INTRODUCTION

Glaucoma, a group of optic neuropathies, is the leading neurodegenerative cause of blindness [1]. It is commonly characterized by a slow progressive degeneration of the axon and then the soma of retinal ganglion cells [2, 3]. Among patients with diverse glaucoma types worldwide, primary open-angle glaucoma with elevated intraocular pressure (IOP) is the most common; however, the prevalence of normal tension glaucoma with IOP in the normal range is increasing [4]. The pathogenesis of glaucomatous optic neuropathies is therefore most likely explained by two distinct mechanisms, the mechanical or pressure theory and the vascular theory [5, 6]. IOP has been confirmed as the only treatable risk factor for glaucoma till now. Elevated IOP contributes to progressive...
damage of the axon and soma of retinal ganglion cells and eventually to visual field loss [3, 7]. Non-IOP-related mechanisms such as vascular abnormality may play an important role in the pathogenesis of normal tension glaucoma.

It is widely accepted that if glaucoma is recognized in its early stages, subsequent visual loss can be limited or prevented via currently available treatments. Most cases of glaucoma, however, given its clinically subtle early signs, are not disclosed until vision has already been irreversibly lost [3]. Currently, efforts are being made to screen glaucomas underlying mechanisms and other treatable risk factors, aiming to enable early diagnosis and detection.

Thereby, the present study aims to discover crucial episodes useful in early detection of glaucoma, focusing on the retina itself. For this, the spatiotemporal sequence of retinal degeneration caused by elevated IOP was functionally and morphologically assessed. We used a rat model of primary open-angle type chronic glaucoma induced via unilateral episcleral venous cauterization, according to the diagnostic criterion that primary open-angle glaucoma is a chronic, generally bilateral, but asymmetrical disease [3]. Functional assessment through flash electroretinography (ERG) was conducted to detect alteration in the electric reactivity of the conducting neurons in the photo-transduction pathway in the glaucomatous retinas: that of both photoreceptors and ON bipolar cells was evaluated based on changes in a- and b-wave amplitudes [8-11], and that of the RGCs based on changes in photopic negative responses (PhNRs) [10, 11]. Morphological assessment of retinal degeneration was largely performed via immunohistochemical analyses using \( \text{Ca}^{2+} \) homeostasis-related proteins, such as calcium-binding proteins and the calcium-sensing receptor. The divalent cation \( \text{Ca}^{2+} \) works as a secondary messenger in many signaling pathways, such as membrane excitability and synaptic transmission, in the central nervous system including the retina [12]. Calcium-binding proteins (CaBPs) bind calcium to increase the cells buffering capacity as a constituent of the termination mechanisms of intracellular calcium signal, including calcium buffering, extracellular extrusion, and intra-organelle sequestration. CaBPs are usually expressed in a cell-specific manner and thereby used for identification of specific neuronal types, especially types of amacrine cells, a type of associated retinal neuron. In our study, the immunohistochemical characteristics of CaBPs were utilized for determining changed amacrine cells in glaucomatous retinas, in addition to indistinct oscillatory potentials in flash ERGs. The calcium-sensing receptor (CaSR), as a member of the G-protein-coupled receptor C family, is activated by local extracellular \( \text{Ca}^{2+} \) or by other physiologically relevant polycationic molecules as a first messenger, and therefore currently accepted to play a role in maintaining local ionic homeostasis in the brain [13, 14]. Despite ubiquitous distribution of CaSR in the brain [15], there is nearly no report on CaSR expression in the retina. We demonstrate, using immunohistochemical analysis, CaSR expression in the retinal ganglion cells and its alteration pattern under ocular hypertension, followed by electron microscopy of CaSR-immuno-stained retinal preparation for the identification of apoptotic retinal ganglion cell death in glaucomatous retinas.

MATERIALS AND METHODS

Animals

Sprague-Dawley rats aged 8 weeks were used. All experimental procedures complied with the regulations of the Catholic Ethics Committee of the Catholic University of Korea, which conforms to the US National Institute of Health (NIH) guidelines for the Care and Use of Laboratory Animals (NIH Publications No. 80-23), revised in 1996. Animals were maintained at a 12 h light/dark cycle and fed standard chow.

Elevation of IOP

Elevated IOP was induced via venous cauterization. Briefly, animals were anesthetized with a mixture of tiletamine plus zolazepam (Zoletil; Virbac, France)/xyazine hydrochloride (Rompun; Bayer, Germany) (50/15 mg/kg ip injection). Proparacaine (0.5% concentration) was topically eye-dropped. Three of four episcleral veins in the right-side eyeball were cauterized with a 30-gauge tip cautery (Bovie Medical Co., USA) under a surgical microscope (Olympus, Tokyo, Japan). Animals were grouped and cared for 1 and 3 days, and 1, 2, 4, and 8 weeks after operation. Twenty animals each were assigned for each experimental time point and, 15 animals for normal control and 5 animals for aged-matched control.

IOP measurements

IOP was cumulatively measured using a rodent tonometer, TonoLab (Tono-Pen XL<sup>®</sup>, Medtronic Solan, Jacksonville, FL, USA) in glaucoma and contralateral control eyes at all time points after operation. Five consecutive readings via a single measurement were averaged and corrected based on a linear equation obtained from IOP calibration with the pressure transducer. Data were analyzed by averaging the IOP measurements for each time point.

ERG recording

Rats were dark-adapted overnight before ERG recordings. Under dim red light, rats were anesthetized with intraperitoneal injection of zolazepam and xylazine, and placed in a Ganzfeld dome laying on the stage to ensure a stable position for recording. After moistening with a hydroxypropyl methylcellulose gel, gold-ring
contact-lens electrodes were placed on the eyes and, two reference electrodes and one ground electrode, subcutaneously in the ears and in the tail, respectively. ERGs were recorded with a universal electrophysiological system (UTAS-3000 MF; LKC Technologies, Gaithersburg, MD, USA). Luminescence intensity was calibrated by the manufacturer and controlled by computer system. Scotopic and photopic ERG responses were each obtained from both eyes simultaneously. Scotopic responses are the mechanisms that are specifically adapted for dim or night vision [16]. Rats were first tested for rod-mediated scotopic ERG under dark adaptation exposed to flash intensity of 0.99 cd·s/m². To obtain photopic responses, light-adapted rods with a saturating background (30 cd/s/m²) for 10 min presented flashes of intensity of 75.36 cd·s/m² (4 dB). For each condition, 3–5 responses to the flash stimuli were averaged, with a stimulus interval of 15 s. A-wave amplitude was measured from the baseline to the maximum a-wave peak, and that of b-wave, from the maximum a-wave peak to the maximum b-wave peak. PhNR was recorded under stimulus conditions: a brief 0 dB blue flash (peak wavelength 470 nm) at an intensity of 10 cd/m² against a blue background (peak wavelength 470 nm) of 10 cd/m² (photopic units). PhNR amplitude was measured from the baseline to the trough of the negative peak following the b-wave. The peak latency of the PhNR was measured as the elapsed time between the instant of the flash and the point of maximal descent following the b-waves.

**Tissue preparation**

Eyeballs were enucleated under anesthesia with 4% chloral hydrate. The posterior halves of the globes were immersed in 4% paraformaldehyde in 0.1 M phosphate buffer (PB), pH 7.4. Whole retinas were dissected and immersed in the same fixative for 2 h. After fixation, retinas were immersed in 30% sucrose, refrigerated overnight, and then flash-frozen in liquid nitrogen and stored at -70°C for preservation.

**Immunohistochemistry**

Retinal pieces were trimmed out from the central portion of the superior quadrant and rinsed with 0.01 M phosphate buffered saline (PBS), pH 7.4. After thorough rinsing, retinal pieces were embedded in 4% agar and cut into 40-µm-thick vertical sections. The retinal sections were collected in culture wells and processed for immunofluorescent microscopy. To block nonspecific binding sites, sections were treated with buffer B (1% BSA, 0.2% gelatine bovine, 0.05% saponin/in PBS) for 3 h on ice. Sections were then incubated with antibodies against monoclonal mouse anti-Calcium Sensing Receptor (CaSR, Sigma-Aldrich, St.Louis, MO, USA; dilution 1:3000) and one of following antibodies: polyclonal rabbit anti-calbindin D28K (Swant, CH-1723 Marly, Switzerland; dilution 1:2000), polyclonal rabbit anti-calretinin (Millipore, Temecula, CA, USA; dilution 1:5000), polyclonal rabbit anti-parvalbumin (Swant; dilution 1:5000) overnight at 4°C. Immunoreactivity was visualized with Alexa Fluor 488 donkey anti-mouse and rabbit conjugated IgG (Life technologies, Grand Island, NY, USA; dilution 1:2000). After rinsing in PBS, sections were mounted on a glass slide with mounting medium.

**Confocal microscopy**

Immunofluorescent staining was evaluated via confocal laser-scanning microscopy (LSM 510 Meta, Carl Zeiss Co. Ltd., Germany). Fluorescent images were captured with green laser (excitation 488 nm, emission 490–555 nm) at 400× magnification power. Captured images were converted to JPEG.

**Western blot analysis**

Western blotting was performed on homogenized retinal extracts. Protein concentration in each sample was assayed using the Pierce™ BCA protein assay. Duplicate sets of protein standards containing 0, 0.05, 0.1, 0.2, 0.25, 0.5 and 1 mg/ml BSA were assayed using the same method. Tissue-sample aliquots corresponding to 25 µg of total protein were heated at 100°C for 10 min with an equivalent volume of 2× sample buffer (containing 4% SDS and 10% mercaptoethanol) and loaded onto 5–12% SDS-polyacrylamide gels. The proteins were electro-transferred onto a polyvinylidene difluoride membrane in Tris-glycine-methanol buffer and treated for 1 h in a blocking solution. The membrane was then incubated with antibodies in the blocking solution overnight at 4°C. Primary antibodies were the same as those used in immunohistochemistry; mouse monoclonal anti-β-actin (Sigma-Aldrich, dilution 1:100,000) was used as the control. The antibody-treated membrane was rinsed with 0.1% Tween-20 in PBS for 30 min, and then, incubated with biotinylated donkey anti-mouse IgG or anti-rabbit IgG (Vector Laboratories, Burlingame, CA, USA; dilution 1:1,000) for 2 h. The blot was washed, processed for analysis using an enhanced chemiluminescence (ECL) detection kit (Amersham, Arlington Heights, IL, USA), and visualized using Fujifilm Image Reader LAS-4000 (Fujifilm, Tokyo, Japan). Each experiment was repeated at least thrice, and representative blots are shown. Densitometric analysis was performed using Multi Gauge V3.0 (Fujifilm). Data are presented as mean±SE and statistical significance was determined using Student’s t-test.

**Terminal deoxynucleotidyl transferase dUTD nick end labeling (TUNEL) assay**

Apoptosis was evaluated using the TUNEL method. Retinal
sections (40-µm-thick) were embedded in 4% paraformaldehyde, washed and stained (In Situ Cell Death Detection Kit, Fluorescein; Roche Applied Science, Mannheim, Germany). Briefly, after washing with PBS for 30 min, sections were incubated with 0.1% Triton X-100 / 0.1% sodium citrate solution for 2 min on ice for better permeability. Sections were then incubated with terminal dUTP enzyme at 37°C for 1 h for tailing reaction, and counterstained with 4',6-diamidino-2'-phenylindole (DAPI) for cell nuclei. The reaction was examined using a confocal microscope (LSM 510 Meta).

Electron microscopy

To examine the ultrastructure of normal and 8-week glaucoma retinas, CaSR-immunostained retinal sections were fixed with 2.5% glutaraldehyde (in PB), washed with PB, and post-fixed with 1% osmium tetroxide (in PB). Sections were then washed with PB and dehydrated in an ethanol gradient. After dehydration, sections were infiltrated with Epon 812, flat-mounted on transparent film, and polymerized by curing at 60°C. Ultrathin sections (70~90 nm thickness) were collected on a one-hole grid and the ultrastructure was observed using an electron microscope (JEM-1010; JEOL, Tokyo, Japan).

RESULTS

IOP was elevated to 20 mmHg on the first day after operation and thereafter sustained in the range of 17~20 mmHg throughout the experiment (Fig. 1). IOP and ERG were cumulatively surveyed throughout all experimental time points to elucidate the temporal correlation between IOP elevation and functional damage on each component of ERG. Immunochemical changes were studied at the given time points from 1 week post-operation for spatial correlation between elevated IOP and its morphological impact on specific neuronal populations.

Electroretinography (ERG) of scotopic responses

ERG recordings of scotopic responses in glaucoma were depicted together with that of a normal retina aged 8 weeks (Fig. 2). In all time points, ERGs of scotopic response in glaucoma retinas showed clear reduction in a- and b-wave amplitudes compared to those of normal retina. Oscillatory potentials before the b-wave peak, even though very slight, were also reduced. At 8 weeks post-operation, ERG of glaucoma retinas showed reduced amplitudes than that of normal retina.

ERG of photopic responses

ERG recordings of photopic responses in glaucoma retinas were compared at all time points with those of normal retina (Fig. 3). ERGs of photopic responses in all retinas showed negligible a-wave amplitudes and oscillatory potentials. However, b-wave amplitude in glaucoma retinas was slightly reduced at 1 and 3 days, thereafter slightly recovered and sustained.

Photopic negative responses (PhNRs)

PhNRs in glaucoma retinas were compared with those in normal control retina at each time point (Fig. 4a). PhNRs amplitude in glaucoma retinas significantly reduced from 1 day, recording the lowest at 2 weeks (p<0.01), to 8 weeks (p<0.05) (Fig. 4b).

CaSR and CaBPs immunoreactivity

CaSR strongly immunolocalized into the neurons in the ganglion cell layer (GCL) and nonspecifically into both the neurons in the inner nuclear layer (INL) and the rod and cone layer at normal state (Fig. 5a). CaSR immunoreactivity in the GCL showed no large change in 1-week glaucoma retina, but clearly reduced after 2-week glaucoma especially in ganglion cells (Fig. 5b~d). Calbindin immunolocalized into the horizontal cells in the outermost part of the INL, into the amacrine cell soma in the inner half of the INL and their ramifying processes in the inner plexiform layer (IPL), and to the displaced amacrine and ganglion cells in the GCL (Fig. 5c). Calbindin immunoreactivity showed no large change in 1- and 2-week glaucoma (Fig. 5f and g); however, it remarkably reduced in 8-week glaucoma in all immunoreactive neuronal types (Fig. 5h). Calretinin immunolocalized into the amacrine cells in the INL and their processes ramified in three laminae in the IPL, and into the neurons in the GCL (Fig. 5i). In 1- and 2-week glaucoma, there was no large change in calretinin immunoreactivity (Fig. 5j and k), but calretinin immunoreactive amacrine cells and neurons in the GCL slightly reduced in cell number and immuno-
reactivity in 8-week glaucoma (Fig. 5l). Parvalbumin immunolocalized to AII amacrine cells in the INL and their processes in the IPL (Fig. 5m). Parvalbumin immunoreactivity in the whole-cell profile of AII amacrine cells did not reduce in 1-week glaucoma (Fig. 5n). However, immunoreactivity clearly reduced in both lobular appendages in OFF lamina of the IPL and the dendrites near the GCL after 2-week glaucoma (Fig. 5o and p).

**CaSR and CaBPs expression**

CaSR and CaBP expression levels were evaluated via western blot.
in glaucomatous eyes. CaSR expression levels began to increase in 1-week glaucoma and peaked at 8 weeks (Fig. 6a and b). Changes in calbindin and calretinin expression levels present a shallow concave curve and a mild convex curve showing a bottom and a peak at 2 weeks, respectively (Fig. 6c and d). Parvalbumin expression levels showed no large change throughout the glaucomatous retinas (Fig. 6e).

Neuronal apoptosis
Neuronal apoptosis was assayed using TUNEL (Fig. 7) and

Fig. 3. Photopic ERGs were evaluated. In (a), representative ERGs of scotopic responses of glaucoma (G, red colored waveforms) at each time point (G1d, G 1day; G3d, G 3day; G1w, G 1week; G2w, G 2week; G4w, G 4week and G8w, G 8week) are compared with that of normal control retina (Normal, black colored waveform). In (b), a- and b-wave amplitudes in normal retina and glaucoma retina at each time point were evaluated with a linear graph.
Fig. 4. Photopic negative response (PhNR) was evaluated. In (a), representative PhNRs of glaucoma retina (G) at each time point (G1d, G 1day; G3d, G 3day; G1w, G 1week; G2w, G 2weeks; G4w, G 4weeks; G8w, G 8weeks) are compared with that of normal control retina (N). In (b), PhNR amplitudes in the retinas at normal state (Normal) and at each time point of glaucoma retinas (1d, 3d, 1w, 2w, 4w, 8w) are compared. PhNR amplitude at 2 week glaucoma shows significant reduction (p<0.01), compared to normal retina. *p<0.05 and **p<0.01, paired Student’s t-test.

Fig. 5. Confocal microscopic analysis of normal and glaucoma retinas processed for CaSR and calcium-binding proteins immunohistochemistry. Immunohistochemistry for CaSR (a–d), calbindin (e–h), calretinin (i–l) and parvalbumin (m–p) was performed in normal retina (Normal: a, e, i, m), and in 1 week (G1w: b, f, j, n), 2 week (G2w: c, g, k, o), and 8 week (G8w: d, h, l, p) glaucoma retinas. CaSR immunoreactivity in the neurons in the ganglion cell layer (GCL) clearly reduced in G2w and G8w retinas. Calbindin immunoreactivity in amacrine cells in the inner nuclear layer (INL) gradually reduced in glaucoma retinas and that in the axons of ganglion cells increased. Calretinin immunoreactivity in amacrine cells showed no large change in glaucoma retinas. Parvalbumin immunoreactivity in the entire AII amacrine cell profile gradually reduced in glaucoma retinas. IPL, inner plexiform layer; ONL, outer nuclear layer; OPL, outer plexiform layer. Scale bar indicates 50 µm.
Dual Analyses of Glaucomatous Retina

In electron microscopy (Fig. 8). In normal control retina and experimental control retina of 8-week glaucoma, there were no TUNEL-positive cells (Fig. 7a–d). TUNEL positivity appeared in a few photoreceptors in the outer nuclear layer (ONL) and in several ganglion cells in the GCL only in 8-week glaucoma retina (Fig. 7e and f).

In electron microscopic observation of 8-week glaucoma retina, dying photoreceptors showed the classical apoptotic feature of pyknotic nucleus (Fig. 8a). Degenerating retinal ganglion cells showed necrotic features representing the pyknotic and tortious nucleus, endoplasmic reticulum with enlarged cisternae, and swollen mitochondria in the electron-dense cytoplasm (Fig. 8b).

**DISCUSSION**

The finding of this study show that elevated IOP affects the sensitivity of both photoreceptors and RGCs, and that the invoked retinal damage progresses along two paths. Pressure-induced ischemic insult initiated from the outer retina propagates to the inner retina; and pressure-injured retrograde degeneration starting from the axon of RGCs in the optic-nerve head transfers to their soma in the inner retina. Interestingly, the two paths lead to apoptotic death of the photoreceptors and necrotic apoptosis of the RGCs.

In a full-field flash ERG, the a-wave reflects a massive response from the photoreceptors [11, 12, 17], whereas the b-wave, reflects the summed activity of ON-bipolar cells [18], and an interaction of the ON-bipolar cells and the Müller cells [20]. Based on this in our study, a-wave amplitude changes that show significant reduction in scotopic and negligible reduction in photopic responses indicate that elevated IOP affects the rods and cones simultaneously. In the same manner, b-wave changes that show significant and less significant reduction in scotopic and in photopic responses, respectively, indicate that elevated pressure affects largely the rod function. In another rat glaucoma model, it has been reported that amplitude decreases were not significant prior to ~3 months of ocular hypertension [21]; and, in another model, a significant decrease in ERG amplitudes at 4 weeks was reported [22]. Together with the drastic reduction in 8-week glaucoma that we report here, all three reports show different critical time points probably due to differing modelling methods and evaluation criteria. Given our finding that apoptotic photoreceptors death with drastic a-wave amplitude reduction occurs in the last time point, it can be further deduced that sustained elevated pressure raises ischemic conditions, leading the sensitivity compromised photoreceptors to die via apoptosis. Evidence supporting this hypothesis have been sought in ischemic experiments [23–25]. Milder ischemia by permanent occlusion of the carotid arteries has been introduced to affect the outer retina more readily [23], whereas acute retinal ischemia by elevation of IOP to 120 mmHg led to marked reduction in a-wave amplitudes [24] and to apoptotic photoreceptor death [25]. Despite these results, the significant reduction in scotopic a-wave can be gener-
ally explained by the fact that rats have rod-dominant retinas, as documented by Osborne et al. [26]. There are, however, some clinical studies showing substantial glaucomatous damage to the outer retina. In flash ERGs of glaucoma patients, glaucomatous changes such as reductions and delays were larger and increased with disease severity [27], with a-wave amplitude significantly lowered [28]. With these results, the authors suggest that glaucomatous damage affects photoreceptors as well as RGCs due to both pressure stress and choroidal blood-flow insufficiency. Altogether, a-wave amplitude may be considered as a useful indicator for early detection of glaucoma degeneration.

PhNR in flash ERG has been previously suggested as an indicator in early diagnosis of glaucomatous optic neuropathy, wherein it was shown that PhNR drops even when visual sensitivity losses are mild in macaque glaucoma experiment [29]. Thereby, the authors concluded that PhNR arises from RGCs and their axons, but its delay could be mediated by glia. Another report describes significant decreases in PhNR amplitude with latency unaffected, occurring in early stages when IOP remarkably increased [11]. Largely in agreement with these results, our PhNR tracing showed significant reduction in the first week in only glaucoma and more remarkable reduction at 2 weeks in both retinas. It may therefore indicate that the earlier reduction means pressure-induced damage to the RGC sensitivity, and the later reduction in both retinas, additional damage to supportive glial structure in the optic-nerve head and its fast crossing-over to contralateral via electrical connection. The hypothesis of glial involvement was also consolidated by some morphological evidence from human glaucoma specimens and animal models. It has been found that elevation of IOP produces not only axonal changes, but also non-axonal pressure transducer, such as extracellular matrix and astrocytes, changes in the optic-nerve head [30]. On one hand, it was reported that astrocytes and Müller cells may be involved in the pathophysiology of RGC death or retinal repair in a rat model of acute ocular hypertension [31]. In addition, it has been more recently shown that astrocytes fortified by dense cytoskeletal filaments in the optic-nerve head are the target of raised IOP in rat glaucoma model [32]. Furthermore the
authors proposed that the damage in the axons is rather metabolic than mechanical, because of the localized loss of metabolic support for RGC axons from the astrocytes.

Because of the high energy requirement for crucial roles in phototransduction, photoreceptors enrich numerous mitochondria in their inner segments, while RGCs do that in the global part of their axons [33]. Considering this, elevated-IOP-induced damage seems likely to affect both types of cell, causing mitochondrial dysfunction and invariably leading to apoptotic death. In accordance with this notion, we found that light-microscopically determined apoptosis occurred in photoreceptors, in addition to RGCs. However, apoptotic features of the two cell types were quite different in electron-microscopic observation, although direct matching with light microscopy is missing. Apoptotic features of photoreceptors were classical, whereas those of RGCs, nearly necrotic. Regarding the necrotic features of apoptotic RGCs, the death event of pressure-affected RGCs may be in agreement with the recently suggested theory of necrotic apoptosis or necroptosis [22, 34-36]. According to Degterev et al. [35], necroptosis occurs when death receptor-mediated extrinsic apoptosis pathway is triggered in the absence of intracellular apoptotic signaling, showing necrotic death morphology. On one hand, it has been suggested that necrosis might be regulated, showing the appearance of controlled apoptotic processes such as mitochondrial dysfunction, enhanced production of reactive oxygen species, and ATP depletion after damage-induced lesions [22, 36]. It was also shown that when essential apoptosis-regulating proteins are inhibited, the cell-death pathway could be shifted to necrosis, thereby demonstrating that different cell-death pathways cross-regulate each other [22]. In contrast, it was shown that the activation of death receptors leads to activated NF-κB to promote cell survival or to induce apoptosis and the third pathway, necroptosis [34]. The authors [34] also identified that a death-domain-containing kinase, receptor-interacting protein (RIP) 1 participates in all three cell-fate pathways. Based on these several lines of evidence [22, 34-36], we carefully suggest that RGC death affected by elevated IOP in rat might be caused by necrotic apoptosis as adjudged by necrotic morphology and DNA breakdown. This RGC death phenomenon might be caused by two routes: one is a direct effect of pressure on RGC axon in the retina, and the other is an indirect effect of pressure-injured astrocytes supporting RGC axons in the optic-nerve head.

Morphological evaluations of three kinds of calcium-binding proteins showed slightly different change patterns in both immunoreactivity and total protein amounts. Parvalbumin immunoreactivity in AII amacrine cells [37], it may be indicated that the dendrites and ON cone bipolar cells are more susceptible than the lobular appendages and OFF cone bipolar cells, respectively, to pressure-induced damage. Calbindin and calretinin immunoreactivity in amacrine cells was clearly reduced during glaucoma, well in accordance with previously published results [38, 39]. Total protein levels of calbindin and calretinin in glaucomatous retinas, however, showed decrease and increase, respectively. In case of calbindin, the increase in protein levels in 8-week glaucoma may be due to upregulated expression not in horizontal cells but in the RGC axons. Published results that showed no change in the number of calbindin-immunoreactive horizontal cells in glaucoma [39] and in ischemic [40] retinas support this assumption. Similarly, the increase in calretinin protein levels may be caused by upregulated expression in RGC axons. Previously, it was reported that calbindin and calretinin may participate in delaying the onset of cell death [41], and have distinct function in localized calcium signaling in RGCs with clear compartmentalization, such as calretinin distribution close to the plasma membrane in the absence of calbindin [42]. Based on this, fast propagation of axonal damage to the control retina and calretinin overexpression in our study may suggest that axonal damage is worsened by neighboring cells, for example astrocytes.

The CaSR is synthesized and anchored in the rough ER, terminally glycosylated in the Golgi apparatus, and delivered to the plasma membrane to detect dynamic changes in extracellular calcium cationic concentration [43]. With regard to the CaSR's function, it is documented, on one hand, that an extracellular calcium signal produced in a neighboring cell by calcium cation mobilization could be detected in nearby cells expressing the CaSR [44], but on the contrary, that activated CaSR plays a role in the long-term control of apoptosis by protecting apoptotic cells [45]. Together, our results, showing reduction in CaSR immunoreactivity in glaucomatous RGCs soma despite increased protein levels, may indicate targeting of the protein from the rough ER to the plasma membrane to perform its roles in both intercellular communication and cell protection from apoptotic death in response to altered extracellular environment caused by glaucomatous stimuli.

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