RESEARCH ARTICLE

Inner retinal preservation in the photoinducible I307N rhodopsin mutant mouse, a model of autosomal dominant retinitis pigmentosa

Antonia Stefanov$^{1,2}$ | Elena Novelli$^1$ | Enrica Strettoi$^1$

$^1$Institute of Neuroscience, Italian National Research Council – CNR, Pisa, Italy
$^2$Regional Doctoral School of Neuroscience, University of Florence, Florence, Italy

Correspondence
Enrica Strettoi, Italian National Research Council – CNR, Institute of Neuroscience, Pisa, Italy.
Email: enrica.strettoi@in.cnr.it

Abstract
Rod-cone degenerations, for example, retinitis pigmentosa are leading causes of blindness worldwide. Despite slow disease progression in humans, vision loss is inevitable; therefore, development of vision restoration strategies is crucial. Among others, promising approaches include optogenetics and prosthetic implants, which aim to bypass lost photoreceptors (PRs). Naturally, the efficacy of these therapeutic strategies will depend on inner retinal structural and functional preservation. The present study shows that in photoinducible I307N rhodopsin mice (Translational Vision Research Model 4 [Tvrm4]), a 12k lux light exposure eliminates PRs in the central retina in 1 week, but interneurons and their synapses are maintained for as long as 9 weeks postinduction. Despite bipolar cell dendritic retraction and moderate loss of horizontal cells, the survival rate of various cell types is very high. Significant preservation of conventional synapses and gap junctions in the inner plexiform layer is also observed. We found the number of synaptic ribbons to gradually decline and their ultrastructure to become transiently abnormal, although based on our findings intrinsic retinal architecture is maintained despite complete loss of PRs. Unlike common rodent models of PR degeneration, where the disease phenotype often interferes with retinal development, in Tvrm4 mice, the degenerative process can be induced after retinal development is complete. This time course more closely mimics the timing of disease onset in affected patients. Stability of the inner retina found in these mutants 2 months after PR degeneration suggests moderate, stereotyped remodeling in the early stages of the human disease and represents a promising finding for prompt approaches of vision restoration.

KEYWORDS
bipolar cell, dendrites, retinitis pigmentosa, rhodopsin, synapse

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Degeneration of retinal rods and loss of scotopic vision followed by secondary degeneration of cones progressing to severe, blinding visual impairment are the main clinical manifestations of retinitis pigmentosa (RP), a family of inherited diseases with an estimated incidence of 1:4,000 (Hamel, 2006). RP can be caused by mutations in any of up to 65 different genes identified so far (RetNet: https://sph.uth.edu/retnet/sum-dis.htm). The first symptoms typically manifest during young adulthood, after the retina is fully developed. However, numerous experimental studies on RP have been conducted using naturally occurring or transgenic rodent models (i.e., rd1, rd10, P23H rhodopsin mutant mice), in which photoreceptor (PR) death initiates concomitantly with the late phases of retinal differentiation and refinement of the synaptic circuitry, complicating comparisons to the human pathology. Developing retinal neurons may be intrinsically more vulnerable to the effects of PR death or, conversely, show increased adaptation to the consequences of the PR degeneration.

Translational Vision Research Model 4 (Tvrm4) mice, isolated at the Jackson Laboratory in 2010, bear a single nucleotide polymorphism (SNP) in the rhodopsin gene (rho), resulting in an I307N mutation (substitution of isoleucine to asparagine at Position 307) in the seventh transmembrane domain of the rhodopsin protein (RHO) (Budzynski et al., 2010). The mutation does not affect the structure or function of RHO in the retina if mice are maintained in normal, ambient light levels. The mutant RHO is folded normally and transported to rod outer segments. However, if the retina is briefly exposed to a very strong light (i.e., 1,000 lx), the phototransduction cascade leads to a lower binding rate of the mutant opsins to 11-cis-retinal and a higher than normal activation of transducin by unbound opsins (Budzynski et al., 2010). Free opsins are known to have a toxic effect on PRs and light induction causes a massive degeneration of rods, particularly in the central retina where light reaches the highest intensity. Cone death accompanies rod degeneration as a secondary phenomenon (Budzynski et al., 2010) as observed in RP (Léveillard et al., 2004). The observed phenotype of Tvrm4 mutants is similar to human class B1 RP due to mutations of RHO with a dominant inheritance pattern. Patients also exhibit focal PR degeneration which can be triggered or accelerated by exposure to bright light flashes and a moderate disease progression (Cideciyan et al., 1998; Gal, Apfelstedt-Sylla, Janecke, & Zrenner, 1997; Mendes, Van Der Spuy, Chapple, & Cheetham, 2005; Rakoczy et al., 2011).

Previous studies in our laboratory confirmed normal retinal morphology in heterozygous (HT) Tvrm4 mice kept in normal light conditions (animal house standards) and report that 2 days after photoinduction at 12,000 lx for 2 h, a dramatic decrease in the a and b waves of the electroretinogram (ERG) is seen. By ~1 week, PRs are completely lost from the central retina and condensed, DNA-filled vacuoles remain. PR death is accompanied by Müller glial activation, also confined to the central, degenerating retina. Similarly, lba1-positive microglial cells with a highly activated, amoeboid morphology, invade the outermost layers of central retina, whereas resident microglia retain a normal morphology in peripheral areas (Gargini, Novelli, Piano, Biagioni, & Strettoi, 2017). The first sign of morphological remodeling of the inner retina is retraction of dendritic arbors from rod bipolar cells (RBCs), decreased metabotropic glutamate receptor 6 (mGluR6) immunoreactivity in the outer plexiform layer (OPL), and displacement of mGluR6 puncta in the inner nuclear layer (INL). It is reported that the degenerating central area does not spread over time; but becomes progressively smaller 1 month after the initial photoinduction. Consequently, the amplitude of the scotopic ERG recovers partially, likely because PR cells at the mid-periphery of the retina can regenerate damaged outer segments (Gargini et al., 2017).

Outer retinal degeneration is well described in various rodent models of RP, while our knowledge about inner retinal response to PR death, especially in mature animals, is significantly more limited, particularly regarding the maintenance of synaptic connections in the inner plexiform layer (IPL) and functional consequences of synaptic rearrangements (Euler & Schubert, 2015; Greferath et al., 2015; Soto & Kerschensteiner, 2015; Strettoi, 2015). Improvements in restoration of light sensitivity in degenerated retinas (e.g., by prosthetic implants or optogenetic expression of photosensitive molecules) will require a better understanding on how PR loss affects inner retinal neurons and their fundamental circuitry. It is particularly important to document remodeling and fate of neurons of the retinal vertical pathway, as bipolar cells (BCs) are a primary target of vision restoration strategies. BCs are very numerous and play a crucial role in parallel processing of visual signals and proper transfer of information to ganglion cells (GCs). Clearly for this strategy to succeed, the GC cells need to survive as they are the singular output of electrical signals from the retina.

In the present study, we used the I307N Tvrm4 mice to study inner retinal remodeling in an adult model of RP, at early stages following complete PR degeneration.

2 | METHODS

2.1 | Animals and experimental design

Animal experimental protocols were approved by the Italian Ministry of Health (Protocol #14/D-2014, CNR Neuroscience Institute) and by the Ethical Committee of the CNR Institution. Animals were treated in accordance with the Association in Vision and Ophthalmology statement for the use of animals in research. Experiments were conducted on HT Tvrm4 mice (RhoTvrm4/Rho+) carrying the I307N mutation on the rho gene and on C57Bl/6J (WT) littermates, both strains originally from Jackson (Bar Harbor, ME). Genotyping was carried out according to Budzynski et al. (2010). Before photoinduction, both mouse pupils were dilated with one drop/eye atropine if the eyes were used for immunohistochemistry, and WT animals were used as controls. For electron microscopy analysis, the right pupils were dilated, while the left eyes served as internal controls (CTRL). Nonilluminated areas at the retinal periphery were also used as internal controls as their morphology was also unchanged. A total of
30 (15 mutants and 15 WT) adult mice (aged 3–5 months) were used for immunohistochemistry; six additional HT mice were used for electron microscopy. Retinal structure was examined at 3, 6, and 9 weeks after photoinduction. A minimum of n = 3 animals were used per experimental group.

### 2.2 | Photoinduction protocol

Following 4 hr of dark adaptation in a black cage, Tvrm4 mice (aged 3–5 months) received bi- or unilateral (right eye-only) eye drops of 1 μl, 0.5% atropine (Allergan). After 10 min, mice were placed in the photoinduction chamber: a black box (140 × 30 cm² size; 30 cm height) lined with mirrors on the inner sides and covered with a lid holding four fluorescent bulbs (Philips Master TL5 HE 28 W/840, length 115 cm; 16 cm in diameter; Correlated Color Temperature (Nom) 4,000 K Luminous Efficacy (@Max Lumen, Rated) 104 lm/W cool White mercury lamps). Intensity of light was monitored by an inbox light meter sensor (mod. LX103, Lutron, Digital Instruments), at 30 s intervals. Photoinduction duration was controlled by a digital clock; during the procedure animals were fully awake and freely moving. Temperature did not change significantly during light exposure (Gargini et al., 2017). Animals were exposed to 12,000 lx light for 2 min, after which the animals were kept in the dark for one additional minute (“cooldown” time) postinduction (PI), then returned to the local animal facility where they were kept in ambient light conditions (below 60 lx) on a 12/12-hr light/dark cycle, with water and food ad libitum until the time of retinal examination.

### 2.3 | Tissue preparation and immunohistochemistry

Retinal tissue was harvested at 3, 6, or 9 weeks PI. Mice were anesthetized with intraperitoneal Avertin (3-bromo-ethanol in 1% tert-amyl alcohol; 0.1 ml/5 g body weight) and euthanized by quick cervical dislocation following enucleation of eyes, previously labeled on the dorsal pole with a surgical skin marker (Secureline Surgical Skin Marker, Aspen Surgical). Eyecup preparation was performed in 0.1 M phosphate buffer (PB) by removing the anterior segments (cornea, iris, lens) followed by immediate fixation in 4% paraformaldehyde (PFA) in 0.1 M PB, pH 7.4, for 30 min or 1 hr at room temperature. Fixation was followed by 3 × 5 min washes in 0.1 M PB, infiltration in 30% wt/vol sucrose dissolved in 0.1 M PB and infiltration in Tissue-Tek O.C.T. compound (4583, Sakura, Olympus, Italy). Sample freezing in tissue molds was achieved with isopentane in dry ice; samples were stored at −20 °C until use. Frozen eyecups were later processed for either whole mount preparation or cryosectioning.

Frozen, vertical sections were cut at a cryostat at 12 μm thickness and collected on Super Frost slides for immunohistochemistry (IHC). For whole mount preparations, defrosted eyecups were extensively washed in phosphate buffered saline (PBS); the retina was gently detached from the sclera and flattened by making four radial cuts toward the optic nerve head maintaining a reference on the dorsal pole.

For IHC, both retinal whole mounts and sections were incubated in: blocking solution (0.03% Triton-X 100 + 5% serum from the species in which the secondary antibody was generated, in 0.01 M PBS), for 2 hr at room temperature if sections and overnight at 4°C if whole mounts; primary antibody (Ab) solution (diluted in PBS + 0.01% Triton-X 100 + 1% serum) overnight at 4°C if sections and 3 days at 4°C if whole mounts; in fluorescent secondary Ab solution—diluted as the primary Ab—for 2 hr at room temperature if sections and 2 days at 4°C if whole mounts (Barone, Novelli, Piano, Gargini, & Strettoi, 2012). Secondary antibodies were conjugated with Alexa Fluor 488 or Rhodamine Red X (Table 1).

### 2.4 | Antibody characterization

Technical information regarding the antibodies used in this study are summarized in Table 1.

Rhodopsin antibody recognizes an epitope at the N-terminus of the rhodopsin molecule (Hargrave et al., 1986; Hicks & Barnstable, 1987; Silver et al., 1988). In rat retinal protein extract immunoblotting, two closely spaced 39 kDa bands as well as bands at 78 and 115 kDa with lower intensity representing the rhodopsin monomer and its aggregates are marked (Fekete & Barnstable, 1983). The antibody specifically labels rod PR cells (but not cones) in the rat retina via immunohistochemistry (Barnstable, 1980). In the mouse retina the product labels rod outer segments exclusively (Roof, Adamian, & Hayes, 1994) and the staining observed in our study was consistent with these reports.

Protein kinase Ca (PKCa) antibody recognizes an 80 kDa polypeptide of PKC in bovine brain lysates (manufacturer’s data sheet). PKC is a well-established marker of RBCs in the rat (Grefeath, Grünert, & Wässle, 1990) and mouse retina (Barone et al., 2012; Havercamp, Ghosh, Hirano, & Wässle, 2003) and the same staining consistency was observed in our study.

CD68 is homologous to the mouse antigen macrosialin. Apart from nonhematopoietic tissues like liver and kidney, where CD68 expression is the highest, the antigen can be found on the outer surface of macrophages, monocytes, neutrophils, basophils, and large lymphocytes. The CD68 (KP1) mouse monoclonal antibody was raised against human alveolar macrophages and it stains a 101 kDa and a 70 kDa band on immunoblots of whole cell lysates (manufacturer’s data sheet). In the mouse retina CD68 is a marker of microglia and it is often used in conjunction with Iba-1 (Santos et al., 2008), resulting in a similar staining pattern to the one reported here.

C-terminal binding proteins (CtBP1 and CtBP2) are a class of corepressors capable of interacting with certain transcription factors via a PLDLSL sequence motif. Immunoblotting of mouse brain and retinal tissue lysates revealed that the B-domain of RIBEYE also labels the 48–50 kDa CtBP2 band besides being present both in PR and BC ribbon synapses in the mouse retina based on IHC (Dieck et al., 2005). Our observations confirm similar immunoreactivity.
Glutamine synthetase (GS) is an enzyme responsible for catalyzing the amination of glutamic acid and plays a pivotal role in glutamate neurotransmitter recycling. GS immunoblotting of sheep or rat brain lysates shows a strongly labeled 45 kDa band (manufacturer’s data sheet), reflecting the fact that mammalian GS is composed of eight 45 kDa subunits. Müller glia is reported to show intense GS immunoreactivity (Riepe & Norenburg, 1977) identical to what observed here.

SMI32 reacts with a nonphosphorylated epitope in neurofilament H of most mammals. In western Blots, SMI32 antibodies show two bands at 180 and 200 kDa, which tend to merge into a single band on 2D blots (manufacturer’s data sheet). In the retina of most mammalian species, SMI32 antibodies label the soma and axons of GCs (Lewis & Nixon, 1988).

Glutamic acid decarboxylase (GAD) catalyzes the synthesis of γ-aminobutyric acid (GABA), a major inhibitory neurotransmitter. GAD is present in the mammalian brain in two forms: the 65 kDa GAD65 and the 67 kDa GAD67, respectively, more abundant in axon terminals and generally distributed throughout the neuron (Kaufman, Houser, & Tobin, 1991). In the mouse retina, GAD67 labels GABAergic amacrine cells (ACs) in the INL and IPL with intense immunofluorescence (Müller, Azar, de los Santos, & Brecha, 2017). Patterns of staining found here coincide with previous reports.

Cone Arrestin or Arrestin-C is encoded by the Arr3 gene in murine species. Arrestin-C is predominantly found in inner and outer segments of cone PRs and the IPL of the retina (Gurevich & Benovic, 1995). Cone Arrestin antibody evaluated by the manufacturer by immunoblotting of mouse retinal tissue lysate labeled a 50 kDa band. Immunohistochemical characterization of Cone Arrestin expression pattern performed by Zhu et al. (2002a) and (2002b) reported similar staining intensities and pattern observed in the present study.

Calbindin D is a member of the intracellular calcium-binding proteins family characterized by EF-hand motifs. Calbindin D antibodies recognize a single band of approximately 27 kDa in immunoblots (Airaksinen et al., 1997). In the mouse and rat retina, Calbindin D antibody labels horizontal cells and a subpopulation of ACs (Hamano et al., 1990). Staining pattern shown in this study confirms previous reports.

### Table 1

| Primary and secondary antibodies used in this study |
|-----------------------------------------------------|
| Mouse monoclonal primary antibodies | Antibody manufacturer, catalog #, RRID | Dilution rate |
|--------------------------------------|----------------------------------------|---------------|
| Rhodopsin                            | Sigma-Aldrich, Italy; O4886             | 1:1000        |
| Protein kinase Ca (PKCa)             | Sigma-Aldrich; P5704 RRID:AB_477375    | 1:800         |
| CD68 (KP1)                           | Santa Cruz Biotechnology; sc-20060 RRID:AB_627158 | 1:250         |
| C-terminal binding protein-2 (CtBP2/RIBEYE) | BD biosciences; 612044            | 1:800         |
| Glutamine synthetase (GS)            | Merck-Millipore, Italy; MAB302         | 1:1000        |
| Nonphosphorylated neurofilaments (SMI32) | Covance, CA; SMI-32R RRID:AB_10719742 | 1:800         |
| Glutamic acid decarboxylase-67 (GAD67) | Merck-Millipore; MAB5406               | 1:500         |

| Rabbit polyclonal primary antibodies |
|--------------------------------------|
| Cone Arrestin                        | Merck-Millipore: AB15282              | 1:5000        |
| Calbindin D                          | Swant Ltd., Switzerland; CB38a        | 1:500         |
| Connexin36 (Cx36)                    | Invitrogen, Carlsbad, CA; 51-6200     | 1:500         |
| Protein kinase Ca (PKCa)             | Sigma-Aldrich; P4334                  | 1:800         |
| Glial fibrillary acidic protein (GFAP) | Sigma-Aldrich; G9269 RRID:AB_477035  | 1:1000        |
| Ionized calcium binding adapter molecule 1 (IBA1) | Wako Chemicals GmbH, Germany; 019-19741 | 1:500         |
| Disabled-1 (DAB1)                    | Kind gift of Prof. Botond Roska, Basel, Switzerland | 1:200         |
| RBPMS                                | PhosphoSolutions, Aurora, CO; 1830-RBPMS RRID:AB_2492225 | 1:500         |
| Tyrosine hydroxylase (TH)            | Merck-Millipore: AB152                | 1:1000        |

| Sheep polyclonal primary antibody    |
|--------------------------------------|
| Secretagogin (SCGN)                  | BioVendor GmbH, Germany; RD184120100  | 1:2000        |

| Goat polyclonal primary antibody     |
|--------------------------------------|
| Choline acetyltransferase (ChAT)     | Merck-Millipore; AB144P RRID:AB_11212843 | 1:500         |

| Secondary antibodies                 |
|--------------------------------------|
| Donkey anti-mouse Alexa Fluor 488    | 715-545-151 RRID:AB_2341099            | 1:800         |
| Donkey anti-rabbit Alexa Fluor 488   | 711-546-152                            | 1:800         |
| Donkey anti-goat Alexa Fluor 488     | 705-546-147 RRID:AB_2340430            | 1:800         |
| Donkey anti-sheep Alexa Fluor 488    | 713-546-147                            | 1:800         |
| Donkey anti-mouse Rhodamine Red X    | 715-295-151                            | 1:800         |
| Donkey anti-rabbit Rhodamine Red X   | 711-295-152                            | 1:800         |

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Calbindin D is a member of the intracellular calcium-binding proteins family characterized by EF-hand motifs. Calbindin D antibodies recognize a single band of approximately 27–28 kDa in immunoblots (Airaksinen et al., 1997). In the mouse and rat retina, Calbindin D antibody labels horizontal cells and a subpopulation of ACs (Hamano et al., 1990). Staining pattern shown in this study confirms previous reports.
Connexin 36 (Cx36), a prime component of neuronal gap junction connexins is known to play a pivotal role in the rod pathway; it was first identified in the rat inferior olive and confirmed to be strongly expressed in neural tissue, including the retina (Condorelli et al., 1998; Demb & Pugh, 2002; Söl, Degen, Teubner, & Willecke, 1998). Western blot detects monomeric Cx36 migrating at 36 kDa in homogenates of retina, olfactory bulb, and inferior olive (Rash et al., 2000). In the mouse retina Cx36 immunolabeling is concentrated in the IPL. Staining patterns found in this study match previous reports (Ghinia, Novelli, Saigo, Badea, & Strettoi, 2019).

The glial fibrillary acidic protein (GFAP) is a fundamental protein component of intermediate filaments of CNS macroglia (Eng, 1985). The GFAP positive band on immunoblots of mouse, rat, and human brain lysates appears at approximately 50 kDa (manufacturer’s data sheet). In the healthy mouse retina, GFAP is a reliable marker of astrocytes, while in the diseased tissue, it is widely used as a marker of Müller glial hypertrophy (Eisenfeld, Bunr-milam, & Sarrhy, 1984). Our observations confirm similar immunoreactivity.

Iba1 is a calcium-binding protein with a molecular weight of 17 kDa specifically expressed in macrophage/microglia (manufacturer’s data sheet). A widely used microglial marker in mouse retina (Gargini et al., 2017; Guadagni et al., 2019) produces a staining pattern similar to what observed in our results.

Mouse disabled-1 (Dab-1) cytoplasmic adaptor protein is the product of the disabled-1 gene and is expressed throughout the CNS during embryonic development. Cell lysates of differentiated, P19 brain cells were found to express 60, 80, and 120 kDa mouse Dab-1 isoforms by immunoblot analysis (Howell, Gertler, & Cooper, 1997). In the mature mouse retina, Dab-1 was confirmed to be expressed in all ACs by IHC (Rice & Curran, 2000) with staining patterns similar to what we report here.

RBPMS or RBPMS protein with multiple splicing also known as HERMES belongs to the family of RNA binding proteins possibly playing a role in transcription and translation through binding to RNA. RBPMS antibody recognizes a 24 kDa protein band on immunoblots (manufacturer’s data sheet). RBPMS is specifically expressed in retinal GCs (RGCs) in multiple mammalian species (Kwong, Caprioli, & Piri, 2010; Rodriguez, de Sevilla Müller, & Brecha, 2014). Patterns of staining found here coincide with previous reports.

Tyrosine hydroxylase (TH) is the enzyme responsible for the biosynthesis of the catecholamines dopamine and norepinephrine from tyrosine and is a useful marker for dopaminergic and noradrenergic neurons. By western blot, the TH antibody selectively labels a single band of about 62 kDa (reduced), corresponding to the protein itself (manufacturer’s data sheet). Anti-TH is a robust marker for dopaminergic ACs (DAs) of multiple mammalian species including mice (Wulke & Schnitzer, 1989). We report a similar staining pattern here.

Secretagogin (SCGN) belongs to the EF-hand (E-helix-loop–F-helix-hand) superfamily of calcium-binding proteins. The sheep anti-SCGN antibody recognizes a 32 kDa band on western blots of mouse cerebellum and retina extracts. It is a marker for Type 2, 3, 4, 5, 6, and possibly 8 cone bipolar cells (CBCs) (Puthussery, Gayet-Primo, & Taylor, 2010). We observed staining results similar to what was found in previous publications.

The choline acetyltransferase (ChAT) enzyme is responsible for the synthesis of acetylcholine (Ach). The goat-anti-ChAT antibody recognizes a 70/74 kDa protein band on immunoblots (manufacturer’s data sheet). In the mouse retina, ChAT antibodies specifically label cholinergic or starburst ACs (Jeon, Strettoi, & Masland, 1998). Our observations confirm a similar staining pattern.

### 2.5 Transmission electron microscopy

To achieve optimal preservation of mouse retina, we implemented a protocol combining methods from several laboratories (Ghinia et al., 2019; Morrow, Furukawa, Raviola, & Cepko, 2005).

Animals were deeply anesthetized with intraperitoneal (0.1 ml/5 g body weight) Avertin injections, eyes quickly explanted, and mice euthanized by cervical dislocation. Enucleated eyes—with a blue mark on the dorsal pole—were immediately transferred in oxygenated artificial cerebrospinal fluid where the cornea, iris, and lens were removed, and the eyecups were immersed in primary fixative (1% vol/vol glutaraldehyde +2% vol/vol PFA in Sorenson’s buffer M/30, pH 7.4 + 5% wt/vol sucrose) for 2 hr at 4°C. Following primary fixation, the eyecups were cut into four equal pieces corresponding to the retinal quadrants (dorsal, nasal, ventral, and temporal) without detaching the retina from the sclera and transferred into secondary fixative (3% vol/vol glutaraldehyde in Sorenson’s buffer M/30, pH 7.4 + 5% wt/vol sucrose) overnight at 4°C. Afterward, samples were extensively washed in buffer at room temperature (Sorenson’s buffer M/30, pH 7.4 + 5% wt/vol sucrose–10 × 5 min) and immersed in 3% wt/vol potassium ferrocyanide +2% (wt/vol) OsO4 in H2O, at 4°C for 2 hr. Osmication was followed by washing steps (0.05 M maleate buffer, pH 5.15–10 × 5 min) and by staining en bloc with 1% (wt/vol) uranyl acetate (in 0.05 M maleate buffer, pH 6) for 1 hr at 4°C. Blocks were then washed at room temperature (0.05 M maleate buffer, pH 5.15–5 × 5 min), dehydrated with a graded series of ethanol and infiltrated first with propylene oxide (2 × 15 min), then with a mixture of propylene oxide and epon–araldite resin 50% (vol/vol) catalyzed with 2% DMP30 overnight at room temperature. Final epon–araldite embedding was performed in flat molds at 65°C. Sectioning was performed using a Leica Ultracut ultramicrotome and a diamond Knife. Ultrathin sections, 90 nm thick, were collected on single-hole copper grids (Formvar Support Film Slots, 2 × 1 mm2 Cu grids; FF2010-CU; Electron Microscopy Science, Inc.). No grid staining was necessary given the high contrast of the specimens and the sensitivity of the EM camera.

### 2.6 Imaging and analysis

#### 2.6.1 Cell counts in retinal whole mounts

Retinal whole mounts were imaged with a Zeiss Axioplan light microscope equipped with a Zeiss Axiocam color camera and a Plan Neofluvar 1.25x/0.035 objective to measure retinal surface prior and post confocal microscopy. Image acquisition for cell number quantification was performed with a Leica TCS-SL confocal microscope.
equipped with 488 and 453 lasers, using a Plan Apochromat 40x/1.40 oil objective, employing identical laser and imaging settings for all samples. Four retinal fields (250 × 250 μm² each) were acquired at each eccentricity of the tissue (dorsal, nasal, ventral, and temporal), covering ~1/15 of the whole area of the retina and spanning the entire INL thickness on the z axis. Confocal stacks were saved as TIFF files and transferred to a MetaMorph image analyzer, where cell counts were performed. Cells were identified on single focal planes, labeled and followed across the entire z stacks. Average numbers of cells/mm² were multiplied for the corresponding retinal surface, taking a value averaged between the surface before and after confocal imaging. Statistical comparisons of experimental groups were executed in SigmaPlot applying one-way analysis of variance (ANOVA) test with the Holm–Sidak post hoc method.

2.6.2 | Image thresholding analysis

To estimate the area occupied by axonal endings of BCs in retinal whole mounts and vertical sections, or the area covered by immunolabeled synaptic contacts in the IPL, images were obtained with a Leica TCS-SL confocal microscope using identical laser and imaging settings for all samples, using Plan Apochromat 40x/1.40 or 100x/1.30 oil objectives. High definition projection images were acquired (typically spanning 4 μm across the z axis); brightness and contrast were adjusted with the Zeiss software ZENPRO 2012 or with Adobe Photoshop using consistent settings. An identical region of interest (ROI) was applied to each image for quantification using an image thresholding analysis tool of MetaMorph software. First, each ROI was processed by subtracting the background; then, pixels exceeding the threshold were measured automatically and corresponding areas normalized to mm². Data from different ROI were averaged for each experimental group. Statistical evaluation was done in SigmaPlot using one-way ANOVA tests with the Holm–Sidak post hoc method.

2.6.3 | Whole mount reconstructions

Retinal whole mounts were imaged with an Imager.Z2 Zeiss microscope (Zeiss, Milan, Italy) using EC Plan-Neofluar 5x/0.16 M27, 10x/0.3 M27, and 20x/0.50 M27 objectives; images were tiled with ZEN module “Tiles & Positions” software to reconstruct the entire retinal surface. TH positive DACs were counted on tiled images of entire retinal surface obtained with a 10x objective.

Additional images of vertical sections were obtained with the same Zeiss Imager.Z2 microscope equipped with an Apotome2 device (Zeiss) using an EC Plan-Neofluar 20x/0.50 M27 objective.

2.6.4 | Transmission electron microscopy analysis

Images were obtained using a JEOL 1200EX II electron microscope (JEOL, Milan, Italy) equipped with a charge-coupled device Olympus Veleta Megaview camera (Olympus, Münster, Germany) at a magnification of 40,000×, covering systematically the entire width of the IPL represented in each section. A total of 400 transmission electron microscopy (TEM) micrographs (12.96 μm²/micrograph) per retinal sample (corresponding to a sampled area of 0.0052 mm²/sample) were imaged. Image files were saved as TIFF and transferred to a MetaMorph image analysis workstation, where synapses were counted manually. Criteria applied to identify conventional synapses were the presence of synaptic vesicle clusters abutting the presynaptic membrane, the occurrence of synaptic densities and the unequivocal presence of pre- and postsynaptic processes. For ribbon synapses, identification was based on the presence of an osmiophilic ribbon, either bisecting the presynaptic wedge or freely floating in the cytoplasm but unequivocally tethering synaptic vesicles; the presence of an arciform density at the active zone; and the obvious presence of presynaptic and postsynaptic processes. Counts were averaged per experimental group and statistically analyzed in SigmaPlot applying one-way ANOVA tests with Holm–Sidak post hoc method.

3 | RESULTS

3.1 | PRs of the Tvrm4 mouse retina 3 and 6 weeks PI

By 3 weeks PI by light, PR death in the central retina is largely complete, and no rows of nuclei are visible in the outer nuclear layer (ONL). Surviving, intact rods, and cones can only be found at the far retinal periphery. A similar phenotype is also observed at 6 weeks PI (Figure 1). Large bodies that stain intensely with DNA-binding molecules are visible in the outer retina. These products of condensation of dying PRs persist in the subretinal space after PR death and remain visible at 9 weeks PI, the latest time point examined. Observations on remodeling were conducted in PR-free, central retinal areas. The retina of wild type, age matched mice, littermates of Tvrm4 animals, born by crossing HT Tvrm4 mice, were used as controls, as specified in the text.

3.2 | Rod BCs

Typical atrophy and retraction of dendritic arbors (described by Gargini et al. (2017)) is first observed in RBCs at 7 days PI. By 3 weeks, RBCs in the central, most degenerated part of the retina, are completely devoid of dendrites, their progressive shortening and retraction being accompanied by concurrent shrinkage of RBC axonal endings in the IPL. These changes are still seen at 6 weeks PI (Figure 1). In addition to examining the morphological changes, we determined the number of surviving RBCs, by counting cell bodies in retinal whole mounts. The overall RBC number did not show any change during this time period (one-way ANOVA, p = .295; n = 3 retinas/experimental group; 16 images/retina: 4/retinal eccentricity) demonstrating a survival rate of nearly 100% (Figure 2) despite regression of dendritic trees (Figures 1 and 2). A significant shrinkage of axonal endings was found at the same time points and quantified in retinal whole mounts by analyzing the surface covered by
RBC axonal arborizations in sublamina 5 of the IPL. A decrement larger than 50% was observed in the surface occupied by RBC axonal arbors and lying within the PR devoid area (one-way ANOVA, \( p \geq .0001, n = 3 \) retinas) compared to the peripheral retina where PRs are preserved (Figure 2). This confirms that the regression of axonal arbors is secondary to the degeneration of PRs.

3.3 Horizontal cells

Our analysis of remodeling of second-order neurons also consisted of examining HCs in retinal whole mounts stained with anti-Calbindin D. Analyses of retinas at 6 weeks PI demonstrated that HCs completely disappeared from islands of variable sizes and shapes in the central retina, creating transition zones with extremely sharp edges between regions of surviving and dead cells. Interestingly, these islands were not precisely overlapping with PR-lacking areas and HCs were typically missing in the transition zones of PR degeneration (Figure 3a–g); conversely, numerous HCs with hypertrophic processes were preserved in the central retina next to these islands albeit no PRs were present in the corresponding ONL. HC body counts revealed a 37% decrease in cell density (cells/mm²; \( t \) test, \( p \geq .01 \); \( n = 3 \) retinas/experimental group; 16 images/retina: 4/retinal eccentricity) (Figure 3h), over the central retinal area and in the total number (Figure 3i) of HCs relative to WT controls at 6 weeks PI. Calbindin D and Neurofilament-200 (NF200) antibody staining of vertical sections confirmed the observation of retained HC bodies and dendrites in central retinal areas devoid of PRs and of decreased HC density in transition zones. Morphological abnormalities (mostly hypertrophy) were first observed in the axonal arbors of HCs (NF200 positive) postsynaptic to rods, while dendrites and cell bodies were preserved for longer times (Figure 3e–g), thus confirming the typical rod-to-cone degeneration pattern of this mutant. There is no obvious explanation for the higher severity of HC remodeling in transition zones compared to what is observed in the central retina.
3.4 Cone BCs

CBCs were studied on vertical sections stained for anti-secretagogin (SCGN), known to label as many as six different types of CBCs (Types 2, 3, 4, 5, 6, and possibly 8) stratifying at various depths in the outer and innermost tiers of the IPL. Similar to other cell types, quantification was done at 3 and 6 weeks PI. In the central retinal areas, lacking PRs, CBC dendritic trees appeared to have lost continuity and complexity, although retraction was not as severe as observed in RBCs and became very evident only at 9 weeks PI (Figure 4a–c). No statistical difference was revealed in the number of cell bodies in any of the studied experimental groups, indicating a survival rate of 100% of CBCs (Figure 4d).

However, we found a gradual, significant decrease in the thickness of the three layers revealed by SCGN antibodies in the IPL and corresponding to the axonal arbors of the labeled CBCs (one-way ANOVA, $p \geq .001; n = 3$ retinas). Reduced SCGN staining in sublamina b of the IPL (i.e., SCGN+ Layer 3) increased over time (Figure 4e), despite preservation of the thickness of the whole plexiform layer.

3.5 Amacrine cells

Several antibodies were used on vertical retinal sections to label cholinergic (ChAT), GABAergic (GAD67), dopaminergic (TH), and all ACs...
DAB1, and Calbindin positive ACs. IHCH confirmed in most cases the preservation of the aforementioned cell types, which comprise large size amacrines (i.e., cholinergic cells) as well as small field, glycinergic amacrines, such as the AII. No clear morphological changes were evident in the degenerated retina at 3 and 6 weeks PI (Figure 5). Laminar organization of AC processes also appeared to be...
normal, as highlighted by the correct positioning of the cholinergic bands at one-third and two-third of depth in the IPL and the regular distribution of the three Calbindin positive bands. A noticeable exception was observed in the TH+ DACs, which showed hypertrophic processes in the OPL, exclusively in the central retina and precisely overlapping with the area devoid of PRs. Observations conducted 3 and 6 weeks PI demonstrate that the TH+ hypertrophic processes target and densely entangle some but not all the DNA condensed bodies that accumulate in the outer retina during PR degeneration (Figure 6a–j). Cell body counting reveals no statistical difference in control and Tvrml cases (d). IPL thickness measurements were done on the same micrographs with a MetaMorph longitudinal measuring tool (10 random measurements/micrograph done along the full IPL thickness in the vertical plane (** p < .001; data are shown as average and SE; one-way analysis of variance [ANOVA]) [Color figure can be viewed at wileyonlinelibrary.com]

**Figure 4** Vertical sections from retinas of WT controls (a) and Tvrml mice, 3 (b) and 6 (c) weeks postinduction (PI), with secretagogin (SCGN) staining of cone bipolar cells (CBCs). Three distinct plexa (1, 2, 3) of CBC axonal arbors are visible in the IPL. A gradual, significant decrease in the thickness of SCGN+ axonal arborizations becomes evident over time, mostly due to plexus 3 shrinkage, despite preservation of total IPL width. (d,e) CBC quantitative data. Three vertical sections/animal were analyzed taking 10 micrographs/section, covering five fields, each 150 x 150 μm². Cell body counting reveals no statistical difference in control and Tvrml cases (d). IPL thickness measurements were done on the same micrographs with a MetaMorph longitudinal measuring tool (10 random measurements/micrograph done along the full IPL thickness in the vertical plane (** p < .001; data are shown as average and SE; one-way analysis of variance [ANOVA]) [Color figure can be viewed at wileyonlinelibrary.com]

### 3.6 | Ribbon synapses and gap junctions in the IPL

To examine the distribution of connectivity markers in the IPL of Tvrml mice, we used RIBEYE/CtBP2 staining for ribbon contacts and examined vertical retinal sections at 3, 6, and 9 weeks PI (n = 3 for each group). Confocal micrographs were acquired and edited with identical settings. Immunoreactivity in the IPL was quantified to estimate the area covered by RIBEYE+ puncta using an image thresholding analysis in an ROI spanning the whole IPL width. A significantly increased area (one-way ANOVA, p = .010) occupied by RIBEYE+ puncta was revealed 3, but not 6 and 9 weeks after photoinduction with respect to WT controls (Figure 7a–e). A similar method was used to estimate the density of Cx36+ puncta, revealing gap junctions expressing this particular isoform of connexin. Thresholding analysis showed significantly larger areas attributable to Cx36+ signals in the IPL (one-way ANOVA, p ≥ .001) 3, 6, and 9 weeks after photoinduction with respect to WT controls (Figure 7f–j).

### 3.7 | Ultrastructural analysis of synaptic contacts in the IPL

To investigate synaptic architecture in the IPL of areas of complete PR degeneration, we performed TEM studies to estimate the density of conventional and ribbon synapses across the whole width of the IPL in retinal samples obtained at 3 and 6 weeks PI (n = 3 for each group) (Figure 8). To limit internal variability, data from each light-induced retina were compared those obtained from the contralateral
Vertical sections from noninduced control (left column) (a,d,g,j), 3 weeks (central column) (b,e,h,k) and 6 weeks (third column) (c,f,i,l) Tvrm4 retinas. Antibody stainings highlight amacrine cell morphologies and organization: (a–c) Calbindin (red); (d–f) ChAT (green); (g–i) DAB1; and (j–l) GAD67. Blue: Hoechst nuclear counterstaining. The pattern of immunoreactivity in the inner retina is similar to the control at both time points. No major changes are observed in the general morphology and laminar organization of stained bodies and processes [Color figure can be viewed at wileyonlinelibrary.com]
FIGURE 6  Vertical retinal sections of Tvrm4 mice stained with anti-TH (red) and Hoechst (blue) in a non-induced control eye (left column) (a) and 3 weeks (b) or 6 weeks (c) following light exposure. (d–j) Whole mount micrographs of Tvrm4 retinas as above. The focal plane of (d–f) is in the IPL; (g–j) are taken at the ONL/OPL border. (g,h) The inner limit of the ONL and the outer limit of the OPL, respectively, with arrows pointing at TH-positive processes in the outer retina of a control mouse. In induced retinas (e,f,i,j), the inner plexus (e,f) originating from DACs shows a normal morphology while processes in the OPL are hypertrophic and denser (i,j). Exuberant processes are observed entangling DNA bodies residual from dead PRs. (k) Counts of DAC+ cell bodies (3 retinas/exp. group; counts performed on tiled images of retinal whole mounts) revealed no significant changes 3 and 6 weeks PI. Tvrm4 data are compared to the published number of DACs in WT B6/J (CTRL) mice (Whitney, Raven, Ciobanu, Williams, & Reese, 2009) (data are shown as average and SE; one-way analysis of variance [ANOVA]) [Color figure can be viewed at wileyonlinelibrary.com]
(noninduced) retina of the same animals. This was previously shown to exhibit a normal phenotype, identical to that of WT mice (Gargini et al., 2017). Ultrastructural analysis demonstrated general preservation of number and appearance of conventional synaptic contacts in the IPL at both 3 and 6 weeks PI compared to noninduced controls.

RBC axonal endings could be unequivocally identified by TEM based on their large size, position in sublamina 5 of the IPL and typical cytological features (pale cytoplasm, presence of large number of vesicles, mitochondria, and synaptic ribbons). TEM studies confirmed that RBC axonal endings were visibly smaller with respect to noninduced CTRL specimens.

For ribbon synapses, a decrement in the overall number was observed beginning at 3 weeks PI, although the decrement only became statistically significant at 6 weeks PI. Specifically, we found 9% less ribbons 3 weeks PI and 16% less ribbons 6 weeks PI compared to noninduced CTRLs (Figure 8j) (one-way ANOVA, p = .034). More noticeable than the change in number, we found in photoinduced specimens the majority of ribbons displayed a globular/spherical shape, very different from the elongated appearance of normal ribbons and more reminiscent of the immature morphology observed during retinal development (Regus-Leidig, Dieck, Specht, Meyer, & Brandstätter, 2009) (see Figure 8). Quantitative analysis demonstrated that ~72% of all ribbons were globular by 3 weeks PI and 61% of them were similarly rounded at 6 weeks PI, compared to 10% in noninduced controls. In addition, the number of ribbons not anchored to the presynaptic membrane was found to be significantly higher than normal at 3 and 6 weeks PI, while the number of anchored ribbons obviously showed the opposite tendency. Despite

**FIGURE 7** Anti-CtBP2/RIBEYE staining of ribbon synapses in vertical sections of (a) WT retinas and in Tvr-m4 specimens obtained 3 (b), 6 (c), and 9 (d) weeks postinduction (PI), respectively. (e) Quantitative analysis of CtBP2/RIBEYE puncta. A total of three vertical sections/animal and of 10 micrographs/section were acquired. Imaged fields (78 × 78 μm² acquired with a 100x oil objective) covered both the center (five fields) and the periphery (five fields) of the vertical retinal sections. A significant increase in the density of RIBEYE+ puncta in the IPL was revealed measuring the thresholded area (ThA) 3 weeks PI (**) p ≤ .001; data are shown as average and SE; one-way analysis of variance [ANOVA]). (f–i) A similar analysis of IPL gap-junctions labeled with anti-Cx36 (f–i) shows a significant increase in puncta density 3 (g), 6 (h), and 9 weeks (i) PI compared to (f) WT controls (** p ≤ .001; data are shown as average and SE; one-way ANOVA) [Color figure can be viewed at wileyonlinelibrary.com]
these ultrastructural changes, typical dyad synapses established by RBC axonal endings onto two dendritic processes known to represent the All/A17 AC combination were recognizable and well preserved in all experimental groups (Figure 8).

3.8 | Ganglion cells

Previous studies on GCs in retinal degeneration rodent models predict minimal morphological alterations in RGCs at early stages of the
retinal disease progression (Damiani, Novelli, Mazzoni, & Strettoi, 2012; Mazzoni, Novelli, & Strettoi, 2008). Indeed, we did not observe any change in the fundamental anatomical features of GCs. Specific staining of cell bodies with RBPMS antibodies showed normal density of somas and appropriate confinement in the homonymous layer. Dendrites and axonal bundles of GCs, labeled by SMI32 (neurofilament) antibodies, also revealing the somas of large-size GCs, showed absolutely normal distribution in retinal preparations obtained 3 and 6 weeks PI (Figure 9), with no obvious differences with control specimens.

Altogether, the inner retina of Tvrm4 mice, in which PR degeneration was triggered during adulthood, demonstrated significant cell survival, morphological integrity and preservation of synapses and gap junctions after PR degeneration. Nearly 100% survival rate was found among BCs, in spite of some decrement in HC number. The number of conventional synapses in the IPL was unaffected and the area covered by Cx36-positive gap junctions showed an increment, albeit we detected a tendency in the total number of ribbon contacts to decrease over time. Quantitative data are summarized in Table 2.

FIGURE 8  Semithin vertical sections of Translational Vision Research Model 4 (Tvrm4) mouse retina stained with toluidine blue in a noninduced, control eye (a) and 3 (b) and 6 weeks (c) following light exposure. (d–i) TEM micrographs illustrating synaptic ribbons (arrowheads) and conventional chemical synapses (arrows) involving rod bipolar axonal endings (RBE) in a noninduced control retina (d,g) and in Tvrm4 preparations examined 3 (e,h) and 6 (f,i) weeks postinduction (PI). Arrows indicate the direction of neurotransmission. Atypical globular ribbons (d–f, arrowheads) and classical synaptic arrangement involving RBE with elongated ribbons and dendrites of All and A17-like amacrine cells (IAC) can be observed in all preparations (e–i). A summary of conventional and ribbon synapse density in the IPL is shown in (j). TEM micrographs (400/sample, 12.96 μm²/micrograph)/sample, corresponding to a sampled area of 0.0052 mm²/retina) were imaged. The number of conventional synapses (Conv. s.) is not affected by photoinduction. The total number of ribbons shows a significant decrement 6 weeks PI (** = p ≤ .01; one-way analysis of variance [ANOVA]). Also, significantly more ribbons have a globular shape (Glob. r.) (** = p ≤ .001; one-way ANOVA), 3 and 6 weeks PI, compared to non-induced controls (CTRL). Globular ribbons represent 72% of all ribbons 3 weeks PI but decrease to 61% 6 weeks PI. The number of nonanchored ribbons (NA. r.) is significantly higher 3 and 6 weeks PI, while the number of anchored ribbons (A.r.) shows the opposite tendency. All data are shown as average and SE

FIGURE 9  Retinal vertical sections of Tvrm4 mice stained green for anti-RBPMS (a–c) and anti-SMI32 (d–f) along with Hoechst nuclear marker (blue). (a,d) Noninduced retinal periphery, with intact ONL. Tvrm4 samples of central retina shown 3 (b,e) and 6 (c,f) weeks postinduction (PI). Cell bodies of ganglion cells, brightly labeled by anti-RBPMS antibodies, form a continuous layer in both illuminated and control areas (a–c). Dendrites, large-size cell bodies (arrows) and fascicles of optic nerve fibers revealed by anti-SMI32 staining show normal morphological features in all experimental groups [Color figure can be viewed at wileyonlinelibrary.com]
TABLE 2  Summary of quantitative data collected at 3 and 6 weeks PI and shown as averages ± SE

|                         | HT3w                  | HT6w                  | WT     |
|-------------------------|-----------------------|-----------------------|--------|
| RBCs (avg num/retina)   | 223,383 ± 12,290      | 221,659 ± 12,000      | 246,416 ± 31,593 |
| SCGN + CBCs (avg num/retina) | 110,322 ± 4,538      | 106,822 ± 6,358       | 109,478 ± 5,607 |
| HCs (avg num/retina)    | ND                    | 13,078 ± 1,835        | 18,104 ± 2,624  |
| DACs (avg num/retina)   | 608 ± 5               | 606 ± 11              | 617 ± 8.3 (Whitney et al., 2009) |
| RIBEYE/CtBP2 (avg ThA [pixels]/100 μm²) | 1,227 ± 169         | 478 ± 175             | 519 ± 366 |
| Connexin36 (avg ThA [pixels]/100 μm²) | 469 ± 15             | 495 ± 210             | 394 ± 45  |
| Conventional synapses (avg num/0.005 mm²) | 324 ± 40            | 369 ± 42              | ND      |
| Ribbon synapses (avg num/0.005 mm²) | 95 ± 7              | 87 ± 4                | ND      |
| Globular ribbons (avg num/0.005 mm²) | 68 ± 5              | 53 ± 6                | ND      |

Abbreviations: avg, average; DAC, dopaminergic amacrine cell; ND, nondetermined; PI, photoinduction; RBC, rod bipolar cell; SCGN, secretagogin; ThA, thresholded area.

4 | DISCUSSION

High survival rates of rod and cone BCs demonstrate that the early effects of PR degeneration in the fully developed, adult retina of Tvrn4 mice are no more severe than in the developing retina of other retinal degeneration models (i.e., rd1 and rd10 mice), which are considered representative of human RP, but where there is overlap of PR degeneration and late phases of retinal development. Actually, at 6 weeks PI, the number of rod and cone BCs in Tvrn4 mutants remained close to 100%, unlike that found in rd1 and rd10 mice, where the RBC number dropped significantly by 3 months of age (Gargini et al., 2007). Surprisingly, however, and differently from what was observed in rd10 mice, where HC death overlaps spatially with PR loss, HCs of Tvrn4 mice die in small islands of the mid-central and mid-peripheral retina, not necessarily coinciding with the areas devoid of PRs. Remodeling and degeneration of HCs are most severe not in the center of the light exposure zone, but in the “transition” zones between high PR death and survival, respectively, where HC-free islands are separated by sharp edges from spots of surviving cells, which may persist in the central, most damaged retina, although with decreased intensity of fluorescence staining and hypertrophic processes.

As observed in the BC axonal terminals, an arrest in growth of RBC axonal arborizations in rd1 mice was previously reported (Strettoi et al., 2002), while the atrophy of RBC axonal endings in rd10 mutants was considered a secondary effect (Barhoum et al., 2008). Recent data also confirm that RBC axon terminals remodel, gradually losing complexity and undergoing progressive reduction as retinal degeneration proceeds (Phillips, Otteson, & Sherry, 2010). Indeed, Tvrn4 mice show these features as early as 3 weeks PI in the central, degenerated retina, where the complexity of RBC axonal endings decreases significantly. SCGN positive axonal endings in the IPL shrink progressively as well, but mainly in the ON sublamina of the IPL, as indicated by a thinning in the width of SCGN positive arborizations located in the innermost plexus (n.3) of the IPL. Altogether, these results confirm that rod-to-cone PR degeneration not only affects RBCs but gradually extends to other components of the ON pathway, which can be considered as a highly sensitive indicator of inner retinal remodeling.

Quantitative analysis of HCs at 6 weeks PI revealed a 28% reduction in the number of cells throughout the retina of Tvrn4 mice. This finding is in agreement with reports describing rd10 mutants at ~3 months of age (Gargini et al., 2007). Surprisingly, however, and differently from what was observed in rd10 mice, where HC death overlaps spatially with PR loss, HCs of Tvrn4 mice die in small islands of the mid-central and mid-peripheral retina, not necessarily coinciding with the areas devoid of PRs. Remodeling and degeneration of HCs globally reflecting the progression of typical rod-cone degeneration. These findings demonstrate that dendritic remodeling following PR death neither depends on the age of phenotype onset nor on the underlying genetic mutation or pattern of inheritance; rather, it is a secondary manifestation caused by PR death and represents a true hallmark of RP likely caused by deafferentation.
There is no obvious explanation for the higher severity of HC remodeling in transition zones of PR degeneration. An interesting hypothesis is that local signals (i.e., inflammatory molecules such as cytokines and chemokines, released from activating microglia), thought to contribute to cone death and to retinal abnormalities secondary to PR loss (Guadagni et al., 2019) are higher at the boundaries with regions where PR death is not yet complete, as though a wave of adverse signals preceded the actual phase of cell loss. Large-size neurons, similar to HCs, likely extend across these boundaries and suffer from the adverse effects of diffusible factors. Further investigation is necessary to address the nature of these noncell autonomous events. Yet, one has to remember that the peripheral, non-induced retina of Tvr4 mice can be considered entirely normal, unlike the peripheral retina of other mutants, where the phenotype is simply delayed with respect to the central retina. Local signaling is likely to bear intrinsic differences in Tvr4 and other paradigms of RP.

Morphological analysis of the IPL stained for anti-CtBP2/RIBEYE revealed excellent preservation of synaptic ribbons for up to 9 weeks PI in adult Tvr4 mice with respect to WT controls, with a significantly larger area occupied by RIBEYE positive puncta 3 weeks PI. This indicates the possibility of a peak in synaptic remodeling of the IPL at this time point, which might be driven by the peripheral, PR preserving retinal areas in the attempt of saving the flow of remnant visual signals despite extensive PR death in the central retina. Again, it is possible that long-range, diffusible signals (i.e., acting through intervening glia and blood vessels) may support these attempts of "reconstructive" remodeling, as BCs directly connected to dying PRs are not likely to extend their range of action beyond the light-induced area. Previous data reported a recovery phase in the Tvr4 retina at 4 weeks after photoinduction based on ERG recordings and histological results (Gargiuni et al., 2017). This was speculated to result from only partially damaged PRs at the edge of light-irradiated area, which could regenerate outer segments in 3–4 weeks provided a connecting cilium was still present. Increased density of synaptic contacts in the IPL could be triggered by this phase of PR recovery in the attempt to preserve the flow of information through the retina or anyway as an effect of a plastic stage of remodeling. The occurrence of migrating, remnant PRs converging in retinal foci has been reported (Busch, Gorgels, & Van Norren, 1999; Lin, Masland, & Strettoi, 2009; Paulus et al., 2008) and could explain the need to accommodate extra synaptic contacts in the IPL.

The IPL of Tvr4 retinas at 3 and 6 weeks PI was studied with the aid of TEM, confirming that rod bipolar endings have smaller sizes, and yet they are engaged in the typical RBC–All amacrine–A17 amacrine synaptic arrangements, which could be easily recognized. Quantitative analysis of synaptic contacts revealed no significant changes in the number of conventional (inhibitory) synapses in the overall width of the IPL; however, the total number of ribbon synapses displayed a decreasing tendency over time, with significantly fewer synaptic ribbons at 6 weeks PI compared to non-induced control retinas. Moreover, approximately 70% of all ribbons exhibited an abnormal globular or amorphous shape at 3 weeks PI while only 10% globular ribbons were found in the controls. Yet, the percentage of anomalous ribbons decreased to 60% by 6 weeks PI. Significantly larger areas covered by RIBEYE positive puncta in the IPL at 3 weeks PI observed by confocal microscopy can be explained by a larger area occupied by the same number of ribbons, implicating an increase of their average size. Most likely, globular ribbons observed at 3 weeks PI result in larger fluorescent profiles in the IPL, causing an increase in the area over the threshold. The decrease in the number of (globular) ribbons at 6 weeks PI agrees with our immunofluorescence observations.

Globular ribbons can be observed during early retinal development (Regus-Leidig et al., 2009) as precursors preceding full synaptic maturation and anchoring of these organelles at the presynaptic membrane. The globular form found here suggests altered turnover of the ribbons, as previously reported during artificial changes introduced in the dark–light cycle (Adly, Spiwoks–Becker, & Vollrath, 1999; Hull, Studholme, Yazulla, & von Gersdorff, 2006). Spherical (often free-floating) ribbons reminiscent of the globular organelles we observed have been described in the PR terminals of double-knockout mice for complexes (Cpx) 3 and 4, two regulatory proteins of vesicle exocytosis (Reim et al., 2009). Cpx3/4 double knockouts display retinal morphological and functional abnormalities altogether indicative of visual defects. Indeed, structural changes in ribbons have been demonstrated to affect coordinated multivesicular release of glutamate at the synapse (Mehta, Snellman, Chen, Li, & Zenisek, 2013). It is possible that RBCs of Tvr4 retina, deprived from rod input, undergo a downregulation of the dendritic and axonal machinery responsible for sensing and releasing glutamate, respectively. Additional functional studies are required to test the excitability and capability of synaptic transmission of BCs, in view of possible optogenetic therapy targeting these cells. However, our observations suggest overall stability of IPL ribbons, insofar 9 weeks PI the area covered by RIBEYE+ puncta in the IPL did not decrease further based on our IHCH data.

Analysis of CX36 distribution revealed a significantly larger surface occupied by immune positive puncta in the IPL of induced mice at all the time point studied, compared to WT controls, suggestive of an increase of gap junction density following PR degeneration. In degenerating retinal areas, Cx36 positive puncta are visibly distributed more extensively into the OFF layers of the IPL compared to the control samples (see Figure 7). Gap junctions were demonstrated to contribute to the spontaneous electrophysiological oscillations recorded from animal models of RP and human RP patients and pharmacological blockade of gap junctions was proven to decrease aberrant oscillatory activity and restore light responses driven by surviving PRs in the degenerating rd10 retina (Toychiev, Ivanova, Yee, & Sagdullaev, 2013). The increased expression level of Cx36 positive gap junctions observed in induced Tvr4 mice could represent a maladaptive change, causing oscillatory activity of inner neurons. It would be functionally relevant to prove the existence of oscillatory activity in Tvr4 mice, by multielectrode array (MEA) recordings from GCs through the time course of degeneration, and to verify the contribution of IPL gap junctions to inner retinal oscillations.

Because cell-to-cell coupling and the conductance of gap junctions are highly dependent on and regulated by dopamine levels in the vertebrate retina (Witkovsky, 2004), we studied the DAC network in...
Tvrm4 mice to correlate the morphology of DACs with changes in Cx36 immunofluorescence. TH labeled DACs showed normal appearance in the inner retina, with normal general architecture, process networks and number of cell bodies 3 and 6 weeks PI. However, processes reaching out to the OPL were found to be markedly hypertrophic in the central retina. Outer DAC dendrites were observed entangling dead PRs, represented by DNA-filled, membrane-bound bodies, and constituting typical side products of photoinduction. The area of hypertrophic processes precisely and exclusively colocalized with the area of light conditions (Witkovsky, 2004). It seems logical that dopamine release from DA amacrine reached a peak during the acute phase of inducing light exposure. However, there is no obvious explanation of the hypertrophy of the outer dopaminergic plexus observed here and of the obvious contiguity of DA dendrites and DNA-condensed profiles in the outer retina; moreover, DA dendrite hypertrophy persists well beyond the acute phase of photoinduction.

Calbindin D, ChAT, GAD67, and DAB1 antibodies were used to study GABAergic, cholinergergic, and glycinergergic (All) ACs in the retina of Tvrm4 mice 3 and 6 weeks PI. These markers revealed good preservation of all AC types based on anatomical observations. In particular, laminar organization of AC dendrites and cell body layering appeared normal. These results match previous reports of long-term AC survival rate in rd10 mice, where all AC types were found in normal physiological and morphological organization, except for All ACs, which were reported to be reduced in number and disorganized in distribution in rd10 animals of 40 days of age (Barhoum et al., 2008). Tvrm4 mice observed 6 weeks PI, a stage corresponding to the phenotype of P40 and older rd10 mice, showed considerable preservation of All ACs in the central retina, despite complete loss of PRs. Since All ACs are key members of the rod and cone signaling pathways and are considered a suitable target for optogenetic therapy, their long-term survival holds promising opportunities for vision restoration in RP. Once again, good preservation of All amacrine morphology indicates a high capacity of the retina to maintain a normal structural organization even when the disease phenotype manifests in adulthood and plasticity typically associated with a young age has been lost.

Previous studies on GCs in retinal degeneration rodent models predict no morphological alterations in these cells at relatively early stages of the retinal degeneration (Mazzoni et al., 2008; Damiani et al., 2012; Greferath et al., 2015). We did not observe alterations of the functional anatomical features of GCs in Tvrm4, light induced mutants. RBPMS and SMIM2 staining showed normal cell density, organization of cell bodies, dendrites, and axonal bundles of these cells in vertical sections both at 3 and 6 weeks PI. Again, we point out to the need of a complete electrophysiological characterization of the Tvrm4 retina, particularly at the GC level, in view of previous reports about the presence of a spontaneous, aberrant, oscillatory activity in the local field potentials of GCs in both rd1 and rd10 retinas recorded with MEA (Goo et al., 2011; Stasheff, Shankar, & Andrews, 2011).

Based on remodeling data obtained from adult Tvrm4 mice in the present study—confirming and expanding previous work in other animal models of RP—we conclude that the process of inherited, rod-to-cone degeneration triggers morphological remodeling of inner retinal neurons first acting on components of the rod system and of the ON signaling pathway, as proven by dendritic retraction from RBCs in the OPL and (later) of ON BC axonal ending shrinkage in the ON sub-lamina of the IPL. Contacts in the IPL are highly preserved, albeit with some ultrastructural abnormalities of synaptic ribbons. Preliminary observations from our TEM analysis in the IPL of Tvrm4 mice 3 and 6 weeks PI show ultrastructural signs typical of autophagy and/or parapoptosis (Wei et al., 2015), including the presence of swollen mitochondria with broken or diminished inner membrane system with clear signs of invaginations, empty vacuoles as well as concentric multilamellar compartments. Experimental evidence of autophagy in the IPL would suggest a compensatory response of inner retinal neurons following PR death and loss of input from the outer retina, consequently indicating the possibility of reversing inner retinal remodeling. Quantification of, for example, degenerating mitochondria by TEM analysis and biochemical assessment of LC3-II (autophagy marker) and related proteins could provide indications about autophagy in the IPL. At cellular level, engineering Tvrm4 mice expressing fluorescent marker proteins like GFP under the promoter of ON BC specific genes would allow single cell transcriptome analysis looking for autophagy-related expression patterns.

Noticeably, the phenotype of Tvrm4 mutants shares a number of features with non-neovascular “dry” age-related macular degeneration (ARMD), including light sensitivity (de Jong, 2006; Marc et al., 2008), focal occurrence of abnormalities in the central retina (de Jong, 2006), rod-to-cone PR death (Cucurio, Medeiros, & Millican, 1996), and the subsequent dendritic remodeling of second-order neurons (Jones & Marc, 2005; Sullivan, WoldeMussie, & Pow, 2007). However, clear documentation of inner retinal remodeling in human ARMD is only available for late stages of the disease (Jones et al., 2012), when vision restoration based on remnant retinal tissue is virtually impossible. Therefore, results obtained from photo-induced Tvrm4 mice may bear important possibilities to translate findings to early and mid-stages of ARMD and provide evidence of inner retinal ability to resist to PR degeneration, at least in the relatively short run. This encourages timely therapeutic approaches for this devastating disease.

Good preservation of the inner retina in Tvrm4 mice, an excellent model of RHO AD RP Type B1, offers the opportunity to test the efficacy of vision restoration on adult animals by optogenetic tools targeting ON BCs, that is, driving the expression of ChR2 (Legali et al., 2008; Macé et al., 2015), MAG460 (Gaub et al., 2014), ReaChR (Sengupta et al., 2016), ChrimsonR (Klapoetke et al., 2014), or naturally occurring retinal ON BC specific genes would allow single cell transcriptome analysis looking for autophagy-related expression patterns.

Good preservation of the inner retina in Tvrm4 mice, an excellent model of RHO AD RP Type B1, offers the opportunity to test the efficacy of vision restoration on adult animals by optogenetic tools targeting ON BCs, that is, driving the expression of ChR2 (Legali et al., 2008; Macé et al., 2015), MAG460 (Gaub et al., 2014), ReaChR (Sengupta et al., 2016), ChrimsonR (Klapoetke et al., 2014), or naturally occurring retinal cone opsins (Berry et al., 2019). This is particularly important in view of the present difficulty to treat dominant (gain-of function) mutations by gene therapy directed to PRs: maintenance of the inner retina for sometimes after total PR loss offers therapeutic opportunities that exploit second-order and higher order neurons to restore vision.

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DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

ORCID
Enrica Strettoi https://orcid.org/0000-0002-6101-599X

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