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Original Research Communication

The thioredoxin encoded by the Rod-derived Cone Viability Factor gene protects cone photoreceptors against oxidative stress

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Abstract

Aims: Rod-derived cone viability factor long (RdCVFL) is an enzymatically active thioredoxin encoded by the nucleoredoxin-like-1 (*Nxn1*) gene. The second product of the gene, RdCVF, made by alternative splicing is a novel trophic factor secreted by rods that protects cones in rodent models of retinitis pigmentosa, the most prevalent inherited retinal disease. It acts on the cones by stimulating aerobic glycolysis through its interaction with a complex containing basigin-1 and the glucose transport GLUT1. We studied the role of *Nxn1* in cones after its homologous recombination using a transgenic line expressing Cre recombinase under the control of a cone opsin promoter.

Results: We show that the cones of these mice are dysfunctional and degenerate by 8 months of age. The age-related deficit in cones is exacerbated in young animals by exposure to high level of oxygen. In agreement with this phenotype, we found that the cones express only one of the two *Nxn1* gene products, the thioredoxin RdCVFL. Administration of RdCVFL to the mouse carrying a deletion of the *Nxn1* gene in cones reduces the damage produced by oxidative stress. Silencing the expression of RdCVFL in cone-enriched culture reduces cell viability showing that RdCVFL is a cell-autonomous mechanism of protection.

Innovation: This novel mode of action is certainly relevant for the therapy of retinitis pigmentosa since the delivery into cones of the *rd10* mouse, a recessive model of the disease, rescues cones.

Conclusion: Our work highlights the duality of the *Nxn1* gene which protects the cones by two distinct mechanisms.

Introduction

Cone photoreceptors have been shown to be particularly vulnerable to oxidative stress (10,22,23,36,37). In that respect, it is of interest to notice that the nucleoredoxin-like one (*Nxn1*) gene, which encodes by alternative splicing rod-derived cone viability factor (RdCVF), also encodes RdCVFL that contains the thioredoxin fold as well as the catalytic CXXC domain (4,26). Thioredoxins participate in redox homeostasis by reducing oxidized thiol groups in target proteins (28). Thioredoxins are oxidized during that reaction and must then be reduced by thioredoxin reductases that may be redox sensors of oxidative environment (38). Specifically, RdCVFL interacts with the microtubule associated protein TAU and prevents its oxidation *in vitro*. It also prevents its phosphorylation and aggregation in the retina (10,15). RdCVFL protects rod photoreceptors against photo-oxidative damage (12) and reduces the oxidation of polyunsaturated fatty acids induced by photoreceptor degeneration in the *rd10* mouse (6). The *rd10* mouse is a model of retinitis pigmentosa, the most common form of inherited retinal degenerations. We have shown that delivering RdCVFL to the retina under the control of a ubiquitous promoter does not prevent cone degeneration, unlike RdCVF (11). RdCVF production results from intron retention following transcription of the *Nxn1* gene. The phylogenetic conservation of an in frame stop codon in RdCVF messenger produces a shorter protein, truncated within its thioredoxin motif (8). The 109 residues of RdCVF are identical to the N-terminal region of the RdCVFL protein which is 217 amino acids long. RdCVF has no thioloxidoreductase activity even though it contains the thioredoxin catalytic site sequence CXXC (26). The effect of RdCVF on cones is mediated by its interaction with basigin-1 (BSG1), a splice variant of the *Bsg* gene expressed specifically by photoreceptors (1). The interaction of RdCVF with BSG1 at the surface of the cones stimulates glucose uptake by accelerating glucose transport by GLUT1. Cone rescue is the result of the production of metabolites by aerobic glycolysis that are probably used for cone outer segments renewal to sustain cone

function. RdCVF is a promising therapeutic for retinitis pigmentosa, for which the degeneration of rods by a cell autonomous mechanism leads to night blindness, a mild condition, but is irremediably followed by the non-cell autonomous death of cones, which leads to loss of daylight vision (7,27).

We envisioned that in healthy organisms, the active thioredoxin RdCVFL would be the sensor of luminance in rods participating, somehow in the production of a proper level of trophic factor RdCVF to cones adapted to light environment. Here, we challenged this view by inactivating the *Nxn1* specifically in cones. We show that cones expressed the thioredoxin RdCVFL that is involved in their protection against oxidative stress generated during aging. Our work supports further development of a novel double-therapeutic approach for retinitis pigmentosa.

Results

Specific disruption of *Nxn1* gene in cones.

To inactivate the *Nxn1* gene in cones, we crossed the *Nxn1cd/cd* mouse line that carries two LoxP sites framing exon 1 of the gene (10) with the Cone-Cre mouse, a transgenic line that expresses Cre recombinase under the control of the 2.1 kb human OPN1L/MW (red and green opsins) promoter to obtain animals with the [Cone-Cre:*Nxn1cd/cd*] genotype and their controls [Cone-Cre:*Nxn1+/+*] (24). The recombination was validated and quantified using DNA extracted from sorted cells labeled with antibodies against mouse OPN1MW, the red/green opsins (**Supplementary Figure 1**). PCR primer pairs P1/P2 detected the wild-type (wt) alleles of the *Nxn1+/+* sorted cells and the pair P3/R4 the recombined (rc) alleles of the *Nxn1-/-* sorted cells (**Figure 1A** and **1B**). Cells sorted from the [Cone-Cre:*Nxn1cd/cd*] retina carry exclusively the rc allele at 554 base pair (bp) and cells sorted from the [Cone-Cre:*Nxn1+/+*] retina, carry the wt allele at 575 bp. The amplification of the *Pde6b* gene was used as an internal control. In order to quantify the recombination, we performed quantitative PCR analysis of the rc allele on DNA isolated from the whole retina using the primer pair P3/R4. 100% and 0% recombination correspond to the value obtained with *Nxn1-/-* and *Nxn1+/+* retinas respectively (**Figure 1C**). This scale shows that the recombination took place in about 2.5% of the [Cone-Cre:*Nxn1cd/cd*], a value that corresponds approximately to the percentage of MW cones in the mouse retina. We also quantified *Nxn1* gene expression in the retina using RT-PCR. Using primers internal to the recombined locus (P1/P2) we did not amplified the *Nxn1* messenger in the *Nxn1-/-* retina, as expected. We could not differentiate *Nxn1* expression level in [Cone-Cre:*Nxn1cd/cd*] retina from that of the [Cone-Cre:*Nxn1+/+*] mouse showing that the recombination did not occur in the rods (97% of all photoreceptors) and is consequently specific to cones (**Figure 1D**). Similarly, using specific primer pairs, we observed similar expression of RdCVF and RdCVFL messengers in [Cone-Cre:*Nxn1cd/cd*] and [Cone-

Cre:Nxn1cd/cd] retinas (**Figure 1E and F**). Using western blotting, we show that the expression of RdCVFL in the rods, the most numerous photoreceptors can be detected after recombination of *Nxn1* in the cones at 1 month of age (**Figure 1G**).

The function of cones was measured at 8 months of age using electroretinography (ERG). The traces show b-waves when tested by photopic ERG recording (**Figure 2A**). As previously reported, the amplitude of the photopic b-wave is reduced for *Nxn1*-/- mice compared to their control, the *Nxn1*+/+ mice (10) (**Figure 2A and B**). Surprisingly, while the b-wave amplitude for the [Cone-Cre:*Nxn1*+/+] is comparable to that of the *Nxn1*+/+ mice, that of the [Cone-Cre:*Nxn1cd/cd*] resembles that of the *Nxn1*-/- mouse even though the cones (comprising only 3% of photoreceptors in the mouse retina) are surrounded by rods expressing the trophic factor RdCVF. We noticed that the trace of the b-wave differs between the *Nxn1*+/+ mice and the [Cone-Cre:*Nxn1*+/+]. Since the only difference between the two strains is that cones of the [Cone-Cre:*Nxn1*+/+] mice express the recombinase Cre under the control 2.1 kb human OPN1L/MW promoter, the difference might be linked to an unknown effect of the recombinase on ERG response but more likely a phenomenon of transcriptional interference between the locus control region (LCR) of the transgene with the endogenous expression of cone opsins resulting in a reduction of cone opsin expression revealed in the trace of the b-wave (21,34). This effect may be mediated by transcriptional squelching for transcription factors binding to the LCR (16,17). The phenomenon is not strong enough to modify b-wave amplitude and does not interfere with our conclusions. Since cone-photopic ERG is a mixed ON-cone-bipolar and Muller cells response, we used flicker ERG that records specifically cone responses (**Figure 2C**). Here again, [Cone-Cre:*Nxn1cd/cd*] flicker amplitude at 10 Hz and 20 Hz is reduced compared to [Cone-Cre:*Nxn1*+/+] control mice (**Figure 2D, E and Supplementary Figure 2A**). We also look at the implicit time that scores cone photoreceptor response independently of the amount of light delivered to the retina (**Supplementary Figure 2B**). The

implicit time is retarded by 12 ms (56 to 68) in the [Cone-Cre:*Nxn1cd/cd*] mice compared to the [Cone-Cre:*Nxn1+/+*] mice (**Supplementary Figure 2C**). We sacrificed the animals, labeled the cones with a mixture of OPN1MW and OPN1SW antibodies and measured the density of the cones over the entire surface of the retina (1,9). The density of cones was found to be lower and similar for the *Nxn1/-* and the [Cone-Cre:*Nxn1cd/cd*] retinas compared to that of the [Cone-Cre:*Nxn1+/+*] retinas (**Figure 2F and G**). When we measured the density of cones at 4 months of age, we also found it lower for the [Cone-Cre:*Nxn1cd/cd*] retinas compared to that of the [Cone-Cre:*Nxn1+/+*] retinas (**Supplementary Figure 2D**). When we observed the flat-mounted retina at higher magnification, we observed that the labeling of opsins was distinct between the two genotypes in the ventro-nasal part of the retina (**Supplementary Figure S2E**). This is certainly a indication of a dimorphism of cones of the [Cone-Cre:*Nxn1cd/cd*] mice that precedes their degeneration in this area.

The cones are sensitive to oxidative stress even in the presence of rods.

Since the cones in the [Cone-Cre:*Nxn1cd/cd*] retinas are exposed to the trophic activity of RdCVF secreted by rods (1), their death cannot arise from a lack of trophic support. We housed one-month old animals in two different conditions. One of the groups was maintained in normoxic conditions (21% O₂), while the other group was maintained for one week in hyperoxia, in an atmosphere composed of 75% oxygen. At this age, photopic ERG responses of [Cone-Cre:*Nxn1cd/cd*] and [Cone-Cre:*Nxn1+/+*] control mice are more similar than traces from mice at 8 months of age in line with the progression of cone degeneration with age (**Figure 3A**). Hyperoxia reduces photopic b-wave amplitudes for all genotypes but the amplitudes for the [Cone-Cre:*Nxn1+/+*] mice under hyperoxia match those of the [Cone-Cre:*Nxn1cd/cd*] mice under normoxia (**Figure 3A and B**). Interestingly, the reduction in amplitude is similar for both the *Nxn1/-* and the [Cone-Cre:*Nxn1cd/cd*] mice suggesting that the observed deficit is independent of RdCVF. When looking at a pure cone response using flicker ERG, the

phenomenon was exacerbated (**Figure 3C**). Under normoxia, the three groups of animals tested have quite similar responses (**Figure 3D**). Under hyperoxia, the amplitude of the [Cone-Cre:*Nxn1*+/+] flicker ERG is only marginally reduced by O₂ corroborating previously published results (10). Using this assay, the contribution of RdCVF could be observed with a more dramatic reduction of the flicker amplitude at 10 Hz and 20 Hz (**Figure 3C, 3D, Supplementary Figure 3A and B**) for the *Nxn1*^{-/-} compared to the [Cone-Cre:*Nxn1*cd/cd] mice. After sacrifice, the retinas were labeled with a mixture of OPN1MW and OPN1SW antibodies and cone density was measured. Here again, the loss of cone function under hypoxia paralleled a reduction in the number of cones in the [Cone-Cre:*Nxn1*cd/cd] mice (**Figure 3E and F**). The similarity in cone density of the *Nxn1*^{-/-} and the [Cone-Cre:*Nxn1*cd/cd] mice under hyperoxia, while they differ in cone flicker ERG amplitudes, may indicate that in the *Nxn1*^{-/-} mice without trophic support of RdCVF, cones loss their function before they degenerate. We examined the function and the survival of rods in the mice at 1 month of age in the presence or absence of oxidative stress. Scotopic ERG were recorded (**Supplementary Figure 3C**) and a-wave, characteristic of rod function was found unaffected under increasing flash intensity in any conditions (**Supplementary Figure 3D**). We found that the b-wave that results from more downstream signaling at the level of bipolar cells shows a trend toward lower amplitude for [Cone-Cre:*Nxn1*cd/cd] mice as well as reduced amplitude in 75% oxygen (**Supplementary Figure 3E**). The thickness of the outer nuclear layer (ONL), composed of 97% rods was measured by optical coherence tomography in those animals (**Supplementary Figure 3F**). No differences could be observed between the two genotypes. Along the temporo-nasal retinal axis, the ONL was found slightly reduced in the [Cone-Cre:*Nxn1*+/+] mice exposed to 75% oxygen on the temporal side (**Supplementary Figure 3G**). Along the dorso-ventral retinal axis, the ONL was found slightly reduced in the [Cone-Cre:*Nxn1*+/+] mice exposed to 75% oxygen on the ventral side (**Supplementary Figure 3G**). Overall, these results show that the function and the survival of rods are not affected by deletion of *Nxn1* in 3% of photoreceptors, the cones. The

contralateral eye of the mice were enucleated and fixed to perform immunohistochemistry. We observed, as previously reported, that the *Nxn1*^{-/-} retina shows signs of oxidative damage using both anti-acrolein or 4-hydroxynonenal (HNE), two products of lipid peroxidation under oxidative stress (10) (**Figure 4**). Under hyperoxia, both [Cone-Cre:*Nxn1*^{+/+}] and [Cone-Cre:*Nxn1*^{cd/cd}] retinas were labeled with these antibodies, but the acrolein staining was more pronounced in the later group.

The cones express RdCVFL, the thioredoxin protein product of the *Nxn1* gene.

Since [Cone-Cre:*Nxn1*^{cd/cd}] mice have an age-dependant deficit in cone function and survival that can be accelerated by hypoxia, most likely the *Nxn1* gene is expressed by cones in wild-type (wt) retinas. We produced a reporter adeno-associated viral (AAV) vector expressing GFP under the control of the human OPN1L/MW promoter (39). We used an AAV variant called AAV2-7m8 that can target photoreceptors following intravitreal injection (11). On retinal explants of wt mice (BALB/c) injected with this vector, the fluorescence of GFP is co-localized with the expression of OPN1MW revealed in red by immunohistochemistry (**Figure 5A**). Individual GFP-positive cells were collected by micro-aspiration with patch-clamp glass electrodes. Current responses to voltage steps were recorded in some of these cells using the patch-clamp technique. Voltage steps ranged from -100 mV to 40 mV (10 mV increments). Below -60 mV, only weak linear inward currents were observed, and the cells typically exhibited a large outward current at potentials above -40 mV, as expected for cone photoreceptors (**Figure 5B and C**) (32). After patch-clamp recordings the cones were used for single-cell RT-PCR analysis. We used a negative (0) and a positive control that correspond to no cDNA and cDNA from wt retina (BALB/c) at post-natal day (PN) 40 respectively. The GFP cDNA shows that the RNA was collected from cones that do not express the rod photoreceptor marker, *Rho* (**Figure 5D**). We could amplify with specific primers the messenger of the thioredoxin RdCVFL but not that of the trophic factor RdCVF. For further validation, we repeated the analyses on 12

independent GFP-positive cells that all gave the same result (**Figure 5E**). The cones, at least the MW-cones, express the thioredoxin RdCVFL exclusively.

The thioredoxin RdCVFL protects cones against oxidative stress.

To further establish the role of the thioredoxin RdCVFL in the observed deficit of the cones, the AAV2/8 virus, that drives expression of RdCVFL or RdCVF or GFP under a CMV promoter were employed. The subretinal injection of this virus serotype can transduce photoreceptors efficiently (29). We delivered either RdCVF or RdCVFL by subretinal injection of recombinant AAV2/8 vectors into the [Cone-Cre:*Nxnl1cd/cd*] mouse at postnatal days 20 (PN20) and housed the animals for five days, between PN35 and 40, in 75% O₂. The reduction of the photopic b-wave amplitude in uninjected mice shows that the cones of the [Cone-Cre:*Nxnl1cd/cd*] mice were affected by hyperoxia at PN40 (**Figure 6A**). The delivery of GFP or RdCVF had no effect on the ERG b-wave amplitude and on its shape, the later revealed by the contralateral uninjected eye (**Figure 6B, C and D**) in the [Cone-Cre:*Nxnl1cd/cd*] mice. In contrast, RdCVFL prevented the reduction of the amplitude by 48% [(126 – 103) / (151 – 103)] compared to [Cone-Cre:*Nxnl1+/+*] control mice. The flicker ERG was also recorded (**Figure 6G**). Using this readout, the results show a 46% [(110 – 78) / (147 – 78)] protection by RdCVFL (**Figure 6E, F, Supplementary Figure 4A-C**). The implicit time was shortened from 63 ms for animals treated with AAV-GFP to 60.5 ms for those treated with AAV-RdCVFL (**Supplementary Figure 4D**). The data are correlated (**Supplementary Figure 4E**). The animals were sacrificed and the cone density was quantified after labeling cones with peanut agglutinin (PNA), lectin from *Arachis hypogaea*. RdCVFL protected the cones by 63% [(4.7 – 4.2) / (5.0 – 4.2)] (**Figure 6H and I**).

We have previously reported that cones exclusively express the RdCVFL (and not the RdCVF) messenger RNA in cone-enriched culture from chicken embryos (15). Therefore, we

silenced the expression of RdCVFL in the cell culture system by electroporating a miRNA plasmid (1). In order to validate the efficacy of the miRNA molecule, we cloned the open reading frame of chicken RdCVFL under a CMV promoter, 3' to *Renilla reniformis* luciferase. The plasmids were co-transfected in HEK293 cells. 48 h later, the luciferase activity showed that the miRNA is efficient (**Figure 7A**). The viability of the cells was scored after 7 days *in vitro* (DIV) by counting the number of cells transduced by the RFP reporter (**Figure 7C**). Specific silencing of endogenous RdCVFL reduces cell survival in a dose-dependent manner compared to a scrambled miRNA construct (**Figure 7B**).

The thioredoxin RdCVFL protects cones in a model of retinitis pigmentosa.

We have previously shown that in the *rd1* mouse, a model of recessive retinitis pigmentosa, the expression of *Nxnl1* is rod-dependent (33). *Nxnl1* expression by bipolar cells is lost following rod degeneration even if the bipolar cells are not affected in that model. We reasoned that the expression of RdCVFL by cones may also be lost following rod degeneration. We used a similar genetic model, the *rd10* mouse that carries a mutation in the same gene, *Pde6b*, but with slower kinetic of degeneration, thereby allowing functional evaluation (6). We produced novel AAV2/8 vectors for transgene delivery to mouse cones under the control of the human OPN1L/MW promoter. The *rd10* mice were subretinally injected at PN20 in one eye with AAV2/8 vectors and the function of the cones was recorded at PN40, after rods had degenerated. There was a net decrease in cone function in the *rd10* mice compared to wt mice (**Figure 8A and C**). The transduction of cones with recombinant AAV vectors does not affect the shape of the photopic b-wave as seen in the contralateral uninjected eye (**Figure 8B**). Using this paradigm, RdCVF was not able to protect the cones, while RdCVFL protected cones by 36% [(102 – 55) / (187 – 55)]. We also recorded the flicker ERG in those same animals (**Figure 8D, E, F, and Supplementary Figure 5A**). RdCVFL generated a 45% protection. We also noticed that the photopic and flicker ERG measurements are correlated for the 42 *rd10* eyes

included in the experiment (**Supplementary Figure 5B**). After sacrificing the mice, cone density measurements showed that RdCVFL prevented secondary cone death by 47% $[(5.2 - 4.3) / (6.2 - 4.3)]$ (**Figure 8G and H**).

Discussion

We have demonstrated that the thioredoxin RdCVFL is normally expressed by cones as well as rods in the mouse retina. The role of this enzyme is to prevent the damage to cones observed in aged animals. Retinal damage is accelerated when the mice are exposed to a high level of oxygen. This cell-autonomous mechanism is very probably involved in the secondary cone degeneration in retinitis pigmentosa. In this disease, the loss of expression of the trophic factor RdCVF results in the lack of stimulation of aerobic glycolysis through its cell-surface receptor BSG1. The cones are deprived of glucose and their outer segments shorten because of the lack of production of proper glucose metabolites (1,7,25). The reduction of the size of the cone outer segment is apparently not followed immediately by their death explaining why non-functional cones can be observed in the retinas of blind patients affected by retinitis pigmentosa (5). We have previously reported that the function of the cones in *Nxn1*-/- mice deteriorates when the animals are exposed to hyperoxia (10). The deficit likely results from the absence of expression of RdCVFL in the cones rather than the lack of protection by the rods. What could be the mechanism explaining the vulnerability of cones to oxidation? For retinitis pigmentosa, the problem arises from the absence of regulation of choroidal circulation (35). The delivery of oxygen from the blood is maintained in the rod-less retina, even if its consumption is reduced by 97%. Consequently, the retinal oxygen pressure reaches values that approximate hyperoxia. The cones and their outer segments composed of polyunsaturated fatty acids are damaged by reactive oxygen species produced from leakage of the respiratory chain. In the absence of *Nxn1*, the retina shows signs of lipid peroxidation (10). After specific recombination of *Nxn1* gene in the cones, the retina also displayed signs of oxidative damage (**Figure 4**). The viability of the cones of the cone-enriched cultures depends on the endogenous protection by the thioredoxin RdCVFL. In that system, we showed that the survival by ectopic RdCVF is not dependent on redox homeostasis but on aerobic glycolysis (1). Nevertheless, in the retina, the

thiol-oxidoreductase activity of RdCVFL depends on its cycling between a reduced and an oxidized status as any other thioredoxin. Once oxidized, RdCVFL must be reduced by thioredoxin reductases (20). The co-factor of thioredoxin reductases, NADPH, is produced mostly from glucose through the pentose phosphate pathway (PPP) (**Figure 9**). In that respect, in cones, RdCVF likely acts upstream of RdCVFL by providing reducing power. Many of the glycolytic enzymes are negatively regulated by oxidation of thiol groups at cysteine residues, like glyceraldehydes-3-phosphate dehydrogenase and pyruvate kinase (PKM) (2,19). Oxidation of PKM at cysteine 358 (C358) stops the production of pyruvate and results in the accumulation of phosphoenol pyruvate (PEP). PEP, an allosteric inhibitor of triose phosphate isomerase, interrupts glycolysis which produces an elevation of the intracellular concentration of glucose-6-phosphate (G6P). G6P, produced by the action of hexokinases on glucose entry to the cell via glucose transporters, is diverted to produce ribulose-5-phosphate (R5P) in PPP, enzymatic steps catalyzed by glucose-6-phosphate dehydrogenase (G6PDH) and 6-phosphogluconate dehydrogenase (PGD). Both G6PDH and PGD use NADP⁺ as a co-factor and that is reduced to NADPH during the reaction. NADPH is used by glutathione reductase to reduce glutathione and by thioredoxin reductases to reduce thioredoxins. Reduced RdCVFL could reestablish aerobic glycolysis in cones after oxidative stress and we have identified a potential interaction between RdCVFL and PKM (15). One could speculate that in patients suffering from retinitis pigmentosa after the loss of expression of RdCVF produced by rods, cones become non-functional and die within several years as a result of the loss RdCVFL expression and subsequent oxidative damage in cones. We have previously observed that the expression of *Nxn1* in cells neighboring the rods depends on the presence of rods (33). Thus, a therapy aimed at preventing secondary cone degeneration should be pursued using both RdCVF and RdCVFL. While RdCVF could be delivered to cones by any surrounding cells, excluding dying rods, RdCVFL would have to be transduced to the cone themselves to exert the cell-autonomous protection observed here. This might be achieved using ubiquitous or cone-restricted promoters.

Regarding the evolution of RdCVF signaling, it is quite reasonable to propose that the thioredoxin activity precedes the trophic activity of RdCVF, the latter resulting from an alternative splicing event that produces a truncated thioredoxin (18). Cones evolved earlier than rods, the latter appeared during evolution at a period when a reduction in energy consumption favored cephalization (13). The alternative splicing event at the origin of the RdCVF metabolite signaling should be anterior to the diversification of the nucleoredoxin-like family by duplication of an ancestral *Nxn1* gene into *Nxn1* and *Nxn2* (6,8). Interestingly, the lamprey, a jaw-less fish, harbors two types of photoreceptors with cone morphology. One of them is possibly a rod in regard to its spectral sensitivity (31). Prior to its role in rod-cone interactions, the ancestral *Nxn1* gene may have been involved in redox homeostasis and possibly contributed to glycolysis.

Innovation

Retinitis pigmentosa is a blinding disease characterized by the sequential loss of rod photoreceptors followed by the cones. The nucleoredoxin-like 1 (*Nxn*/1) gene encodes by alternative splicing for a trophic factor, rod-derived cone viability factor (RdCVF) that is secreted by rods and protects cones by stimulating aerobic glycolysis. We show that the other product of this gene, the thioredoxin protein RdCVFL, is expressed by cones where it exerts protection against oxidative stress. This suggests that the protection of cone vision in retinitis pigmentosa should benefit from a double therapeutic approach where both products of the *Nxn*/1 gene are administrated.

Material and methods

Animals.

The *Nxnl1*-/- mice on BALB/c background were generated previously (10). The BALB/c (*Nxnl1*+/+) mice were used as wt mice. The *Nxnl1cd*/cd mice on BALB/c background contain loxP sites frame exon 1 of the *Nxnl1* gene, and the Cone-Cre mice with Cre recombinase directed by the human red/green pigment (HRGP) gene were kindly provided by Yun Zheng Le (24). Cone-Cre mice were crossed with *Nxnl1*+/+ mice more than 10 generations to reach BALB/c background. [Cone-Cre:*Nxnl1*+/+] and [Cone-Cre:*Nxnl1cd*/cd] mice were obtained by crossing *Nxnl1cd*/cd mice with Cone-Cre mice. The *rd10* and C57BL/6J mice were purchased from Jackson laboratory. All mice were housed under the light/dark (12/12 h) cycle, 23-25°C room temperature and offered *ad libitum* access to food and water. The project was approved by French Ethics Committee (N°A-75-1863; OGM n°5080 CA-II). And all experiments were performed in accordance with the European Community Council Directives of September 22, 2010 (2010/63/UE).

Retinal cells preparation for FACS cone sorting.

Mouse eyes were enucleated, their retina dissected. Retinas (2 retinas/tube) were dissociated in 450 µl Hank's balanced salt solution (HBSS, no calcium) with 50 µg trypsin for 5 min at 37°C. To stop the reaction, the cells were mixed well with 1,000 µg trypsin inhibitor in 450 µl HBSS (with calcium) and 50 µg Dnase I by using a pipette and centrifuge 3 min at 2,000 g to keep the dissociated cells. The cell pellet was resuspended in 500 µl phosphate buffer saline (PBS), and fixed for 30 min in 2% paraformaldehyde. The fixed cells were then washed with PBS and blocked in PBS containing 0.4% bovine serum albumin (BSA), 1% fetal bovine serum (FBS) and

1 mg/ml saponin to proceed for immunolabeling with rabbit anti-OPN1MW antibodies (1/40, Millipore AB5405) or negative control rabbit anti-IgG (1/40), and then secondary antibodies: R-phycoerythrin (1/50, Life technologies). The labeled cells were washed and maintained in 500 µl PBS for FACS cone sorting.

Analysis of homologous recombination by PCR.

Genomic DNA from FACS-sorted cones was extracted with DNeasy Blood & Tissue Kit (Qiagen, 69504). PCR analysis were carried out using primers P1 (5'-TACCTTGAGTGACACACC-3') / P2 (5'-CCCTCCCTGGTCTTCCCTGGGG-3') for wt allele, and primers P3 (5'-TCCACACACAGCACATCATACG-3') / R4 (5'-TGTGATGGAAGAGGGATGGAAC-3') for recombined allele.

Quantitative-PCR analysis.

Total RNAs of mouse retina were extracted with kit (RNeasy mini kit, Qiagen). The RNA was reverse-transcribed with SuperScript® II reverse transcriptase kit (Life technologies). Samples have been analyzed by quantitative PCR using power SYBR green PCR master mix (Life technologies) in a 7500 standard real-time PCR system (Applied biosystems) with primers P1/P2, or P3/R4, RdCVF (5'-CTACAGAGGAGAACAGGAC-3'; 3'-TGCACAAGTAGTACCAAGGAC-5') or RdCVFL (5'-GCAACAGGACCTTCCCTCA-3'; 3'-CCAGACGCTGGATCTCCTC-5').

Western-blotting.

Western blotting on mouse neural retina was performed as previously described using rabbit polyclonal antibodies RdCVF-N (26).

Electroretinograms.

Mice were anesthetized intraperitoneally with a mixture of ketamine (80 mg/kg) and xylazine (10 mg/kg) diluted in saline. Their pupils were dilated by topical application of 0.5% mydriaticum. Body temperature was maintained at 37°C with a circulating hot water heating pad. The electrical signal was recorded using a pair of electrodes constructed in our platform adapted for use on mice. A gold loop electrode was placed on the center of the corneal surface and maintained with ocrygel (TVM) to ensure good electrical contact. Two stainless steel reference electrodes were inserted subcutaneously in each cheek of the mouse to normalize signal output, and the ground electrode was inserted subcutaneously in the back of the mouse. Recordings were made from both eyes simultaneously. Recording measurements: The light stimulus was provided by a 150 W xenon lamp in a Ganzfeld stimulator (Multiliner vision, Jaeger toennies, Germany). Responses were amplified and filtered (1 Hz-low and 300 Hz-high cut off filters) with a 1 channel DC-/AC-amplifier. To isolate cone responses in photopic, a 5 min light saturation at 10 cds/m² was used to desensitize the rods. The cone photopic ERGs shown represents the average of 10 responses from 10 consecutive flashes at 10 cds/m² intensity. The flicker ERG was also used to isolate cone responses at flash frequencies of 10, and 20 Hz at 3 cds/m² intensity. Data acquisition: The photopic b-wave amplitudes were measured from the base-line to the peak. For scotopic recordings, single flash recordings were obtained at light intensities of 3x10⁻³, 3x10⁻², 3x10⁻¹, 3 and 10 cds/m² using a sampling frequency of 5 kHz, a flash duration of 4 ms, and a frequency stimulus of 0.5 Hz. Data were recorded from 50 ms before stimulus onset to 450 ms post-stimulus. Data acquisition: The scotopic a- and b-wave amplitudes were measured from the base-line to the peak.

Optical coherence tomography (OCT).

Mice aged of 1 month were anesthetized and pupils were dilated. Eye dehydration was prevented by regular instillation of sodium chloride drops. OCT images were recorded on the right eye using a spectral domain ophthalmic imaging system (Spectral domain Optical coherence tomography, OCT, Bioptigen 840 nm HHP; Bioptigen; North Carolina USA). We performed rectangular scans consisting of a 2 mm by 2 mm perimeter with 1000 A-scans per B-scan with a total B-scan amount of 100. Scans were obtained while centered on the optic nerve. OCT scans were exported and analyzed using ImageJ. Outer nuclear layer (ONL) thickness was measured every 100 µm, ventral-dorsal and temporal-nassal, starting 100 µm from the center (optic nerve).

Hyperoxia treatment.

Mice were maintained in a 75% oxygen chamber for seven days with free access to food and water (BioSpherix, NY 13083).

Immunohistochemistry.

The eyes were enucleated quickly and immersed in 4% paraformaldehyde in PBS overnight at 4°C. They were incubated successively in 10% and 20% sucrose at 4°C and then embedded in optimal cutting temperature compound (Sakura finetek, Gentaur, Belgium). Frozen sections of 10 µm thickness were obtained for immunohistochemistry. The sections were first pre-incubated 2×30 min in blocking buffer 5% BSA and tween-20 0.1% in PBS). They were then processed to overnight incubation at 4 °C in 4-Hydroxynonenal (HNE) (Calbiochem, 1/300) or acrolein (Cosmo bio, Tokyo, Japan, 1/150) antibodies. After two washes of 30 min, the sections were

incubated 2 h at room temperature with goat anti-mouse IgG conjugated to alexa 594 (1/500), lectin PNA conjugated to alexa 488 (1/100, Life technologies) and the nuclear marker DAPI (Sigma, 1/1000). The sections were washed 3 times with PBS before ultimately mounted with fluoromount-G (SouthernBiotech). Images have been obtained with a confocal microscope (Olympus FV1000) at 63x objective. Stacks of consecutive images taken at 500 nm intervals were acquired sequentially with lasers: diode 559 nm, argon 488 nm and diode 405 nm and Z projections of serial sections were reconstructed in the software (FV-10-ASW-4.2) coupled to confocal.

Cone density measurement.

Cone counting was performed after ERG recording. The retinas were carefully dissected in pre-warmed (37°C) PBS (Gibco, Life technologies) and transferred to 4% paraformaldehyde overnight at 4°C. The retinas were prepared for immunostaining as described (9). After 1 h 30 min saturation in PBS containing 1% BSA, 0.1% tween and 10% normal goat serum (NGS), the retinas were incubated with a mixture of OPN1SW [1/400, (14)] and OPN1MW antibodies (1/200, Millipore) in the blocking buffer overnight at 4°C. They were washed 3×15 min in PBS containing 0.1% tween-20 and 1% BSA (PBS*) for goat anti-rabbit IgG conjugated to alexa 488 (1/400) incubation for 2 h at room temperature. Then they were re-washed 3×15 min in PBS*, counterstained by 3 h incubation in lectin PNA conjugated to alexa 488 (1/100) (3) in the blocking buffer. After final washing in PBS, the retinas were whole-mounted photoreceptor side-up in fluoromount-G. Cone density was analyzed in an automated system developed (e-conome) as described previously (9). Raw data are available on request.

Single cell recording and harvest.

The AAV2-7m8 OPN1L/MW-GFP injected eyes were enucleated; their retinas were dissected and dissociated 5 min by papain (1 unit/retina). Dissociated cells were then resuspended in RINGER solution (125 mM NaCl, 3.6 mM KCl, 1.18 mM MgCl₂, 22.6 mM NaHCO₃, 0.02 mM NaH₂PO₄, 0.028 mM Na₂HPO₄, 1.2 mM Na₂SO₄, 10 mM Glucose, 0.54 mM EDTA) for whole cell recording. Recording pipettes were pulled from borosilicate glass capillaries (BF100-50-10, Sutter Instruments) with a next generation micropipette puller (P-1000; Sutter Instruments). Single GFP-expressed cell was voltage-clamped to measure membrane currents. The recorded cell was collected in 5 µl PBS in a 1.5 ml Eppendorf tube.

Single cell RT-PCR analysis.

Single cell cDNA was amplified according to the protocol described in Morris *et al.* (30).

AAV injection.

Mice were anesthetized with isoflurane (IsoFlo, France) inhalation. They were first placed in a container connected with the vaporizer (Tec 7, Anestéo, France). Anesthesia was induced by inhalation of 5% vaporized isoflurane within 30 s. During the injection, anesthesia was maintained by mask inhalation of 2% vaporized isoflurane (debit 2 l/min). For intravitreal injection, the micro-syringe (10 µl, Hamilton, Reno, NY) was gently inserted into the vitreous cavity. A total of 2×10^{10} vg (viral genome) AAV in a 1 µl volume was subsequently injected into the vitreous cavity. For sub-retinal injection, a small hole was performed with a 10-0 needle in the superior sclera. The micro-syringe was then gently inserted into the sub-retinal space, and 1 µl of AAV (2×10^{13} vg/ml titer) was injected. The local retinal detachment was checked to confirm the success introduction of the virus.

miRNA construction and electroporation in cone-enriched culture.

The miRNA sequence used was designed with BLOCK-iT Pol II miR RNAi (Life technologies). The miRdCVFL (5'-AGTGTGCCAGGAGAAGCATCA-3'), targets the sequence at positions 512-532 from chicken RdCVFL mRNA 5'-end (XM_423688). This miRdCVFL or its scrambled (5'-GAGTACAAGTCGCGTACGAAG-3') was cloned into pcDNA6.2 GW/EmGFP-miR (Life technologies) vectors. At the same time, chicken RdCVFL open reading frame (ORF) sequence (624 bp) was cloned under the control of a CMV promoter in pRL-CMV (Promega), 3' to the *Renilla reniformis* luciferase to obtain pRL-CMV-RdCVFL-Ren. This plasmid was co-transfected with miRdCVFL or scramble plasmid in HEK293 cells using Lipofectamine 2000 in OptiMEM medium (Life technologies). 48 h later, *Renilla* luciferase activity was measured from cell extracts using dual-luciferase reporter assay system (Promega) to validate the efficacy of the miRdCVFL molecule. For live-dead assay, the delivery of the miRdCVFL from pcDNA6.2 GW/EmGFP-miRdCVFL construct and the RFP-reporter from pCAG-G-RFP plasmid was performed by co-electroporation into cone-enriched-chicken cultures as previously described (1). The RFP-reporter was served as a marker of the transduced cells. Seven days after electroporation, the RFP-expressing living cells were counted on an automated microscopy system.

Statistics.

All measurements were made using blind protocols. Statistical significance was analyzed with Student's t-test to compare the means of 2 groups. When 3 or more groups were compared, one- or two-way-ANOVA Tukey's test has been applied (Prism 6; GraphPad software). In the figures, different levels of significance are indicated as: * for $p<0.05$; ** for $p<0.01$; *** for $p<0.001$; **** for $p<0.0001$.

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List of abbreviations

AAV: Adeno-associated viral vector

ANR: Agence nationale de la recherche

bp: base pair

BSA: bovine serum albumin

Bsg: Basigin gene

BSG1: Basigin-1 protein

CMV: Cytomegalovirus

CNRS: Centre national de la recherche scientifique

DNA: Deoxyribonucleic acid

Dnase I: Deoxyribonuclease I

ERG: Electroretinography

FACS: Fluorescence-activated cell sorting

FFB: Foundation fighting blindness

FBS: Fetal Bovine Serum

GFP: Green fluorescent protein

G6P: Glucose-6-phosphate

G6PDH: Glucose-6-phosphate dehydrogenase

GLUT1: glucose transporter SLC2A1

HBSS: Hank's balanced salt solution

HNE: 4-Hydroxynonenal

Ig: Immunoglobulin

Inserm: Institut national de la santé et de la recherche médicale

miRNA: micro-ribonucleic acid

NADPH: Nicotinamide adenine dinucleotide phosphate, reduced form

NADP+: Nicotinamide adenine dinucleotide phosphate, oxidized form

NGS: Normal Goat Serum

Nxn1: Nucleoredoxin-like-1 gene

Nxn2: Nucleoredoxin-like-1 gene

OCT: Optical coherence tomography

OPN1L/MW: Long (red) and medium (green) wavelength opsins

OPN1MW: Medium (green) wavelength opsin

OPN1SW: Short (blue) wavelength opsin

ONL: Outer nuclear layer

PGD: 6-phosphogluconate dehydrogenase

PCR: Polymerase chain reaction

PEP: Phosphoenol pyruvate

PBS: Phosphate buffer saline

PKM: Pyruvate kinase isoform M

PN: Post-natal day

PNA: Peanut agglutinin

PPP: Pentose phosphate pathway

R5P: Ribulose-5-phosphate

rc: recombinated allele

RdCVFL: Rod-derived cone viability factor long

RdCVF: Rod-derived cone viability factor

RFP: Red fluorescent protein

RNA: Ribonucleic acid

TAU: Microtubule-associated protein τ

UPMC: Université Pierre et Marie Curie

vg: viral genome

wt: wild-type

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Figure legends.

Figure 1.

Homologous recombination of the *Nxnl1* gene in the cones of the [Cone-Cre:*Nxnl1cd/cd*] mice. Position of P1/P2 and P3/R4 primers within the *nxnl1* gene (**A**). PCR analysis in FACS-sorted cones of *Nxnl1*+/-, *Nxnl1*-/-, [Cone-Cre:*Nxnl1*+/-] and [Cone-Cre:*Nxnl1cd/cd*] mice, negative control (0). Top row: the amplification of exon1 (P1, P2). Middle row: the amplification by primers (P3, R4) framing exon1. The rod-specific-phosphodiesterase-6b gene (*Pde6b*) is an internal control (**B**). Real-time PCR quantification with primers (P3, R4) in the DNA extracted from mouse retinas (n=3 mice). Cre-mediated recombination took place in about 2.5% of the retinal cells in [Cone-Cre:*Nxnl1cd/cd*] retina as compared to *Nxnl1*-/- mice (n=3 mice) (**C**). Real-time PCR quantification of *Nxnl1* gene in the *Nxnl1*+/-, *Nxnl1*-/-, [Cone-Cre:*Nxnl1*+/-] and [Cone-Cre:*Nxnl1cd/cd*] mice retina with P1/P2 (n=3 mice) (**D**). Quantification of RdCVF cDNA expression (n=3 mice) (**E**). Quantification of RdCVFL cDNA expression (n=5) (**F**). Western blotting analysis of retinal extracts of mice aged 1 month using anti-RdCVF-N antibodies. ACTB: cytoplasmic actin (**G**). a.u.: arbitrary unit. Error bars correspond to standard of the mean (SEM).

Figure 2.

Cone function and viability of cones of the [Cone-Cre:*Nxnl1cd/cd*] mice are reduced at 8 months of age. Representative photopic ERG recordings (**A**). Amplitudes of photopic ERGs (n=10 mice) (**B**). Representative 10 Hz flicker ERG recordings (**C**). Amplitudes of 10 Hz flicker ERGs (n=10 mice) (**D**). Amplitudes of 20 Hz ERG at 8 months (n=10 mice) (**E**). Cone density on flat-mounted retinas (n=8 eye, one per mouse) (**F**). Representative images of 8-month-old mouse retinas labeled with OPN1MW and OPN1SW antibodies for cone density measurements (**G**). Lower panel: 40 fold amplification of the * area of each retina. Scale 1.25 x: 1 mm; 40 x: 25 μ m. ERG photopic or flicker data are the average of both eyes of each mouse. Error bars correspond to SEM: *: p < 0.05; **: p < 0.01 and ***: p < 0.001.

Figure 3.

Cone function and viability of cones of the [Cone-Cre:*Nxnl1cd/cd*] mice are reduced after exposure of young animals to hyperoxia. Representative photopic ERG recordings (**A**). Amplitudes of photopic ERGs (Normoxia: n=8 mice; Hyperoxia: n=18 mice) (**B**). Representative 10 Hz flicker ERG recordings (**C**). Amplitudes of 10 Hz flicker ERGs (Normoxia: n=8 mice; Hyperoxia: n=20 mice) (**D**). Cone density on flat-mounted retinas (Normoxia: n=7 eyes, one per mouse; Hyperoxia: n=12 eyes, one per mouse) (**E**). Representative images of retina labeled with OPN1MW and OPN1SW antibodies for cone density measurements. Retinas of 1 month old mice under 21% normoxia. Retina of 1 month old mice after 1 week 75% hyperoxia treatment. Lower panel: 40 fold amplification of the * area of each retina (**F**). Scale 1.25 x: 1 mm; 40 x: 25 μ m. Normoxia: 21% O₂; hyperoxia: 75% O₂. ERG photopic or flicker data are the average of both eyes of each mouse. Error bars correspond to SEM: *: p < 0.05, **: p < 0.01, and ****: p < 0.0001.

Figure 4.

Oxidative damage in retinal sections. Oxidative stress was induced in retinal cells after 1 week under 75% O₂. Immunostaining of retinal sections of *Nxnl1*+/+, *Nxnl1*−/−, [Cone-Cre:*Nxnl1*+/+] and [Cone-Cre:*Nxnl1cd*/cd] mice at 1 month: DAPI (blue), PNA (green) and 4-hydroxynonenal (HNE, red) (**A**), or acrolein (red) (**B**). Scale bar: 20 μm.

Figure 5.

Cones express the thioredoxin RdCVFL and not the trophic factor RdCVF. Image of a representative portion of the retina injected with AAV2-7m8-OPN1L/MW-GFP (green) and labeled by anti-OPN1MW antibodies (red and green) (**A**). Scale bar, 20 μm. Current responses to voltage steps of a GFP-positive cell by stepping the membrane potential from −100 mV to +40 mV in 10 mV increments (**B**). Current-voltage (I-V) curve of the same cell (**C**). Single cell RT-PCR of negative control (0, first lane), cone (second lane) and positive control (+, third lane) (**D**). RT-PCR analysis of twelve clamped cones from a total of retinas of 6 individual mice (**E**).

Figure 6.

RdCVFL protects cones against oxidative stress. Amplitudes of photopic ERGs (**A**, n=18; **B**, n=11). Representative photopic ERG recordings (**C**). ERG recordings of the corresponding contralateral uninjected eye (**D**). Amplitudes of 10 Hz flicker ERGs (**E**, n=21; **F**, n=8). Representative 10 Hz flicker ERG recordings (**G**). Cone density on flat-mounted retinas (n=8) (**H**). Representative images of retina labeled with PNA for cone density measurements. Lower

panel: 40 fold amplification of the * area of each retina. Scale 1.25X: 1 mm; 40X: 25 μ m (H). n = number of injected eyes, one per mouse. Grey bars: [Cone-Cre:Nxnl1+/+], black bars: [Cone-Cre:Nxnl1/-] mice. Error bars correspond to SEM: *: p < 0.05, **: p < 0.01, ***: p < 0.001 and ****: p < 0.0001.

Figure 7.

Cone-RdCVFL is implicated in cone survival. Luciferase activity after transient co-transfection of 200 ng miRdCVFL plasmid or its scrambled control with the RdCVFL-*Renilla* luciferase reporter plasmid in HEK293 cells (**A**). The number of live chicken cones (n=30) one week after electroporation with scrambled control (200 ng) or miRdCVFL (200 ng and 400 ng) (**B**). Image of the cone-enriched culture cells electroporated with the RFP reporter (**C**). Error bars correspond to standard deviation in (A) and SEM in (B): *: p < 0.05, **: p < 0.01, ***: p < 0.001 and ****: p < 0.0001.

Figure 8.

RdCVFL expression in cones of the *rd10* mice results in their protection. Representative photopic ERG recordings (**A**). ERG recordings of the corresponding contralateral uninjected eye (**B**). Amplitudes of photopic ERGs (n=7) (**C**). Representative 10 Hz flicker ERG recordings (**D**). Amplitudes of 10 Hz flicker ERGs (n=7) (**E**) and those of 20 Hz flicker ERGs (n=7) (**F**) at PN40. Cone density on flat-mounted retinas (n=7) (**G**). Representative images of retina labeled with OPN1MW and OPN1SW antibodies for cone density measurements. Lower panel: 40 fold amplification of the * area of each retina (**H**). Scale 1.25X: 1 mm; 40X: 25 μ m. n = number of injected eyes, one per mouse. wt: C57BL6/N strain. Error bars correspond to SEM: *: p < 0.05.

Figure 9.

Theoretical effects of reactive oxygen species on aerobic glycolysis. RdCVF: rod-derived cone viability factor, RdCVFL: rod-derived cone viability factor long, red, ^{ox}: oxidized, ^{red}: reduced, BSG1: basigin 1, GLUT1: facilitated glucose transporter 1 (SLC2A1), HEX: hexokinase, GPI: glucose phosphate isomerase, PFK: phosphofructokinase, ALDO: aldolase, GAPDH: glyceraldehyde-3-phosphate dehydrogenase, TPI: triose phosphate isomerase, PGK: phosphoglycerate kinase, PGM: phosphoglycerate mutase, ENO: enolase, PKM: pyruvate kinase, LDH: lactate dehydrogenase, MCT1: facilitated lactate transporter 1 (SLC16A1), Glc: glucose, G6P: glucose-6-phosphate, G6PDH: Glucose-6-phosphate dehydrogenase, PGD: 6-phosphogluconate dehydrogenase, R5P: Ribulos-5-phosphate, PPP: pentose phosphate pathway, PEP: phosphoenol pyruvate, NADPH: nicotinamide adenine dinucleotide phosphate, TrxRs: thioredoxin reductases, PYR: pyruvate, Lact: lactate, OXPHOS: oxidative phosphorylation, ROS: reactive oxygen species.

Supplementary Figure 1. Fluorescence activated cell sorting (FACS). Retinal cells from the [Cone-Cre:Nxnl1cd/cd] mouse were labeled with rabbit anti-OPN1MW antibodies (**C, E**) or rabbit anti-IgG (**B, D**), followed with R-phycoerythrin. Two types of cells were detected by FACS (**A**). All type1 cells (**C**) exhibit low fluorescence intensity as the negative control (**B**). In type 2 cells, the 2% of cells that contain higher fluorescent intensity (**E**) than their negative control (**D**) were collected and analyzed.

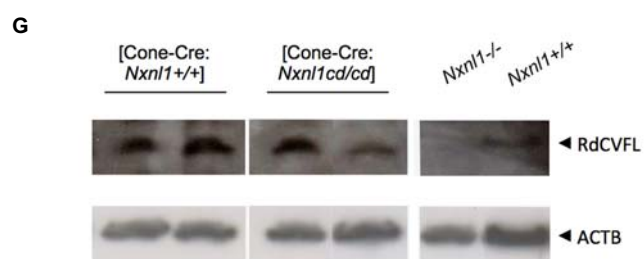
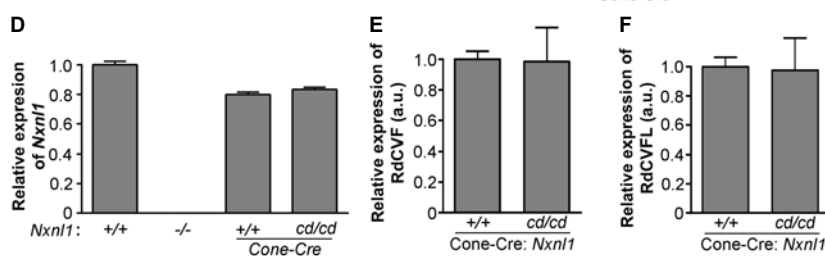
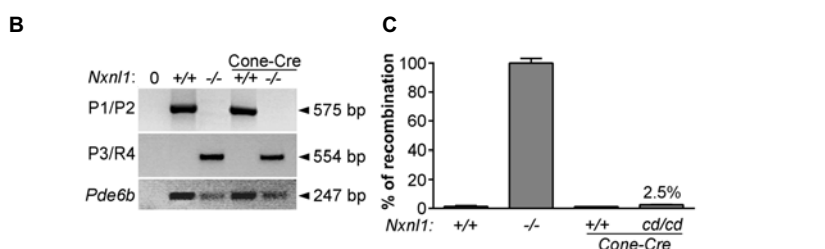
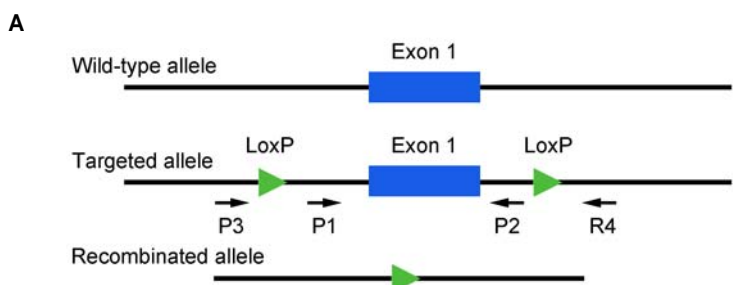
Supplementary Figure 2. Representative 20 Hz ERG recordings at 8 months (**A**). Representation of the b-wave amplitude and the implicit time from ERG photopic traces (**B**). Implicit time of photopic ERGs (n=10 mice) Blue: Cone-Cre-*Nxnl1*+/+, Red: one-Cre-*Nxnl1cd/cd* and green: *Nxnl1*-/- (**C**). Cone density on flat-mounted retinas at 1 month (n=7) 4 months (n=5) and 8 months (n=7). n: eye, one per mouse. Error bars correspond to SD: *: p < 0.05, **: p < 0,01 (**D**). Representative images from flat-mounted retinas at high resolution (x100) of 4-months old mice labeled with a mixture of S- and M-antibodies. Scale bars 60 µm (**E**).

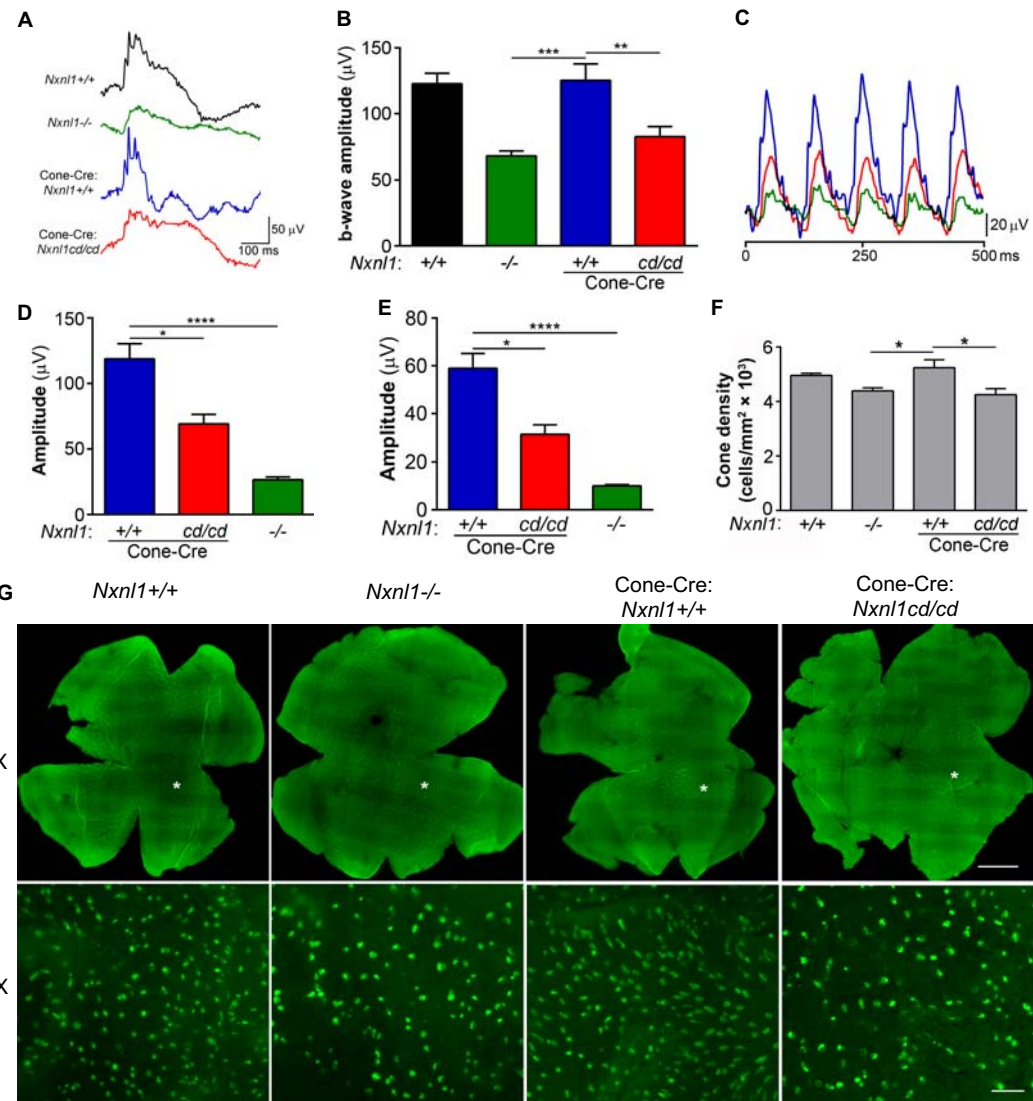
Supplementary Figure 3. Representative 20 Hz ERG recordings at 1 month (**A**). Amplitudes of 20 Hz ERGs at 1 month (normoxia: n=8 mice; hyperoxia: n=20 mice) (**B**). Representative traces of scotopic ERG after 3×10^{-1} cd.s/m² flash intensity at 1 month of age (**C**). Scotopic a-wave amplitudes (**D**, normoxia n=5 mice; hyperoxia +/+, n=6 mice; cd/cd n=8 mice). Scotopic b-wave amplitude. The asterisk indicates that the values are statistically different at 3×10^{-1} cd.s/m² (**E**, normoxia n=5 mice; hyperoxia +/+, n=6 mice; cd/cd n=8 mice). Representative images of the OCT (**F**). Outer nuclear layer thickness along the temporo-nasal axis (**G**, normoxia n=5 mice; hyperoxia +/+, n=7 mice; cd/cd n=9 mice). Outer nuclear layer thickness along the dorso-ventral axis (**H**, normoxia n=5 mice; hyperoxia +/+, n=7 mice; cd/cd n=9 mice). Error bars correspond to SEM: **: p < 0.01, ***: p < 0.001 and ****: p < 0.0001.

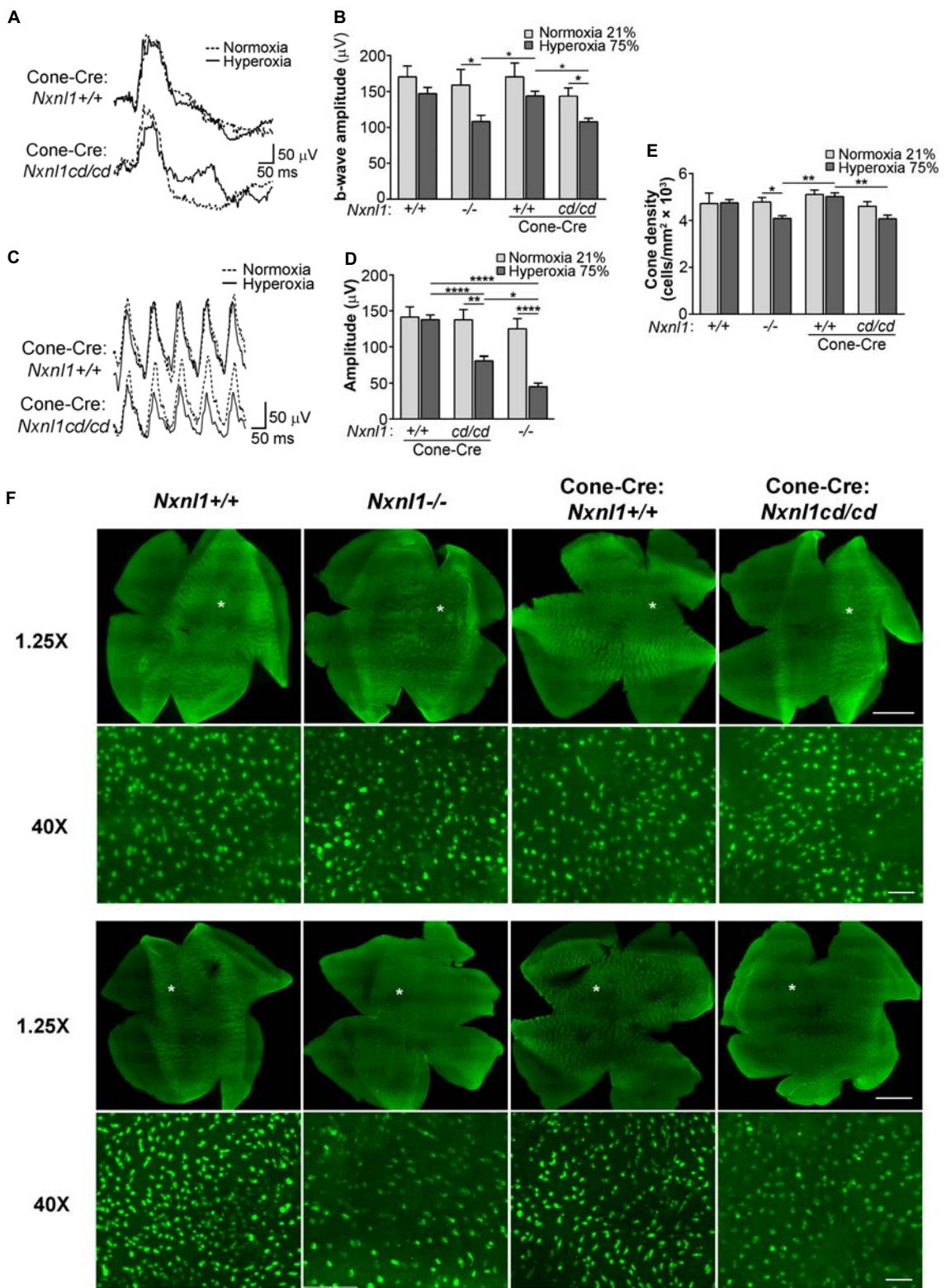
Supplementary Figure 4. Amplitudes of 20 Hz flicker ERGs (**A**, n=20; **B**, n=8) at PN40. Representative traces of 20 Hz flicker ERGs at PN40 (**C**). Implicit time of photopic ERGs (n=8 mice) (**D**). n = number of injected eyes, one per mouse. Correlation between the photopic and 10 Hz flicker ERG. Data plotted for 53 [Cone-Cre:*Nxnl1cd/cd*] measurements of photopic versus flicker 10 Hz ERG are giving a Pearson correlation coefficient of 0.6354. The coefficient is

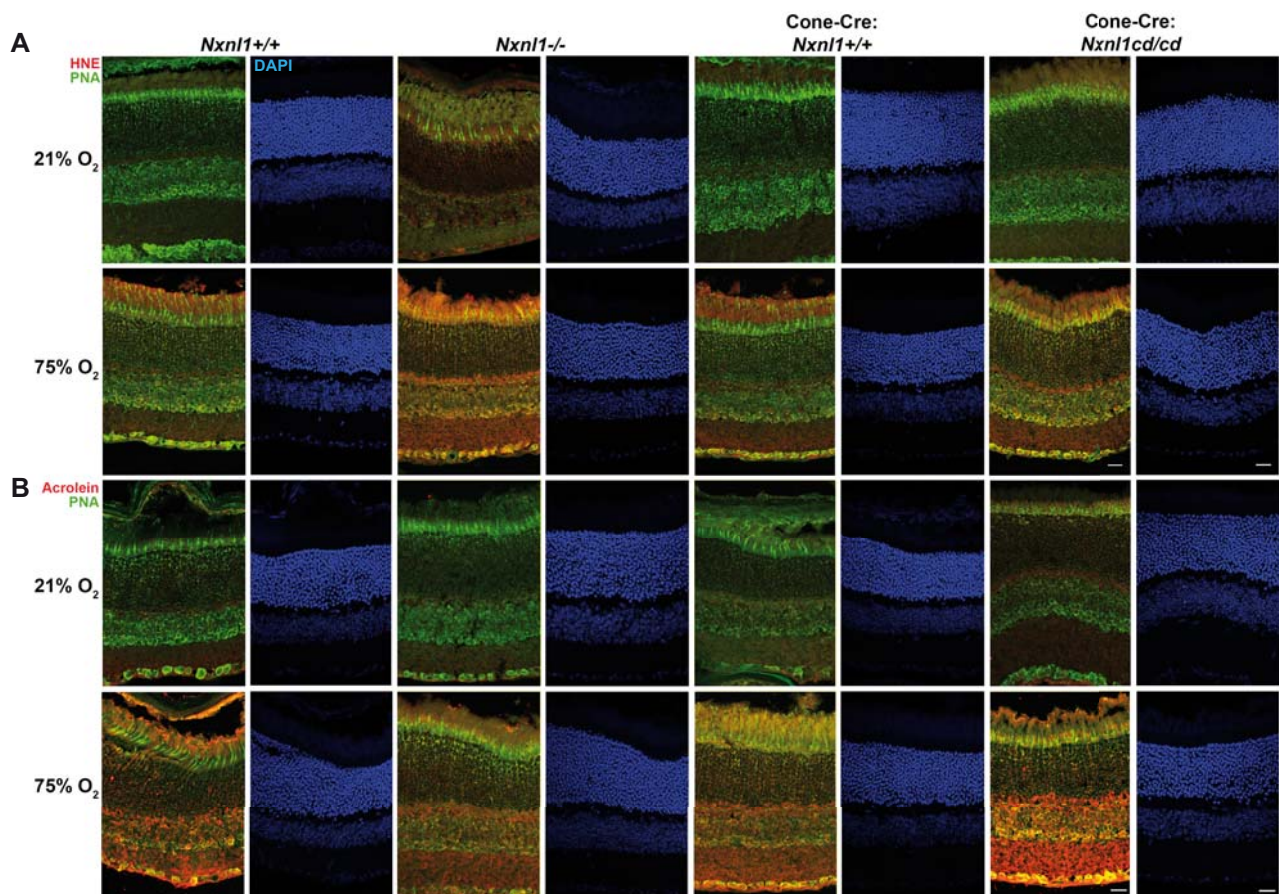
0.5089 when comparing photopic to flicker 20 Hz. n = number of injected eyes, one per mouse. Error bars correspond to SEM: *: p < 0.05, ****: p < 0.0001.

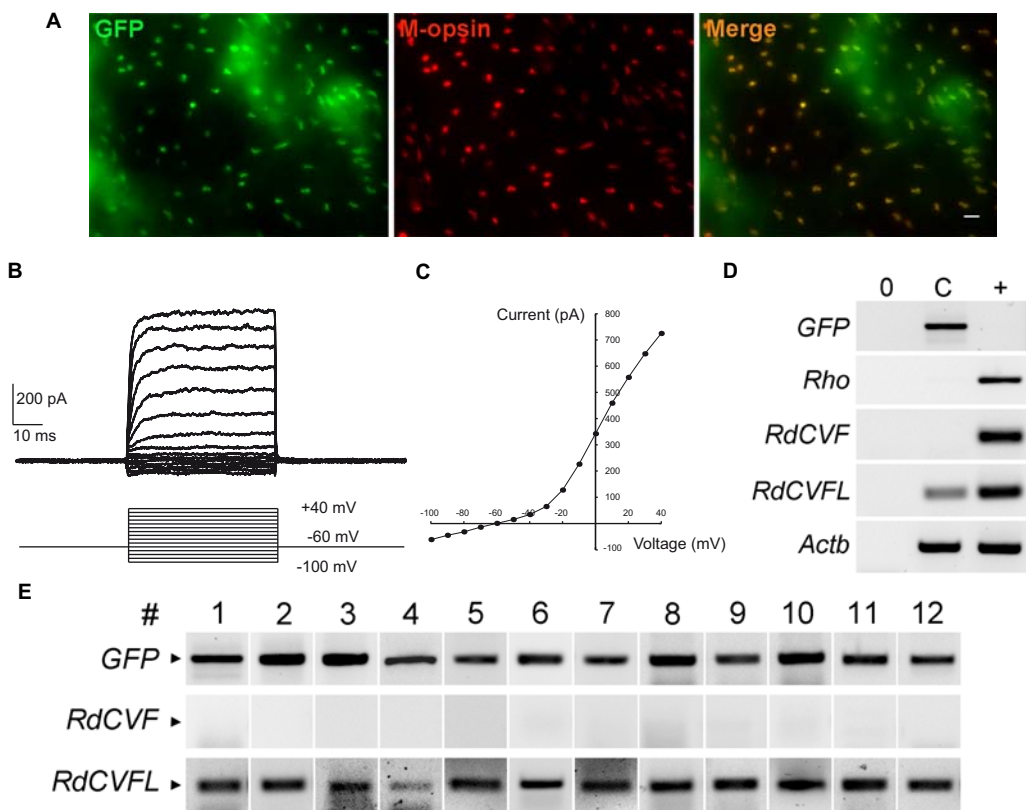
Supplementary Figure 5. Representative traces of 20 Hz flicker ERGs (A). Correlation between the photopic and 10 Hz flicker ERG. Data plotted for 42 *rd10* measurements of photopic versus flicker 10 Hz ERG are giving a Pearson correlation coefficient of 0.503. The outlier in red (113-60) corresponds to a non-injected contralateral eye (B). The coefficient is 0.544 when comparing photopic to flicker 20 Hz.

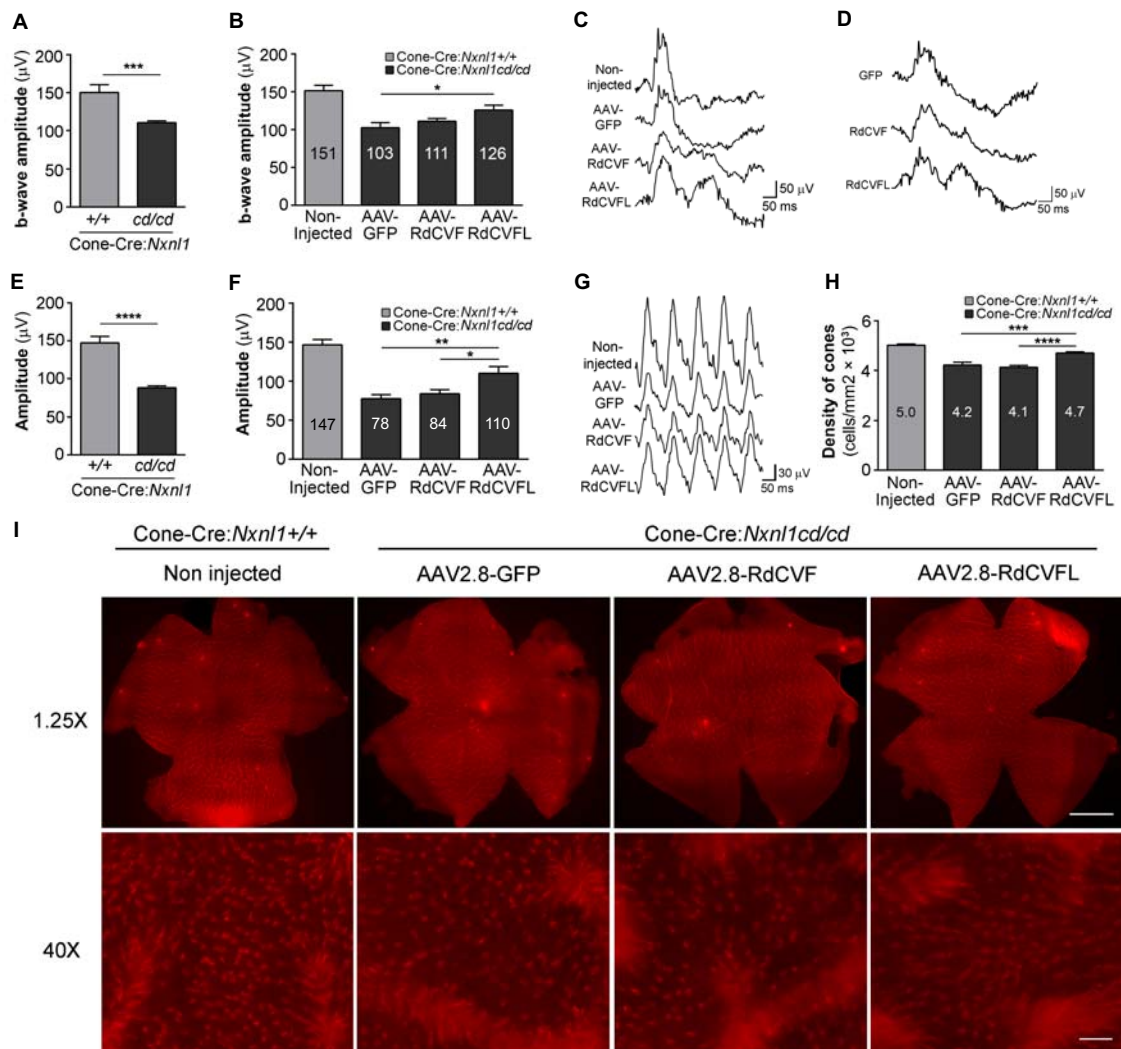


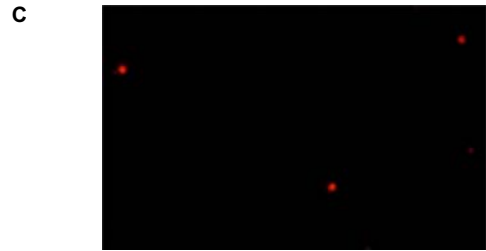
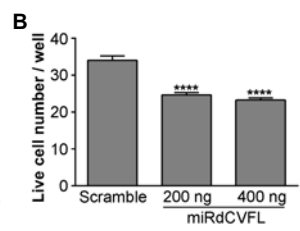
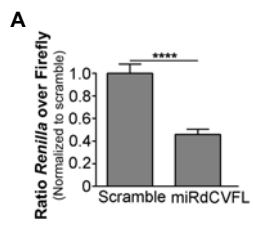


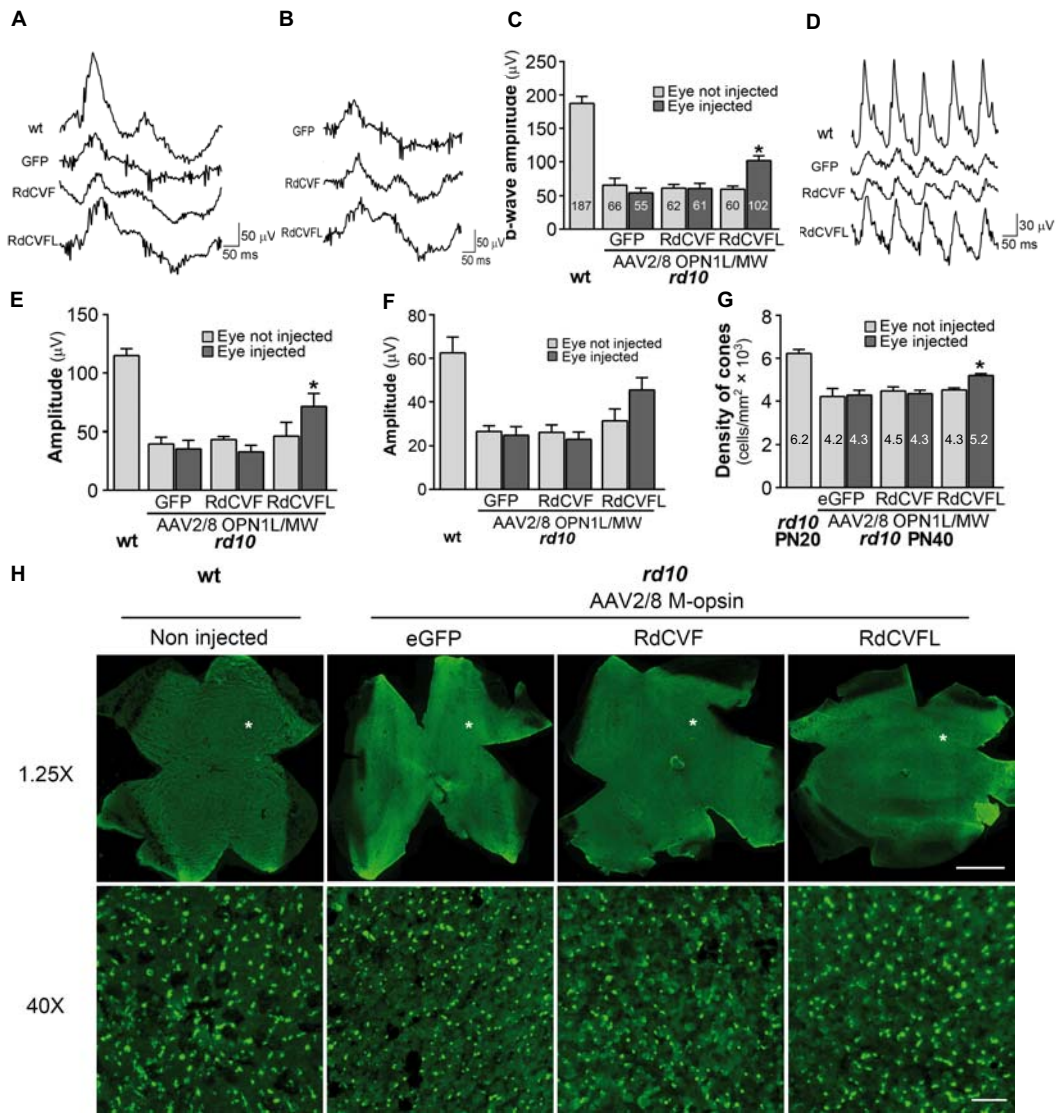


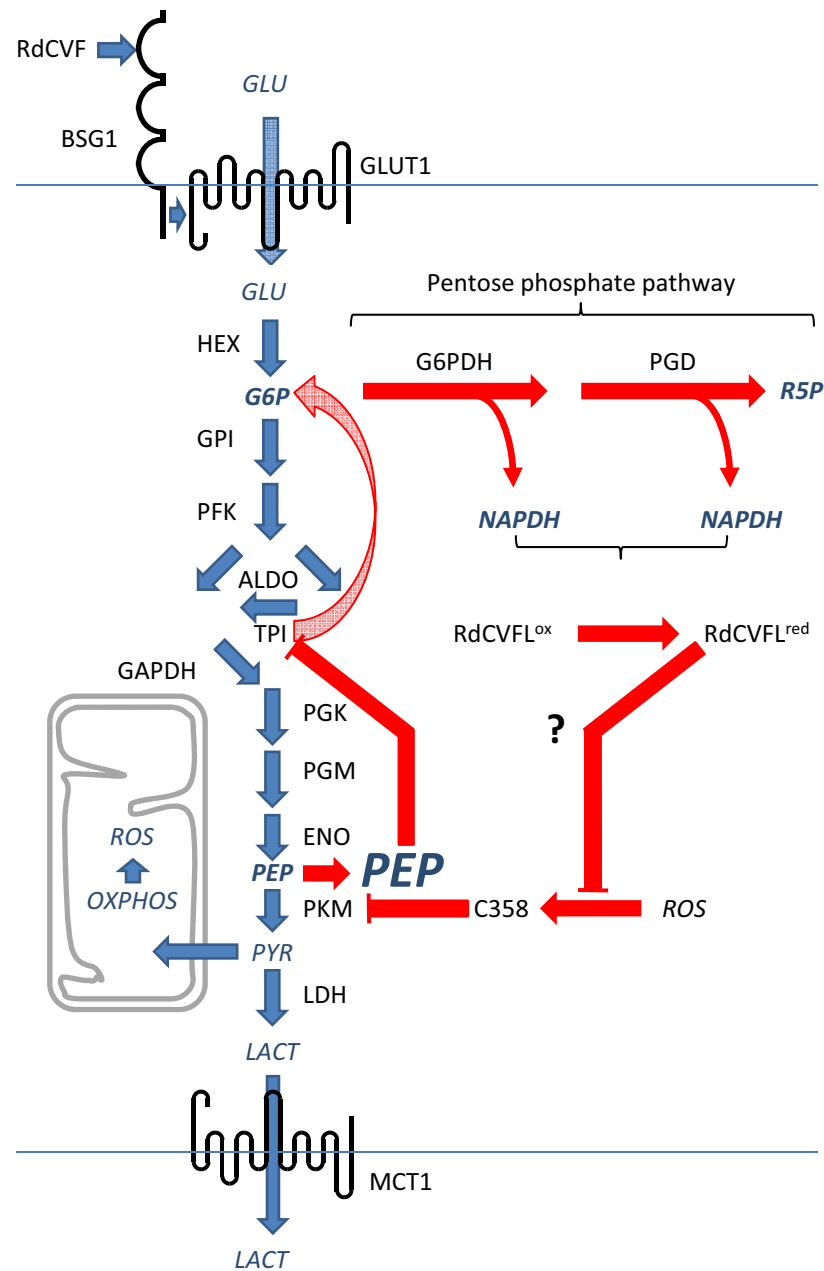


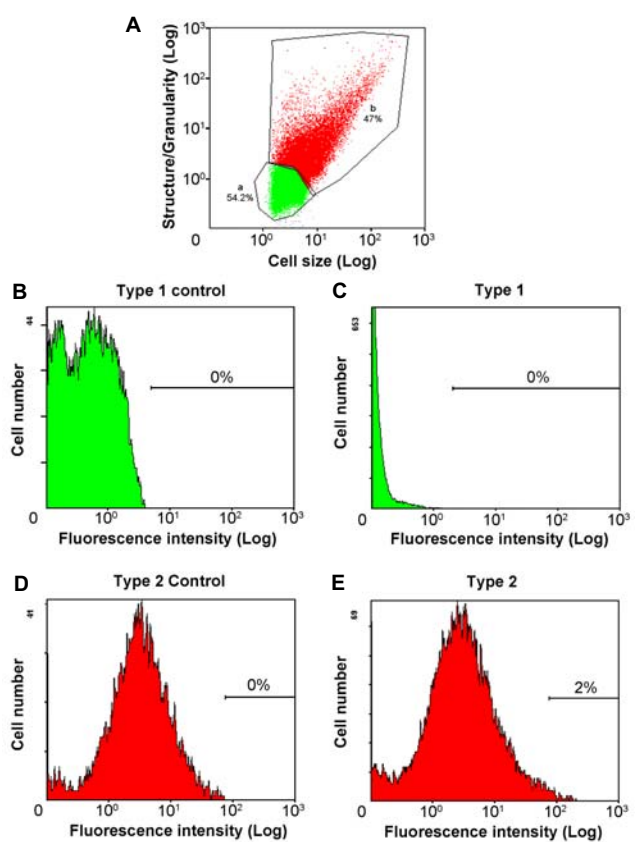


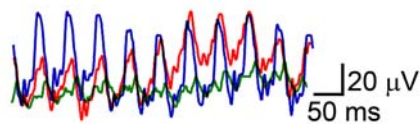
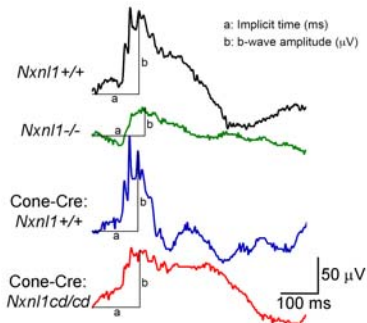
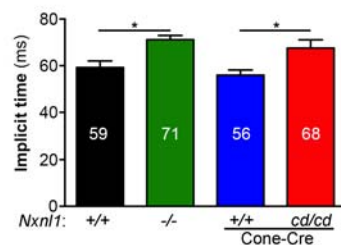
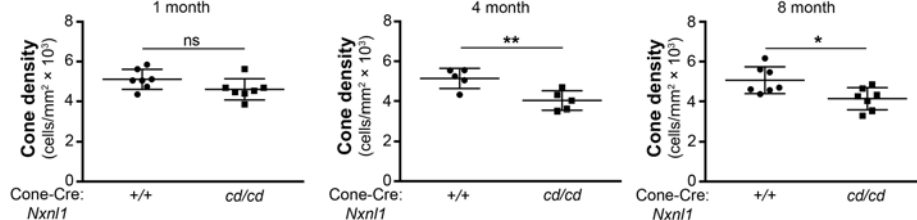










A**B****C****D****E**