or more also had substantially increased survival of RGC cell bodies as compared to Bax+/− controls (Figure 3E). Thus, RGC death and axonal degeneration are clearly distinguished in these experiments.

Other Proapoptotic Molecules Do Not Compensate for BAX Deficiency

In some neuronal cell types, BAX deficiency delays but does not prevent apoptosis [49]. This is because other proapoptotic molecules (e.g., another BCL2 family member, BAK) mediate cell death in the BAX-deficient neurons [50]. To test this possibility in the DBA/2J model, we aged Bax+/− mice to 18 mo. As expected in a complex age-related disease, the severity of glaucomatous damage varies between individual DBA/2J eyes at any age. Nevertheless, by 12 mo of age, the majority of eyes have severe optic nerve damage (see below). Therefore, 18 mo of age is 6 mo after the majority of eyes have severe axon loss. Despite this extensive axonal degeneration, there was no obvious reduction in RGC numbers in any of the 18-mo-old Bax+/− eyes (Figure 3E). This result indicates that other molecules do not substitute for BAX and that BAX is essential for RGC apoptosis in DBA/2J inherited glaucoma.

Homologous but Not Heterozygous BAX Deficiency Alters RGC Number

To test the role of BAX in glaucomatous RGC death, we extensively backcrossed a previously characterized null allele of Bax (Bax(homo)) [44] onto the inbred DBA/2J background. In mammals, approximately twice as many RGCs are produced during retinal development than survive into adulthood [45–47]. As expected from previous studies of retinal development on a different genetic background [39], complete BAX deficiency increased the number of RGC-layer somata in adult DBA/2J mice by 220% (average cell number per 40× field ± standard error of the mean [SEM], number of retinas analyzed: Bax+/−, 199 ± 5.6, n = 7; Bax−/−, 437 ± 15.3, n = 8). In agreement with this, Bax−/− mice had 217% more RGC axons than Bax+/− mice (Bax+/−, 50,504 ± 1,988, n = 8; Bax−/−, 108,907 ± 10,322, n = 4; p < 0.001). Reflecting the increased number of RGC axons and the proportional increase in glial cell types [48], the cross sectional area of Bax+/− optic nerves was significantly increased (average ± SEM, number of optic nerves measured: Bax+/−, 0.157 ± 0.005 mm², n = 13; Bax−/−, 0.278 ± 0.008 mm², n = 16, p < 0.001). In heterozygous Bax+/− mice, RGC number (average per 40× field ± SEM, 212 ± 14.0, n = 5) and the optic nerve area (0.171 ± 0.007 mm², n = 13) was not different from Bax+/− mice (p = 0.352 and p = 0.107, respectively). Thus, heterozygous levels of BAX are sufficient for death of the normal numbers of RGCs during retinal development.

BAX Ablation Preserves RGC Numbers but Does Not Prevent RGC Axonal Degeneration in Glaucoma

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Homologous but Not Heterozygous BAX Deficiency Alters IOP

DBA/2J mice develop a form of pigmented glaucoma that is secondary to a progressive iris disease. Iris pigment and cell
Figure 1. Dying RGCs Have Characteristic Features of Apoptosis

(A–C) A double-labeling assay that identifies fragmented DNA using fluorescently labeled dUTP (A) and detects chromatin condensation by binding of the dye YOYO-1 (B) was used to assess the presence of these hallmarks of apoptosis in glaucomatous DBA/2J eyes at 10–11 mo of age (a time when many RGCs die). A cell in the retinal ganglion cell layer (GCL, arrowhead) has both of these features of apoptosis as indicated by double labeling (C). INL, inner nuclear layer.

(D–F) Electron microscopy provided further evidence for apoptosis. (D) An example of a healthy RGC. (E) Chromatin condensation (a hallmark of apoptosis) along the inner surface of the nuclear envelope in a ganglion cell (arrows). The internal limiting membrane of the retina is indicated by arrowheads. (F) An apoptotic body in the ganglion cell layer (arrows) containing a nuclear fragment with prominent condensed chromatin (asterisk) and other cell remnants.

(G) A TUNEL assay (see Materials and Methods) was used to assess the prevalence of cell death at different ages. TUNEL labeling was not detected at 7 mo (an age prior to glaucomatous cell death) and peaked at 10–13 mo, when most RGCs die. No TUNEL-positive cells were detected in nonglaucomatous, age-matched control mice. These results support an important role of apoptosis in RGC death in spontaneous glaucoma. Scale bar, 1 μm.

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found no differences (average 6
we analyzed IOP in 4-mo-old mice of each genotype and
(Figure 4). In contrast, the IOPs of
difference was statistically significant at 10.5 mo of age
(mice (n = 49 for each genotype; see Materials and Methods).
(A and B) Before the DBA/2J glaucoma damages RGCs, the optic nerves of both Bax+/+ (A) and Bax−/− mice (B) had a normal organization. The
axons appeared healthy with a clear axoplasm and darkly stained myelin
sheath. (C and D) BAX deficiency did not prevent glaucomatous optic nerve
damage. Severe degeneration involving extensive to complete axon loss and
scarring occurred in both Bax+/+ (C) and Bax−/− mice (D). The
majority of mice of both genotypes had this severe degree of damage by
12 mo of age. These experiments show that BAX is not required for
glaucomatous axon degeneration. Scale bar, 50 μm.
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Figure 2. BAX Is Not Required for Glaucomatous Optic Nerve
Degeneration

To assess the effects of Bax deficiency on optic nerve degeneration, we
analyzed PPD-stained optic nerve cross sections from Bax+/+ and Bax−/−
mice (n = 49 for each genotype; see Materials and Methods).
Figure 3. BAX Deficiency Prevents Glaucomatous RGC Death

To determine the effects of BAX deficiency on RGC death in glaucoma,
we analyzed RGC layer cells at stages with and without glaucomatous
optic nerve damage (see Materials and Methods). All shown images are
from a similar region of the superior, peripheral retina.

(A and B) In both Bax+/+ (A) and Bax−/− (B) mice without glaucomatous
optic nerve damage, the retinas appear healthy. The retinas of both
genotypes are similar except that Bax−/− mice have extra RGCs (since
BAX is important in normal developmental RGC death [39]).

(C and D) In contrast, an obvious difference was evident between the
retinas of Bax+/+ and Bax−/− mice that had all suffered severe
 glucomatous damage with 95% or more axon degeneration. As
expected, for Bax+/+ retinas (C) from eyes with 95% or more optic nerve
axon loss, there was a noticeable decrease in RGC layer cells (compare [C]
to [A]). In contrast, retinas from Bax−/− mice with correspondingly
damaged optic nerves (D) had suffered no obvious loss of RGC layer cells
(compare [D] to [B]). This suggests that BAX is required for RGC death in
DBA/2J glaucoma. As is well established for both RGCs and other
neurons, Bax−/− RGCs that survive without axons have a shrunken
morphology [38,82]. This is clearly evident in the Bax−/− glaucomatous
mice (D).

(E) RGC layer cell counts for eyes with 95% or more axon degeneration
confirmed that BAX is necessary for RGC death in this glaucoma. To allow
comparison between genotypes, the percent of surviving cells is shown
(% soma in mice with 95% or more axon loss compared to mice of the
same genotype without glaucomatous damage). At 12 mo of age, Bax+/+ mice had 61.4% ± 3.8% of their RGC layer cells remaining, Bax−/− mice
had no appreciable cell loss (101% ± 5.3%). The RGC layer cells of Bax−/−
mice were also protected (89.2% ± 5.7%). The p values comparing
differences in cell counts between nonglaucomatous and very severely
glucomatous (≥ 95% axon loss) eyes of the same genotype were:
Bax+/+, p < 0.001; Bax+/−, p = 0.207; Bax−/−, p = 0.426. No cell loss was
seen in Bax−/− mice even out to 18 mo (94.8% ± 4.4% cells surviving; p = 0.524 compared to nonglaucomatous Bax−/− mice). These findings show
that BAX gene dosage has an important effect on the susceptibility of
RGCs to glaucomatous death. Scale bar, 50 μm.
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Although iris damage was similar in mice of all three
genotypes, Bax genotype did have an effect on IOP. The peak
period of IOP elevation in DBA/2J mice is from 9 to 12 mo of
age, with the IOP distribution clearly shifting upward
between 3 and 12 mo of age. We monitored IOP at key ages (9,
10.5, and 12 mo). Surprisingly, Bax−/− mice tended to have
lower IOP than either Bax+/− or Bax+/+ mice at 9 mo, and the
difference was statistically significant at 10.5 mo of age
(Figure 4). In contrast, the IOPs of Bax+/− and Bax+/+ mice
were not different. Because of the lower IOP in Bax−/− mice,
we analyzed IOP in 4-mo-old mice of each genotype and
found no differences (average ± SEM, number of eyes examined:
Bax+/+, 13.54 ± 0.45 mm Hg, n = 22; Bax+/−,
13.77 ± 0.33, n = 22; Bax−/− 13.46 ± 0.34, n = 20; p > 0.5).
in IOP elevation was significant at 10.5 mo (\( p < 0.001 \)). The width of the horizontal line (black in [A], gray in [B]) represents the mean IOP ± SEM of a group of wild-type preglaucomatous DBA/2J mice. In both 9- and 10.5-mo-old Bax+/+ and Bax−/− mice that were 3 mo old. The degree of IOP elevation, however, was altered in Bax+/− mice. In both 9- and 10.5-mo-old Bax+/− mice, the average IOP was less than that of Bax+/+ and Bax−/− mice. This reduction in IOP elevation was significant at 10.5 mo (\( p < 0.01 \)). The IOP of Bax+/− mice did not differ from wild type at either 9 or 10.5 mo (\( p > 0.82 \)). By 12 mo, there was no difference in IOP between mice of any genotype (\( p > 0.25 \)).

This result indicates that BAX deficiency does not alter baseline IOP but does have an effect as the IOP increases to glaucomatous levels in older mice.

The lower IOP insult in Bax−/− mice does not account for the survival of their RGCs. This conclusion is supported by the normal RGC numbers remaining in Bax+/− mice with indistinguishable IOP from Bax+/+ mice. Previous studies have shown that BAX deficiency allows RGC survival following axotomy or optic nerve crush [13]. By contrast, even when neuroprotective treatments are administered, only a small number RGCs survive in the short term (4-6 wk) in Bax+/+ mice exposed to severe axon trauma [51,52]. Thus, there is no reasonable explanation for the finding of prolonged survival of RGCs that have no axons other than that BAX is a necessary RGC-intrinsic molecule for apoptosis in this glaucoma model.

**Bax Deficiency Delays Axon Degeneration**

Although we have shown that axon degeneration is not dependent upon BAX, our results clearly identify BAX as an endogenous susceptibility factor for both RGC death and axonal degeneration in DBA/2J glaucoma. As discussed above, complete or partial BAX deficiency had a profound rescuing effect on RGC cell bodies. Importantly, decreasing functional Bax gene dosage also decreased susceptibility to glaucoma by delaying the progression of axon damage (Figure 5). At 10.5 mo of age, the majority of Bax+/+ mice had moderate or severe optic nerve damage (see Materials and Methods), with only 20% being mildly affected. In contrast, 53% of Bax+/− and 44% of Bax−/− mice were only mildly affected at 10.5 mo of age. At 12.0 mo of age, the distribution of optic nerve damage was indistinguishable among mice of the three Bax genotypes (Figure 5). Since mice of each Bax genotype were littermates that were housed in the same cages throughout aging, these results provide compelling evidence that decreasing BAX levels delays optic nerve damage.

The delay of optic nerve damage in Bax+/− mice (note: Bax+/− mice had similar IOP insults to Bax+/+ mice) suggests that partially decreasing BAX levels in RGCs protects RGC axons. However, since complete BAX deficiency limited IOP elevation, a further protective effect of BAX deficiency by lowering IOP is also possible and may explain the trend toward greater axonal protection in Bax−/− mice. Thus, it is possible that either low-expressing or low-activity alleles of BAX may affect glaucoma susceptibility both by limiting and/or delaying IOP elevation and by directly protecting RGCs from damaging effects of harmful high IOP.

**An Integrated Approach Supports a Role of Direct Optic Nerve Injury in Glaucoma**

Comparing the specific pathways active in glaucomatous RGC death to the pathways induced by acute, experimental manipulations can provide information about the initial insult(s) to RGCs in glaucoma. N-methyl-D-aspartate (NMDA) receptor-mediated excitotoxic injury and direct axon injury are two insults that have been proposed to kill RGCs in glaucoma. Acute experimental procedures can be used to mimic these insults. Intraocular NMDA injection is used to mimic excitotoxic RGC insult, and controlled optic nerve crush is used to mimic a direct axon insult [53,54]. To assess the likely roles of these insults in a spontaneous glaucoma, we subjected preglaucomatous DBA/2J mice of differing Bax genotypes to these procedures. This allowed direct comparison of RGC death induced by these distinct excitotoxic and axonal insults to the naturally progressing glaucoma (Figure 6) in a single genetic context. Bax genotype had absolutely no
**Discussion**

**BAX-Mediated Apoptosis Is Important in an Inherited Glaucoma**

Our findings provide important new information about RGC injury and death in glaucoma. BAX deficiency completely prevents RGC death in DBA/2J mice. These results conclusively demonstrate that apoptosis plays a pivotal role in this inherited model of glaucoma. BAX is the first molecule shown to be completely necessary for RGC death in any glaucoma. Considering the protection we demonstrate in this mouse model, it is worth assessing BAX pathways as important targets for new treatments in human glaucoma.

**Distinct Pathways Mediate RGC Death and Axonal Degeneration in Glaucoma**

Intrinsic axonal degeneration pathways have recently been identified [55,56]. The molecular components of these pathways appear to be distinct from those active in classical somal apoptosis [57,58]. Thus, the different compartments of a neuron can degenerate by different molecular processes. In glaucoma, it is not clear whether the same or different degeneration pathway(s) are activated in the cell body and axon. Our study demonstrates that BAX is required for RGC death but not for RGC axonal degeneration in DBA/2J glaucoma. This indicates that the axonal degeneration pathway is distinct from apoptosis in this inherited glaucoma. Our findings clearly demonstrate that axon degeneration is not a consequence of RGC death, since severe axon degeneration occurred in Bax<sup>-/-</sup> mice without RGC death. It is not yet clear whether the RGC apoptosis and axonal degeneration pathways have some common features or are completely distinct. However, for the design of therapeutic strategies for human glaucoma, our studies suggest that both apoptotic and axonal degeneration pathways should be considered.

**Alternative Glaucoma Hypotheses**

The initial RGC compartments that are insulted in glaucoma, as well as the nature of the damaging insults that induce degeneration, are not completely clear. In the excitotoxic hypothesis of glaucoma, elevated IOP leads to...
The elevated intraocular glutamate levels [59]. The elevated glutamate levels are proposed to cause excessive stimulation of glutamate receptors (NMDA type), leading to increased intracellular calcium levels and RGC death. A different glaucoma hypothesis involves direct optic nerve injury. In this hypothesis, high pressure places stress on the optic nerve as the nerve exits the eye through the lamina cribrosa [60]. Important studies report that the first damage to RGCs is evident in the axon segment near the lamina cribrosa in the optic nerve head [61,62], so it was suggested that this is the first site of IOP-induced insult (see Quigley [60]). Although it definitively shows local axonal dysfunction, the occurrence of initial damage in this region does not conclusively indicate that this is the first or only site of neuronal insult. Because of optic nerve head architecture and the stress at the lamina cribrosa, it is conceivable that the axon segment at the lamina cribrosa may take substantial resources to maintain, especially when IOP is elevated. Somal stress may decrease available resources for axon maintenance and repair. Therefore, somal stress or damage may contribute to the abnormalities observed in the optic nerve head. As a group, Bax+/− mice had an indistinguishable IOP insult compared to Bax+/+ mice, but their RGCs did not undergo pressure-induced cell death. Importantly, RGC axonal degeneration was delayed in these Bax+/− mice. Therefore, our data imply that shielding the RGC cell bodies has a protective effect against axon degeneration.

Direct Optic Nerve Damage Resembles Glaucoma

To provide insight to the nature and location of the damaging insults that occur in glaucoma, we compared the effects of BAX deficiency on RGC death in inherited glaucoma to RGC death induced by either direct optic nerve injury or excitotoxicity (all in the genetically uniform DBA/2J strain). Intracocular MDA injection was used to model excitotoxic RGC death, and controlled optic nerve crush was used to mimic direct optic nerve damage [53,54]. Unlike the DBA/2J glaucoma, our experiments show that the excitotoxic insult does not require BAX to induce RGC death. Although these experiments cannot rule out the possibility of an intrinsic excitotoxic mechanism, these results do not support a role of MDA receptor-mediated excitotoxicity as a primary cause of glaucomatous RGC death. Similar to the DBA/2J glaucoma, RGC death following optic nerve crush requires BAX, and both Bax+/− and Bax−/− mice are profoundly protected. Along with our demonstration of an axon intrinsic degeneration pathway, these results further support the hypothesis [60] that direct optic nerve and axon injury is an important pathogenic component leading to RGC death in glaucoma.

Bax Can Modulate Neuronal Susceptibility in Glaucoma

Individual patients have different levels of susceptibility to glaucomatous RGC death [2,63]. Our experiments clearly identify Bax as an important modulator of neuronal susceptibility in DBA/2J glaucoma. BAX deficiency prevented RGC death and delayed optic nerve degeneration in both Bax+/− and Bax−/− mice. These results suggest that the use of BAX inhibitors could potentially be used to delay glaucomatous vision loss. In situations where BAX is important, pharmacologically suppressing BAX activity may significantly slow the progression of glaucoma. Since RGCs were maintained for an extended period after axon degeneration in Bax−/− mice, treatments that inhibit BAX pathways may allow long-term preservation of RGC cell bodies. Such treatments may allow the RGCs of patients to be stored in their own retinas until future treatment strategies are developed that can stimulate axonal growth and restore vision.

Complete BAX Deficiency Limits IOP Elevation

In addition to implicating BAX as a target for direct neuroprotective treatments, the lower IOP of Bax−/− mice suggests that BAX inhibition may delay or limit IOP elevation. These results suggest that apoptotic death of cells affecting aqueous humor drainage contributes to IOP elevation, at least in secondary glaucomas where the drainage structures are insulted by pigment and cell debris. In a previous study assessing neuroprotection by an apoptosis inhibitor in a rat model of glaucoma, the treated rats had lower IOP than the other group [25]. Although not a conclusion of this rat study, the IOP data support a role for apoptosis in IOP elevation. In humans, cell death has been speculated to contribute to common forms of glaucoma (due to loss of drainage structure cells in old individuals and at late stages of glaucoma [64,65]). However, a primary role for ocular drainage pathway cell death during IOP elevation is not clearly established. Importantly, a recent study convincingly demonstrated endoplasmic reticulum stress and subsequent cell death in primary cultures of drainage pathway cells expressing human glaucoma mutations [66]. Together with our finding that complete BAX deficiency delays IOP elevation in a glaucoma setting, these results strongly support further investigation of apoptotic pathways and effects of antiapoptotic drugs on IOP in human glaucoma.

BAX Is a Candidate Human Glaucoma Susceptibility Gene

The profound protection against RGC death and the delay in axon degeneration in Bax−/− mice together suggest BAX as a candidate human glaucoma susceptibility gene. It is important to note that we considered the possibility that a closely linked gene that was transferred from the 129/SV strain (in which the Bax mutation was generated) hitchhiked into the DBA/2J background along with Bax and explains the protection in heterozygotes. We conclude that this possibility is remote on the basis of the following observations. First, the RGCs of wild-type mice of the parental 129/SV strain are not protected from optic nerve crush. Bax heterozygosity protected the animals from both optic nerve injury and axon injury is an important pathogenic component leading to RGC death in glaucoma.
development and progression of some but not other diseases [67–72]. Other factors that control BAX expression could also be important. Lower levels of BAX are associated with a worse prognosis for some types of cancer [73]. Our findings in Bax−/− mice support the hypothesis that quantitative variation in the level of BAX gene product may alter the prognosis of glaucomatous damage in individuals with high IOP. Although further studies are needed to assess this possibility, quantitative variation of BAX activity among human patients may have a substantial effect on susceptibility and disease progression. It is possible that lower-activity alleles may result in slower or less severe damage, whereas high-activity alleles may be detrimental. Characterization of BAX alleles may have important predictive value for disease progression.

Materials and Methods

Animals and husbandry. Mice were housed in a 14 h light to 10 h dark cycle under previously described conditions [74]. The Jackson Laboratory (Bar Harbor, Maine, United States) pathology surveillance program regularly screened for pathogens. All experiments were conducted in accordance with the Association for Research in Vision and Ophthalmology’s statement on the use of animals in ophthalmic research and were approved by our institutional animal care and use committees. Both male and female mice were used. For each age group and genotype, approximately equal numbers of males and females were used. A Bax null allele (Baxem1SjK [44]; herein referred to as Bax−) was backcrossed to B6.129X1-Scr(Cg)-littermates. All three genotypes (Bax+/-, Bax+/+, Bax−/−) were sectioned at 5 μm thickness and subjected to a modified double label protocol that involved in situ end-labeling (equivalent to a TUNEL assay) of fragmented DNA (using BODIPY fluorophores; Molecular Probes, Eugene, Oregon, United States) and detection of condensed chromatin (with the dimeric cyanine dye YOYO-1; Molecular Probes) as published [75]. Samples were analyzed with a confocal microscope. Conventional TUNEL assays were performed as previously described [13] and conducted on the following numbers of DBA/2 mice of each age group: 7 mo (six), 8–9 mo (ten), 10–11 mo (16), 12–15 mo (nineteen, and 15–18 mo (sixteen). Five to 12 controls per genotype at each age, and for each disease model, were intercrossed to produce Bax+/+, Bax+−, and Bax−/− littersmates. All three genotypes were housed together and analyzed simultaneously. DBA/2J mice were from our colony (Sj) that was initiated with mice purchased from The Jackson Laboratory. DBA/1J mice were obtained from The Jackson Laboratory.

Cell death related assays. Eyes from DBA/2J or control DBA/1J mice were fixed and retinas were flat-mounted (strain and genotype matched non-related DBA/2J or control DBA/1J mice and more than 15 control mice of mixed genetic background of each genotype at each age), an optic nerve rating scale was used for the glaucoma progression study (see Figure 5). The indicated damage levels are readily distinguishable upon inspection of the nerve without counting. Nevertheless, axon counts were performed on at least eight randomly selected nerves of each disease grade to provide quantitative information about these distinct stages of disease (see below). Two investigators (masked to genotype, age, and the damage level assigned by the other investigator) assigned a damage level to each nerve. The two investigators assigned the same grade more than 95% of the time (321 out of 335 nerves). For the nerves on which the initial two investigators differed, a third (masked) investigator was utilized. The third investigator’s grade always agreed with one of the initial grades, and the most common assigned grade was used. The number of nerves of each genotype assessed at each age were as follows. For 10.5 mo, Bax+−/− = 49, Bax+/+ = 62; Bax−/− = 58; for 12 mo, Bax+−/− = 52, Bax+/+ = 17, Bax−/− = 46.

The damage levels and typical numbers of normal axons present at each stage (determined through axon counts by an investigator masked to damage grade) follow. The representative axon counts were determined for randomly selected nerves of each grade using the counting procedure described above. In mildly affected nerves, there was very mild or no damage, with healthy axons having a clear margin of demarcation. The axon number was significantly different between optic nerves of each genotype at each age, and for each damage model. Comparisons.

Optic nerve damage. Optic nerves were dissected, processed, embedded in plastic, sectioned and stained with paraffin/enhanced (PDD) as previously described [76], except that the staining time was increased to 35 min. Endothelial and myelin staining with PDD stains all myelin sheaths, but differentially stains the axoplasm of sick or dying axons darkly. Counts of normal-appearing axons were performed using established unbiased counting methods. Prior to beginning axon counts, the optic nerve was maximally magnified, and its cross-sectional area was automatically calculated. Magnification of the same nerve section was increased to 1,000x, and a total of 20 fields at 1,000x were electronically collected. The fields were spaced in a regular fashion across the entire nerve, taking care to avoid field overlap so that the same area was not counted more than once. The 20 fields in each collected picture were on the computer screen so that only the final picture was visible to the operator. For nerves with a large number of axons (mildly and moderately affected nerves), a rectangular box that contained a minimum of 200 axons was then drawn on the twentieth image. For nerves with severe axon loss, a larger box was drawn so that a significant proportion of the nerve could be counted. The software program then “cut” a rectangle centred at the same location in all 20 images. Since the operator could only see the top image, this removed some of the possibility of unconscious operator bias and made the selection of axons to be counted random. Axons were counted manually and marked using the computer. The program tracked the total area counted and the total axon count for all 20 images. The total counted area averaged 12.1%, 14.2%, and 20.5% of the total nerve area for mildly, moderately, and severely affected nerves, respectively. The final counts were calculated as number of axons/axon length.

With this approach, the nerves with 95% or more axon loss were selected for RGC counts by comparing the remaining axon number to the average for unaffected nerves of the same genotype. Because of the large number of mice (approximately 50–70 mice of each genotype at each age), an optic nerve rating scale was used for the glaucoma progression study (see Figure 5). The indicated damage levels are readily distinguishable upon inspection of the nerve without counting. Nevertheless, axon counts were performed on at least eight randomly selected nerves of each disease grade to provide quantitative information about these distinct stages of disease (see below). Two investigators (masked to genotype, age, and the damage level assigned by the other investigator) assigned a damage level to each nerve. The two investigators assigned the same grade more than 95% of the time (321 out of 335 nerves). For the nerves on which the initial two investigators differed, a third (masked) investigator was utilized. The third investigator’s grade always agreed with one of the initial grades, and the most common assigned grade was used. The number of nerves of each genotype assessed at each age were as follows. For 10.5 mo, Bax+−/− = 49, Bax+/+ = 62; Bax−/− = 58; for 12 mo, Bax+−/− = 52, Bax+/+ = 17, Bax−/− = 46.

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Ganglion cell death. Eyes were fixed and retinas were flat-mounted and Nissl-stained with cresyl violet using a modification of the technique reported by Stone [80]. Retinal ganglion cells make up approximately 40%–60% of the neurons in the ganglion cell layer of the mouse retina. The ganglion cell layer is distinguished from the other resident neuron in the ganglion cell layer (the displaced amacrine cell) based on cellular morphology [53,81]. This is especially true during disease, when morphology and marker expression can change dramatically. Consequently, cell loss was distinguished from the change in cell number compared to control eyes (strain and genotype matched non-glaucomatous eyes for the spontaneous glaucoma experiments and the contralateral nonmanipulated eye for the controlled crush and
excitotoxic experiments). RGC density varies greatly with respect to retinal location. Therefore, two 40x fields were counted in each retinal quadrant and care was taken to ensure that the fields were the same distance from the periphery. For each individual eye, the eight counts for each retina were averaged. To assess RGC survival in the spontaneous glaucoma, retinas from eyes with very severely affected nerves that had fewer than 5% surviving axons were compared to retinas from unaffected eyes without glaucomatous nerve damage. RGC number was counted in approximately eight severely affected eyes and eight unaffected eyes of each genotype, except for unaffected control +/+ mice (live eyes) and 18 mo unaffected Bax−/− mice (four eyes).

NMDA injections and controlled optic nerve crush. These experiments were performed as described previously [53]. For NMDA injections, 2 μl of an 80 mM solution of NMDA in balanced saline solution was injected intravitreally into one eye of each mouse using a glass micropipet. After 4 d the eyes were harvested and cells counted. Data were collected from nine Bax−/−, nine Bax+/−, and seven Bax+/+ mice. In each paradigm, cell loss was measured relative to the cell number present in the control eye of each mouse examined.

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