Neuroretinal Cell Death in a Murine Model of Closed Globe Injury: Pathological and Functional Characterization

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PURPOSE. Blunt ocular trauma causes severe retinal injury with death of neuroretinal tissue, scarring, and permanent visual loss. The mechanisms of cell death are not known, and there are no therapeutic interventions that improve visual outcome. We aimed to study the extent, distribution, and functional consequences of cell death by developing and characterizing a rat model of retinal injury caused by blunt ocular trauma.

METHODS. The eyes of anesthetized adult rats were injured by either weight drop or low-velocity ballistic trauma and assessed by clinical examination, electoretinography, light microscopy, electron microscopy, and TUNEL. Projectile velocity was measured and standardized.

RESULTS. Weight drop did not cause reproducible retinal injury, and the energy threshold for retinal injury was similar to that for rupture. Low-velocity ballistic trauma to the inferior sclera created a reproducible retinal injury, with central sclopetaria retinae, retinal necrosis, and surrounding commotio retinae with specific photoreceptor cell death and sparing of cells in the other retinal layers. The extent of photoreceptor cell death declined and necrosis progressed to apoptosis with increasing distance from the impact site.

CONCLUSIONS. This is the only murine model of closed globe injury and the only model of retinal trauma with specific photoreceptor cell death. The clinical appearance mirrors that in severe retinal injury after blunt ocular trauma in humans, and the ultrastructural features are consistent with human and animal studies of commotio retinae. After ocular trauma, photoreceptor apoptosis may be prevented and visual outcomes improved by blocking of the cell death pathways. (Invest Ophthalmol Vis Sci. 2012;53:7220-7226) DOI: 10.1167/iovs.12-9887

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for the Use of Animals in Ophthalmic and Vision Research. Female rats weighing 170 to 200 g were purchased from Charles River Laboratories (Margate, UK), kept on a 12-hour light/dark cycle with a daytime luminance of 80 lux, and fed and watered ad libitum (as in all our previous studies of central nervous system injury18). Wistar rats were used for studies of weight drop and Lister hooded rats for studies of ballistic injury, except where otherwise stated. All injuries were induced under general anesthesia with inhaled isoflurane in oxygen. For electron microscopy (EM) and terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling (TUNEL), animals were killed by perfusion with fixative under terminal anesthesia with intraperitoneal ketamine/medetomidine. Electronretinograms (ERG) were recorded under general anesthesia with intraperitoneal ketamine/medetomidine, and animals were killed by intraperitoneal overdosage of pentobarbital for retinal whole mounts.

**Weight Drop Injury**

Weights were dropped from a height of 50 cm onto the lateral sclera. Six groups of rats were injured by weight drop (2–6 eyes of 2–4 rats per group). Weights of 22.6 g (0.111 J) and 31.3 g (0.154 J) with 6 mm flat tips were dropped onto the central cornea, and rats were killed at 2 hours after injury. Weights of 17.6 g (0.086 J) and 25.0 g (0.113 J) with 5 mm flat tips and 22.6 and 31.3 g with 6 mm flat tips were dropped onto the lateral sclera, and animals were killed at 2 days after injury. To increase the energy of the weight drop method, a 7 cm doubled section of cataputch (Match System superpower catapult latex; Muddy Tackle International, Heanor, UK), stretched by 7 cm, was used to propel a 22.6 g/6 mm tip weight downward; animals were killed at 2 days after injury. Additional Lister hooded rats were injured with a 31.3 g weight/6 mm tip plus catapult latex. The catapult latex gave a measured impact energy of 0.6 J for both 22.6 and 31.3 g weights due to inefficiencies in the system. Velocity was calculated for weight drop as gravitational potential energy; the additional energy of the catapult latex was calculated by distance traveled when fired horizontally from a known height.

**Ballistic Injury**

Injury was created by firing a dome-headed (air gun pellet) or spherical (ball bearing) weight down a 5.5 × 300 mm steel barrel using compressed air held in a 500 mL reservoir with pressure monitoring, released by a solenoid-activated valve (response time <5 ms, flow coefficient [Cv] 491 NL/min; SMC Pneumatics, Milton Keynes, UK). Projectiles were plastic ball bearings (0.095 g), plastic 0.22-caliber pellets (0.91 g), and metal 0.22-caliber air gun pellets (0.5 g). Projectile velocity was measured using a PC sound card.19 Briefly, a Dell Studio XPS (Dell Corporation Ltd., Berkshire, UK) laptop was used to record the sound of the compressed air device being fired using MAGIX Music Editor 3 (MAGIX AG, Reno, NV) at a metal plate at known distance from the muzzle. The resultant waveform was then viewed and the time between air first leaving the barrel and the impact with the metal plate recorded at 10.5, 22, 31.5, and 42 cm for 10 iterations at each distance. Mean velocity was calculated at the different time points.

Wistar rats (n = 15 rats, 26 eyes) were used for cadaveric studies to determine the risk of rupture. Female Lister hooded rats were used to study the effects of a 0.095 g projectile delivered at 20 m/s to the sclera: (1) bilaterally for EM studies before killing at 2 hours (lateral scleral impact; n = 4 animals), 2 days (inferior scleral impact; n = 4 animals), and 14 days (inferior scleral impact; n = 4 animals) after injury; (2) unilaterally to inferior sclera for TUNEL and immunohistochemical studies before killing at 2 days (n = 4 animals); and (3) unilaterally to inferior sclera for ERG studies and retinal whole mounts before killing at 2 weeks (n = 8 animals).

**In Vivo Imaging**

Animals were examined immediately after injury, at 2 days, and at 14 days by indirect ophthalmoscopy.

**Electroretinography**

ERG were recorded (HM9ERG; Ocuscience, Kansas City, MO) at 2, 7, and 14 days after ballistic injury and interpreted using ERGView (Ocuscience). Animals were dark adapted overnight and prepared for ERG under dim red light (>650 nm). Scotopic flash ERG were recorded at −2.5, 0, and +0.5 log units with respect to standard flash and photopic flash ERG with background illumination of 30,000 mcd/m² at 0 and +0.5 log units, using DTL fiber (Unimed Electrode Supplies, Farnham, UK) corneal electrodes with pressure-molded Aclar (Agar Scientific, Stansted, UK) contact lenses and needle skin electrodes (Unimed).

**Electron Microscopy**

The primary tissue fixative was 4% gluteraldehyde in 0.1 M phosphate buffer (PB; pH 7.3) as used in previous studies of commotio retinae.15,16 Eyes were removed and the cornea incised before 24 hours postfixation in the primary fixative at room temperature. A section of injured retina was dissected out, washed in 0.1 M PB, postfixed in 1% osmium tetroxide in 0.1 M PB for 45 minutes, washed in PB, dehydrated in ascending concentrations of ethanol, and immersed in propylene oxide before infiltrating in resin (Durcupan; Electron Microscopy Sciences, Hatfield, PA) for 24 hours and polymerizing at 56°C for 24 hours. Semithin sections (1 μm thick) were cut using glass knives on an “ultracut” ultramicrotome (Reichert-Jung, Vienna, Austria). Sections were stained with toluidine blue. Ultrathin gold sections (70–90 nm) were cut with a glass or diamond knife, floated on distilled water, mounted on formvar-coated 50-mesh copper grids, stained with uranyl acetate and lead citrate, and examined on a JEOL 1200 EX transmission electron microscope (Jeol [UK] Ltd., Welwyn Garden City, UK) fitted with a LaB6 filament at an operating voltage of 80 kV.

**TUNEL and Immunohistochemistry**

The tissue fixative was 4% paraformaldehyde (PFA) in phosphate-buffered saline. Eyes were removed, and the cornea was incised before 24 hours postfixation at 4°C. Specimens were cryoprotected in ascending concentrations of sucrose in phosphate-buffered saline (PBS) at 4°C; the anterior segments were removed, and the retinal cup was embedded in OCT and stored at −80°C. Sections 15 μm thick were cut using a cryostat (Bright Instruments, Huntingdon, UK) and adhered onto SuperfrostTM (Fisher, Loughborough, UK)-coated glass microscope slides. TUNEL FragEL DNA Fragmentation Detection Kit (Merck, Nottingham, UK) was used per manufacturer’s instructions, except that proteinase K was omitted and replaced by 15 minutes in Triton X-100 0.1% in PBS. Immunohistochemistry was performed using primary antibodies to ED1 (1/400, monoclonal; ABD Serotec; Kidlington, UK) and OX42 (1/400, monoclonal; ABD Serotec; and 1/100, polyclonal; Santa Cruz Biotechnology, Inc., Heidelberg, Germany) with Texas red anti-mouse and Alexa Fluor 488 anti-rabbit IgG (both 1/400; Invitrogen, Grand Island, NY) secondary antibodies.

**Retinal Whole Mounts and Counting of Photoreceptors**

Anterior segments and vitreous were removed; eye cups were fixed in 4% PFA in PBS, and the neuroretina was dissected from the RPE except when it was adherent, in which case the RPE was dissected from the choroid. Retinai were permeabilized in 0.1% Triton X-100/4',6-diamidino-2-phenylindole (DAPI) in PBS for 2 hours and mounted with DAPI mountant for cell counting. Images were captured at 63x magnification using a confocal laser scanning microscope (Zeiss, Hertfordshire, UK) running the LSM 510 software version 3.2 (Zeiss). Images were captured from three different areas (at 1/6, 3/6, and 5/6 of the radius) of each quadrant (total = 12 counts per eye) in the middle of the outer nuclear layer, to account for variation of photoreceptor density.
numbers in the different areas, and quantified using the built-in particle counting facilities in ImagePro (Media Cybernetics, Bethesda, MD). Mean photoreceptor density per high-power field (0.146 ± 0.146 mm) was expressed as a percentage of the mean value for uninjured control tissue.

Semithin (1 μm) toluidine blue–stained retinal sections were cut 2 weeks after injury from the optic disc to the ciliary body through the center of the impact site; control tissue was taken from the unjured inferior retina of animals killed at 2 hours after impact to the lateral sclera. Images were captured at 40× magnification (0.25 × 0.3 mm field of view) from five different areas of each section and quantified by a blinded observer using the user-defined manual counting facility in ImagePro. For display, counts were normalized as percentages of the mean value for uninjured control tissue in the same retinal area.

**Statistical Analysis**

Cell count and ERG data were normally distributed and analyzed with parametric tests in SPSS 19 (IBM, Armonk, NY). Means ± standard error were calculated for all samples. Electrophysiological data were analyzed using either three-way (time, intensity, injury) repeated measures ANOVA or generalized estimating equations for sets missing data (type III sum of squares). Nonsignificant interactions were removed from the models, and normality was assessed using residual plots.

**RESULTS**

**Weight Drop Did Not Create a Reproducible Retinal Injury**

Two hours after impact to the central cornea, no injury was seen by clinical examination or light or electron microscopy. After direct scleral impact, retinal pathology was clinically occult, as the clinical examination and macroscopic appearances were normal. Photoreceptor outer segment disruption was induced in two of nine eyes to which 0.6 J was delivered; and, at this energy level, globe rupture occurred in two of nine eyes (Fig. 1).

**Ballistic Trauma Created a Reproducible Retinal Injury**

In cadaveric studies, a chamber pressure of ≥0.15 bar caused rupture in ≥50% of eyes. Weights of 0.91, 0.50, 0.16 (air gun pellets), and 0.095 g (ball bearing) were tested; 0.095 g was chosen to give the highest-velocity injury, 20 m/s at chamber pressure of 0.125 bar.

Immediately after injury, there was retinal pallor underlying the injury site in all eyes and variable vitreous hemorrhage. The pallor resolved over the next 2 weeks to leave an area of retinal atrophy revealing the choroidal vasculature.

Light and electron microscopy of resin-embedded retinal sections at 2 hours after injury showed disruption of photoreceptor outer and inner segments (OS) and disrupted external limiting membrane (ELM) in three eyes (Fig. 2).

**Specific Photoreceptor Death Developed after Ballistic Blunt Ocular Trauma**

By 2 weeks after ballistic injury, a large area of retinal and RPE atrophy developed. There was progressive loss of the ONL approaching the center of the impact site but relative preservation of the inner nuclear and ganglion cell layers, with hypopigmentation and irregularity of the underlying RPE (Figs. 3A, 3B). At the center of the impact site, all neuroretinal layers were absent.

The proportion of photoreceptors surviving increased at increasing distances from the center of the impact site, demonstrated by ONL cell counts on toluidine blue-stained resin sections of the retina radially from the optic disc to the ciliary body, through the center of the lesion site (Fig. 3C).
ONL cell counting on retinal whole mounts confirmed that 18.4 ± 4.6% of photoreceptors had died.

Photoreceptors Died by Necrosis and Apoptosis

By 2 hours after injury, there were occasional apoptotic nuclei in the ONL with chromatin condensation and nuclear blebbing (Fig. 4A). Two days after injury, in the center of the impact site, there was infiltration of macrophages (histiocytic, ED1-positive, OX42-negative; Figs. 4B, 4I–K, Supplementary Material and Supplementary Figs. S1C–H, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.12-9887/-/DCSupplemental), edema, and necrotic nuclei in the ONL (Figs. 4B–E). In the perilesional area there was disruption of outer segments with inflammatory cell infiltration. There were fewer necrotic photoreceptor nuclei at increasing distances from the lesion site but more apoptotic photoreceptor nuclei (Figs. 4E–H; see Supplementary Material and Supplementary Figs. S1A, S1B http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.12-9887/-/DCSupplemental). Centrally in the impact site, many cells with diffuse nuclear and cytoplasmic TUNEL staining were observed (Fig. 4L), probably because direct cellular injury caused necrotic death. However, in areas away from the impact site, the presence of specific TUNEL-positive photoreceptor cell nuclei confirmed apoptosis of these cells (Figs. 4L–O).

Electrophysiological Assessment of Ballistic Injury

To assess retinal function, scotopic and photopic flash ERG series were recorded and the major components (a- and b-waves) observed. The a- and b-wave amplitudes were quantified and compared between injured and control eyes at 2, 7, and 14 days postinjury. The magnitude and latency of all components were intensity dependent, though only amplitude was injury dependent. ANOVA was used to compare scotopic (dark adapted) a-wave amplitudes. Scotopic a-wave amplitude was significantly reduced by injury (Fig. 5A; 1 df, $F = 59.7$, $P < 0.001$, $g = 0.895$), and there was no significant change between the three time points (2 df, $F = 2.706$, $P = 0.129$, $g = 0.279$). Since b-wave onset can obscure the a-wave, gradient of the a-wave's linear portion (leading edge) was measured, giving a trend similar to that observed for amplitude (1 df, $F = 40.9$, $P < 0.001$, $g = 0.854$). Scotopic b-wave amplitude was significantly reduced by injury (2 df, $F = 39.9$, $P < 0.001$, $g = 0.851$), though there was no significant effect on the scotopic b-/a-wave ratio (see Supplementary Material and Supplementary Fig. S2, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.12-9887/-/DCSupplemental; 1 df, $F = 4.917$, $P = 0.062$, $g = 0.413$). Photopic a-waves were frequently undetectable in injured and control eyes, and some data points were missing. Generalized estimating equations were therefore used to analyze photopic b-wave amplitudes, which were significantly reduced by injury (Fig. 5B; unstructured correlation matrix, 1 df, $P < 0.001$), consistent with the outer retinal injury affecting both cones and rods equally.

DISCUSSION

This is the first murine model of closed globe injury to show that low-velocity ballistic trauma creates reproducible.commo-tio retinae, with pathological features that mirror the clinical and OCT findings in humans after severe blunt ocular trauma. At the impact site, there was necrosis of photoreceptors, and centrally there was death of all retinal cells...
Away from the impact site, photoreceptor death occurred by a combination of apoptosis and necrosis. Preservation of the inner retinal layers, together with specific photoreceptor cell death, mirrored human OCT findings. The reason this injury predominantly affects photoreceptors is unclear, but we suggest that shearing forces at the neuroretina–RPE junction during globe deformation make them particularly vulnerable.

The spectrum of cell death ranges between apoptosis and necrosis, with significant overlap. We have shown cells that are clearly necrotic (Figs. 4C, 4D) and cells that exhibit features consistent with apoptosis (Figs. 4E–H, 4L–O; see Supplementary Material and Supplementary Figs. S1A, S1B, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.12-9887/-/DCSupplemental), though it is likely that many photoreceptors undergo a mixed form of cell death. While necrosis is an unregulated form of cell death, apoptosis is mediated by specific cell death signaling pathways, as is the mixed form of cell death, termed necroptosis, and these pathways can be modulated to increase cell survival. Thus, our results imply that in a proportion of photoreceptors after ballistic retinal injury, cell death is mediated by apoptotic or necroptotic...
regulated signaling pathways, with a potential for successful neuroprotective therapies.

In human studies, mild commotio retinae associated with photoreceptor recovery is characterized by outer segment disruption, whereas severe commotio retinae is associated with photoreceptor inner and outer segment disruption observed by OCT. Inner segment disruption was also present in our model, but has not been reported in other (mild, recovering) animal models of commotio retinae. The RPE and photoreceptor outer and inner segments appear as three bands on spectral-domain OCT. The thinner line above them may represent the ELM or the myoid portion of the cone inner segments. Loss of this line on OCT may be a marker of severe structural changes.

In conclusion, we report an animal model of the outer retinal changes that occur after severe commotio retinae in humans, which is the first murine rodent model of blunt ocular trauma. Our results suggest that outer retinal atrophy occurs as a result of photoreceptor apoptosis and necrosis due to a terminal injury to photoreceptor inner segments. The observed photoreceptor apoptosis is translationally relevant because there is the potential to prevent neuronal death and improve visual outcomes with use of antiapoptotic neuroprotective therapies. This model provides an opportunity for both mechanistic and translational therapeutic studies.

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References


