Late-Onset Cone Photoreceptor Degeneration Induced by R172W Mutation in Rds and Partial Rescue by Gene Supplementation

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PURPOSE. R172W is a common mutation in the human retinal degeneration slow (RDS) gene, associated with a late-onset dominant macular dystrophy. In this study, the authors characterized a mouse model that closely mimics the human phenotype and tested the feasibility of gene supplementation as a disease treatment strategy.

METHODS. Transgenic mouse lines carrying the R172W mutation were generated. The retinal phenotype associated with this mutation in a low-expressor line (L-R172W) was examined, both structurally (histology with correlative immunohistoch­emistry) and functionally (electroretinography). By examining animals over time and with various rds genetic backgrounds, the authors evaluated the dominance of the defect. To assess the efficacy of gene transfer therapy as a treatment for this defect, a previously characterized transgenic line expressing the normal mouse peripherin/Rds (NMP) was crossed with a higher-expressor Rds line harboring the R172W mutation (H-R172W). Functional, structural, and biochemical analyses were used to assess rescue of the retinal disease phenotype.

RESULTS. In the wild-type (WT) background, L-R172W mice exhibited late-onset (12-month) dominant cone degeneration without any apparent effect on rods. The degeneration was slightly accelerated (9 months) in the rds+/−− background. L-R172W retinas did not form outer segments in the absence of endogenous Rds. With use of the H-R172W line on an rds+/−− background for proof-of-principle genetic supplementation studies, the NMP transgene product rescued rod and cone functional defects and supported outer segment integrity up to 3 months of age, but the rescue effect did not persist in older (11-month) animals.

CONCLUSIONS. The R172W mutation leads to dominant cone degeneration in the mouse model, regardless of the expression level of the transgene. In contrast, effects of the mutation on rods are dose dependent, underscoring the usefulness of the L-R172W line as a faithful model of the human phenotype. This model may prove helpful in future studies on the mechanisms of cone degeneration and for elucidating the different roles of Rds in rods and cones. This study provides evidence that Rds genetic supplementation can be used to partially rescue visual function. Although this strategy is capable of rescuing haploinsufficiency, it does not rescue the long-term degeneration associated with a gain-of-function mutation. (Invest Ophthal­mol Vis Sci. 2007;48:5397–5407) DOI:10.1167/iovs.07-0663

A product of the retinal degeneration slow (RDS) gene, Rds is a tetraspanning membrane protein expressed in the disc rim of rods and cones.1 It has been extensively studied for the past 25 years, and more than 80 disease-causing-mutations in this gene have been identified (http://www.retina-international.com/sci-news/rdsnmut.htm). One of the most common Rds mutations is a substitution of tryptophan for arginine at position 172 (R172W).2–5 This mutation has been described by multiple laboratories and occurs in a large number of families. Affected persons typically have faltering visual acuity in the third to fourth decade of life.2 Children and young adults carrying the mutation have been reported as asymptomatic6 but may exhibit macular changes. Functionally, patients carrying this mutation have normal rod, but diminished cone, electro­retinographic (ERG) amplitudes and elongated implicit times.

Rds was first identified in association with a naturally occurring mouse model called retinal degeneration slow (rds+/−), which does not form outer segments and has no detectable rod or cone ERG response. The hemizygous (rds+/−−) mouse exhibits a haploinsufficiency phenotype associated with decreased rod and cone ERG responses and whorl-like, malformed outer segments.6 Rds is a structural protein present as a homodimer or a heterodimer complex, or as both, with its homologue, Rom-1 (rod outer segment membrane protein-1).7,8 Results of in vitro studies indicate that the large extracellular D2 loop of Rds (containing more than 70% of disease-causing mutations) is the site of interactions to form Rds/Rom-1 complexes,7 whereas the C terminus of Rds protein is responsible for outer segment targeting of the protein.9 Rds is thought to have several hypothesized roles in the outer segment: (1) Rds/Rom-1 complexes function to keep the ‘‘hairpin rim’’ region of the rod outer segment disks intact10–12; (2) Rds/Rom-1 complexes potentially keep the disks connected to the rod outer segment plasma membrane through interactions with GARPs (glutamic acid-rich protein) and the cyclic nucleotide-gated ion channels of the plasma membrane13; (3) the C terminus of Rds directly complex targeting to the outer segment and helps align outer segment disks9; and (4) the C terminus of Rds may mediate disc membrane fusion,14–17 possibly in direct cooperation with other binding partners.18

Recent evidence from our group has suggested that Rds does not function the same way in cone photoreceptors that it does in rod photoreceptors. This observation came from studies of the cone-dominant Nrl/rds double knockout mouse,19 whose phenotype is distinct from that of both the Nrl+/−− (neural retina leucine zipper)20 and the rds+/−− retina. In con-
In our previous work, we reported the bio-
nature, these outer segments are tubular and do not form lamellae. The reasons for the difference in Rds function in cones and rods are not fully understood, but this observation is supported by the clinical findings that RDS mutations can cause either rod-dominant (retinitis pigmentosa) or cone-domin-

Although the R172W mutation occurs within the D2 loop region of Rds, it is not located in the area required for complex assembly. In fact, it is unclear why the R172W mutation causes this distinct cone-associated phenotype. What isclear is the importance of arginine at position 172 to cone structure. Specifically, each of the three known substitutions at position 172 (R172W, R172G, R172Q) causes a cone- or a fovea-domin-

In the present study, we characterized a new R172W model that closely mimics the human retinal phenotype and tested the efficacy of gene transfer therapy as a strategy to combat diseases associated with RDS mutations. As reported previ-

In Western Blot Analysis, Rabbit polyclonal antibody against residues 331 to 346 of murine Rds C terminus (1:1000 dilution) was used for the detection of Rds, as described. Anti-actin antibody (1:250 dilution; Sigma-Aldrich, St. Louis, MO) was used to control for sample loading. Retinal protein extraction and blot analysis were carried out as previously described.

In Immunohistochemistry, Mice were humanely killed, and whole eyes from transgenic and nontransgenic mice were enucleated and fixed for 2 hours in 0.1 M phosphate buffer (pH 7.4) containing 4% formaldehyde, after which the lens was removed and the remaining tissue was placed in phos-

In Light and Electron Microscopy and Quantitative Morphometry, Enucleated eyes were fixed in buffered mixed aldehydes, processed, embedded in plastic resin, and sectioned, as described in detail previ-

Materials and Methods Generation of the Transgene R172W transgenic mice were generated as described previously. Briefly, full-length mouse RDS cDNA was isolated, and the R172W mutation in the D2 loop and P341Q modification in the C terminus was introduced by site-directed mutagenesis. The cDNA was preceded by a 1.3-kb promoter fragment of the human interphotoreceptor retinoid binding protein (hIRBP) and was followed by a 0.7-kb regulatory fragment of the SV40 polyA. Previous studies show that the P341Q modification exerts no biochemical, structural, or functional effect on Rds and that it allows specific recognition of the transgenic protein with the monoclonal 3B6 antibody (generated by Robert Molday, University of British Columbia, Canada). Three transgenic lines are used in the present study: HR172W was characterized previously. L-R172W is characterized here, and the normal mouse peripherin/Rds (NMP) line used in our rescue studies was characterized previously. All L-R172W mice used in this study were on the C57Bl6 background and were homozygous for the transgene regardless of the rds background. Mice were maintained in the breeding colony under cyclic light (14-hour light/10-hour dark) conditions; cage illumination was approximately 7 foot candles during the light cycle. All procedures were approved by the University of Oklahoma Health Science Center Institutional Animal Care and Use Committee (IACUC) and adhered to the ARVO Statement for the Use of Animals in Ophthalmic and Visual Research (http://www.arvo.org/).

Expression of the Transgene Total RNA was extracted from retinas of transgenic mice using reagent (TRizol, Gibco-BRL, Gaithersburg, MD). Five micrograms retinal RNA was separated by electrophoresis on a 0.7% agarose gel containing 18% formaldehyde. The gel was stained with ethidium bromide to check RNA integrity (judged by the integrity of the 28s and 18s rRNA) and was then transferred to nitrocellulose membrane and was hybridized.

Quantitative Morphometry was the same as described. To quantify rod function but no permanent rescue of cone function. Supplementation with WT Rds provides partial, long-term res-

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Light microscopy, sections (0.75–4.0 μm thickness, stained with 1% toluidine blue) were photographed with a photomicroscope (BH-2; Olympus, Tokyo, Japan) in auto-expose mode using 20× (air) or 60× (oil immersion) DPlanApo objectives. For electron microscopy, ultrathin (silver-gold) sections were poststained with lead citrate and uranyl acetate on-grid and were viewed with an electron microscope...
Photoreceptor nuclei were counted in a microscopic field that spanned $100 \mu m$ within the central superior and central inferior portions of the retina, approximately $300 \mu m$ away from the edge of the optic nerve head. At least three individual eyes taken from separate litters were used for examination at each time point.

**Electron Microscopic Immunogold Cytochemistry**

The procedures used were as described in detail.\textsuperscript{25,28} In brief, ultrathin tissue sections (in LR White plastic, on nickel grids) were briefly rinsed in Tris-buffered saline (TBS; 0.15 M NaCl, 0.05 M Tris) and then were "blocked" for 1 hour at room temperature in blocking buffer (TBS containing 1% [wt/vol] RIA-grade BSA and 5% [vol/vol] normal goat serum [NGS]). The sections were exposed overnight at 4°C to anti-cone opsin mixture antibody (1:2500 in blocking buffer). After brief rinses with TBS, sections were incubated for 2 hours at room temperature with goat anti–rabbit IgG conjugated to 15-nm diameter colloidal gold (Amersham, Arlington Heights, IL), diluted 1:50 in blocking buffer. After brief rinses in TBS and distilled water, sections were treated with 1% aqueous glutaraldehyde (5 minutes) and rinsed with TBS. Sections were then stained with 2% aqueous uranyl acetate (15 minutes) and lead citrate (6 minutes), rinsed again with distilled water, air dried, and exposed to OsO$_4$ vapors for 30 minutes before viewing with an electron microscope.

**Electroretinography**

Electroretinograms were recorded from nontransgenic and transgenic mice ($n = 6$–$10$ each) at different developmental stages to obtain dark-adapted, light-adapted, and specific cone responses, as described.\textsuperscript{19,29} For spectral electroretinography, a two-flash technique was used to record the responses of blue and red/green cones. The first flash was used to desensitize the rod response; after a 2-second interval, the second flash was presented through a specific filter (400 nm for blue cone and 500 nm for red/green cone) to the retina. In this study, the flash intensity was held constant at $3.1 \log \text{cd} \cdot \text{s/m}^2$, and the interflash interval was kept at 2 seconds. Mice were not dark adapted before recording. Each response represented the average of 10 stimulating flashes at the same intensity and filter setting. At least two responses were recorded for each filter to make sure the ERG system was stable. Statistical significance was determined by Student’s $t$-test (two samples) or ANOVA with Bonferroni post hoc test (more than two samples). ERG amplitudes are presented as averages $\pm$ SEM.

**RESULTS**

**Transgene Generation and Protein Expression**

Several transgenic lines were generated and characterized; two have been selected for extensive study. The first was the...
H-R172W line, which expressed the transgene at approximately 75% of the WT level and showed a severe, early-onset, dominant cone-rod dystrophy. Here we describe the phenotype in an additional line (L-R172W) that expressed the transgene at approximately 40% of the WT level.

The first step in characterizing the L-R172W line was to confirm the expression of the transgenic protein. Retinas from L-R172W mice were taken at postnatal day (P) 30 and used for Northern and Western blot analyses to document the presence of the R172W transcript and protein, respectively. As shown in Figure 1, the R172W transcript (Fig. 1A) and protein (Fig. 1B) were present in retinas from transgenic mice of multiple genetic backgrounds. Based on Western blot analysis of nonreducing gels, R172W protein in L-R172W animals was capable of forming dimers and higher-order complexes, even in the absence of native Rds. Although Rom-1 was nearly undetectable in the rds−/− retina, the presence of R172W protein led to a significant increase in the level of Rom-1 (Fig. 1B, right). Based on previously published quantitative analysis, animals homozygous for the L-R172W transgene on the rds null background expressed the R172W Rds protein at approximately 40% of the endogenous Rds protein level. To confirm that the IRBP promoter was capable of driving transgene expression in rods and cones, immunohistochemical labeling of R172W protein from L-R172W mice was undertaken. As shown in Figure 1C, R172W protein was found in rods and cones (labeled for S-opsin).

**Figure 2.** Evaluation of L-R172W retinal structure and function in the WT background. (A) Histologic and morphometric analyses indicate that, at 12 months of age, L-R172W does not affect rod structure. Scale bar, 20 μm. (B) Serial scotopic ERG analyses at 6 months (6M) and 12 months (12M) show that rod function and sensitivity are not significantly altered by the presence of L-R172W. Dashed lines (+/+ ) indicate the upper and lower limits of the normal range (based on SD) of WT responses. (C) Immunohistochemical analysis (using S-cone opsin antibody) indicates that presence of L-R172W leads to a statistically significant decrease in the number of blue cones. (D) Spectral ERG with 400-nm light (S-cones) and 500-nm light (L/M cones) was conducted on L-R172W and age-matched WT animals. L-R172W animals exhibited a late-onset cone defect (*P < 0.05). Shaded column: WT. Open columns: L-R172W. n = 5 to 7 animals per time point and genotype.
FIGURE 3. Evaluation of L-R172W retinal structure and function in the rds+/− background. Because the L-R172W/rds+/− is the model most representative of the human condition in terms of protein levels, we analyzed retinal structure and function in this model. (A) Histologic analysis of 12-month-old R172W/rds+/− retinas shows moderate structural rescue of the rods compared with rds+/−. Scale bar, 20 μm. (B) ERG amplitudes recorded with 400-nm light (S-cones) and 500-nm light (L/M-cones) indicate that L-R172W/rds+/− is associated with statistically significant, dominant, late-onset cone degeneration (both S- and L/M-cones). This degeneration occurs sooner in the hemizygous background than in the WT (see Fig. 2). Shaded columns: hemizygous. Open columns: L-R172W. *P < 0.05 for differences between rds+/− and L-R172W/rds+/−.

L-R172W Mice Exhibit a Dominant, Late-Onset Cone Dystrophy in the Wild-Type Background

Because the human R172W phenotype was dominant, our next step was to examine whether the L-R172W model was associated with a dominant effect on retinal structure or function. L-R172W eyes at 12 months of age were examined histologically and morphometrically. As shown in Figure 2A, at 12 months of age, the R172W protein had no apparent structural effect on the retina; this was confirmed by morphometric analysis. Next, electroretinograms from dark-adapted animals were used to examine rod function. Consistent with the lack of a structural effect on the rods, ERG analysis (Fig. 2B) indicated that, at 6 months of age, L-R172W mice exhibited scotopic a-wave amplitudes well within the normal range for rod function. Although at 12 months of age the values decreased, they did not diverge statistically from the WT range.

To test the possibility that cone structure and function were adversely affected in L-R172W animals, we used immunohistochemistry to quantify cone nuclei at 12 months and light-adapted spectral electoretinography to examine function at multiple time points. There was a statistically significant (25%) decrease in the number of blue (S) cones found in the retinas of 12-month-old L-R172W animals (Fig. 2C). A similar decrease was observed for red/green (L/M) cones (not shown). This deficit corresponded with a significant decrease in the function of blue cones (approximately 25%) and red/green cones (approximately 20%), as measured by spectral electroretinograms (Fig. 2D), even though these mice have two native Rds alleles. This functional alteration was only observed in 12-month-old animals (the latest age tested).

Cone Degeneration in L-R172W Is Accelerated in the rds+/− Background

Because humans affected by the R172W mutation have one WT and one mutant allele, we studied retinal structure and function of L-R172W animals in the rds hemizygous background. As shown in Figure 3A, in some L-R172W/rds+/− mice, retinal structure at 12 months of age was slightly improved when compared with rds+/− (6–7 vs. 3–4 rows of outer nuclear layer (ONL) nuclei, respectively, in the example shown). However, detailed morphometric analysis indicated no general improvement—that is, L-R172W/rds+/− mice, as a group, did not have more photoreceptor nuclei (per 100-μm area) than nontransgenic littersmates at 12 months of age (L-R172W/rds+/− 124 ± 12 nuclei/100 μm vs. rds+/− 129 ± 15 nuclei/100 μm; n = 6). To test the effect of the R172W protein on rod function in the rds+/− mice, we examined scotopic ERG amplitudes. At early time points (2 months), the presence of the L-R172W protein seemed beneficial to rod function: L-R172W/rds+/− mice had significantly larger scotopic a-wave amplitudes than age-matched rds+/− controls (221.4 ± 47.4 μV vs. 149.0 ± 17.0 μV, respectively; P < 0.05). Consistent with observations of human populations, L-R172W mice in the hemizygous background did not exhibit rod-dominant degeneration. Long-term rod degeneration in L-R172W/rds+/− mice was consistent with the standard haploinsufficiency phenotype seen in rds−/− mouse retinas.

On examination of cone function, it was determined that the late-onset defect we observed in the WT background (Fig. 2D) was accelerated in L-R172W animals hemizygous for rds (Fig. 3B). These animals exhibited approximately 20% decreases in blue and red/green cone ERG function at 9 months compared with rds+/− mice. Compared with WT mice, the decrease in cone ERG amplitudes was even more pronounced; blue cone function was decreased by approximately 45% compared with age-matched WT animals, even though the quantity of Rds protein was approximately equivalent between the two groups (two L-R172W alleles plus one native Rds allele approximately equivalent two native Rds alleles).
R172W Rds Does Not Promote Outer Segment Formation

Given our observations that L-R172W can improve rod structure in the hemizygous background (albeit without permanent functional rescue) and that L-R172W stabilizes Rom-1 expression in the absence of native Rds, we proceeded to test the hypothesis that the L-R172W protein could support the formation of photoreceptor outer segments in the \( \text{rds}^{-/-} \) (null) background. A structural and functional developmental study was undertaken; eyes were collected from L-R172W/\( \text{rds}^{-/-} \) and \( \text{rds}^{-/-} \) mice at P10, P20, and P30. The eyes were examined histologically (Fig. 4), but, in spite of its biochemical similarity to native Rds in terms of complex formation with itself and Rom-1 (as shown in Fig. 1 and Ding et al.25), the amount of R172W protein expressed in L-R172W animals, even dysmorphic ones, was not capable of supporting the formation of outer segments during ocular development. Even though they have similar quantities of Rds protein, the phenotype in L-R172W/\( \text{rds}^{-/-} \) mice (no outer segments) was significantly worse than in \( \text{rds}^{+/+} \) mice (retention of dysmorphic outer segments even at 1 year of age; see Fig. 3A). Finally, ERG analysis indicated that L-R172W/\( \text{rds}^{-/-} \) animals did not have more function than \( \text{rds}^{-/-} \) mice (not shown).

Genetic Supplementation of Wild-Type Rds to Rescue the Phenotype in the R172W Retina

Our next step was to test the hypothesis that gene transfer would be a suitable strategy for treating the R172W disease phenotype. For these proof-of-principle experiments, we chose to use the higher-expresser line, H-R172W, because the severe phenotype can be detected at earlier time points than the more clinically relevant phenotype in the lower-expresser line (L-R172W). Genetic supplementation was our chosen gene transfer method; we used the Rds overexpressing transgenic mouse line known as NMP (normal mouse peripherin/Rds). This mouse line has been fully characterized elsewhere.26 Briefly, the NMP transgene is expressed in rods and cones, is not toxic to the photoreceptor, and has been used effectively to rescue the \( \text{rds}^{+/-} \) haploinsufficiency phenotype and the loss-of-function model expressing the C214S mutation in Rds. Mice homozygous for the transgene express NMP protein at approximately 60% of WT levels.26 Heterozygotes of single (H-R172W/\( \text{rds}^{+/-} \)) and double (H-R172W/\( \text{rds}^{+/-} / \text{NMP}^{+/-} \)) transgenic mice were generated on an \( \text{rds}^{+/-} \) genetic background, and structural and functional rescue of the H-R172W disease phenotype by NMP were assessed at multiple time points.
Light microscopy (Fig. 5A) indicated that, when the animals were 3 months of age, supplementation with NMP led to grossly normal retinal morphology and rescue of ONL thickness (in the H-R172W/rds/H11001/H11002 background). Correlative electron microscopy indicated that supplementation with NMP could restore normal rod outer segment (ROS) structures. However, this structural rescue was not sustained when the animals were 11 months of age (data not shown). As previously reported, the NMP/H11001/H11002/rds/H11001/H11002 retinas (shown with WT as controls in Fig. 5B) exhibited infrequent abnormalities in ROS ultrastructure as a result of relative Rds underexpression (approximately 80% of WT) but showed normal ONL thickness.

To examine the ability of NMP to rescue the structural defect seen in cones, retinas from 3-month-old mice were immunogold labeled with antibodies to S-cone opsin. Figure 6 shows representative images of immunogold-positive labeling in cone, but not rod, photoreceptor outer segments. As shown in Figure 6A, supplementation with NMP led to a remarkable rescue of cone outer segment (COS) structural defects associated with the dominant effect of the R172W mutation. Double-transgenic (R172W+/+/NMP+/+/rds+/+) retinas generated compact, well-aligned, left-eye disks, and, though not fully returned to the morphology seen in WT retinas (Fig. 6B), they were significantly improved when compared with the retinal morphology observed in single-transgenic mice (R172W+/+/rds+/+). In marked contrast to the relatively normal H-R172W ROS structure (Fig. 5A), the devastating effect of R172W on cones is evident from the severely degenerated COS structure (Fig. 6A) in these animals.

To determine the functional relevance of NMP-mediated structural rescue of H-R172W+/+/rds+/+ mice, ERG studies were undertaken. Figure 7A shows representative scotopic and photopic ERG wave forms from single- and double-transgenic mice taken at 1 month of age. As reported previously, the introduction of the H-R172W transgene onto an rds/H11001/H11002 background (H-R172W+/+/rds+/+) alleviates rod functional defects associated with the haploinsufficiency phenotype as early as 1 month of age, whereas expression of the R172W mutant protein causes a significant (P < 0.05) reduction in cone function (Fig. 7B). This early-onset phenotype makes H-R172W a perfect model for our rescue studies. In H-R172W mice, the enhancement in rod function persisted over time, whereas cone function continued to decline in an age-dependent manner. As shown in Figure 7B, the addition of one NMP allele significantly improved rod and cone ERG amplitudes; at 1 month of age, NMP/H-R172W double-transgenic mice had scotopic and photopic amplitudes (90%–91% of WT), significantly higher than H-R172W single-transgenic mice (71% and 50% of WT, respectively). Rescue continued in the short term; at 3 months of age, H-R172W animals carrying an NMP allele had scotopic and photopic ERG amplitudes 82% and 73% of WT levels, respectively, whereas H-R172W animals without...
NMP continued to degenerate (56% vs. 34% of WT responses, respectively). At later time points (11 months), we found double-transgenic (H-R172W+/−/NMP+/−/rds−/−) animals continued to have significantly (P < 0.05) better rod function than single-transgenic (R172W+/−/rds−/−) mice (Fig. 7C, left panel). Although Rds supplementation showed a statistically significant (P < 0.001) amelioration of the cone defect in H-R172W+/−/rds−/− mice at 1 and 3 months of age (Fig. 7B), this rescue did not persist over the long-term, and cone ERG amplitudes in the H-R172W+/−/NMP+/−/rds−/− were not different from H-R172W+/−/rds−/− by the age of 11 months (Fig. 7C, right panel).

**DISCUSSION**

Here we present the phenotype of a transgenic line (L-R172W) that expresses the R172W mutation in Rds at approximately 40% of the WT level. We first confirmed the presence of the R172W transcript and protein level in this line. Functionally, L-R172W mice in the WT background did not exhibit rod-dominant degeneration at either 6 or 12 months compared with WT controls, an observation supported by a lack of morphometric change in the ONL. Conversely, cone function and numbers were significantly reduced in mice carrying the R172W allele. In the rds−/− background, L-R172W cone degeneration was accelerated (compared with the WT background), but again no dominant effect was observed in rods. Furthermore, the presence of the R172W protein could not rescue the rds null phenotype during development and failed to support the formation of photoreceptor outer segments. The L-R172W mouse exhibited a phenotype remarkably similar to that seen in patients carrying the R172W mutation. In the second part of our study, we used the H-R172W line to determine whether genetic supplementation of WT Rds (NMP) would lead to functional or structural rescue of the early-onset dominant phenotype. We observed significant structural and functional rescue of rods and cones at early time points (1–3 months), but at a later time point (11 months), cone rescue was not sustained and rod rescue was only partially sustained.

Our early observation of the R172W protein as incapable of providing significant structural rescue to the rds−/− retina merits further discussion. In our previous work,25 we reported that at early time points, the R172W protein from the H-R172W model was capable of mediating significant structural rescue of ROS in the rds−/− background. H-R172W+/−/rds−/− retinas have outer segments that look similar to those in the rds−/− mice. This partial rescue could be interpreted as a positive function of the R172W allele in rods; however, because so much R172W protein is present in that mouse line, the adverse effects of the R172W mutation were seen much earlier, and the cone degeneration was correspondingly more severe. Thus, we may say that the effects of R172W are clearly dose dependent: at low doses (L-R172W), the dominant cone defect appears later in life (9–12 months) and is moderate (approximately 20% reduction in cone function); however, at high doses (H-R172W), the dominant defect appears very early (1 month) and is devastating (approximately 75% reduction in cone function25). These trends persist regardless of the rds background and clearly highlight the importance of a physiologically relevant model for studying disease pathogenesis. The H-R172W line was able to provide us with invaluable biochemical information, but the clinically more relevant phenotype of the lower-expresser line (L-R172W) leads to more accurate interpretations of disease progression and mechanism.

In this study, the functional phenotype of the L-R172W mouse model closely resembles that seen in human patients harboring this same RDS mutation. The R172W mutation in humans causes a late-onset disease characterized by visual deficits and a decrease in cone function. Although humans affected by this mutation typically do not exhibit any statistically significant decrease in rod function, there is evidence that their rod function tends to be toward the lower limits of the normal range (Fishman G, personal communication, 1999), similar to that observed in our mouse model. Our morphometric observations indicating a decrease in the number of cone nuclei is a first step toward understanding the mechanism of disease progression in patients with the R172W mutation. The fact that the observation is limited to cones further supports
FIGURE 7. Rescue of rod and cone ERG function in H-R172W transgenic mice. (A) Representative rod and cone ERG wave forms at 1 month of age illustrate the improvement in rod and rescue in cone function ($P < 0.001$) in double-transgenic (H-R172W+/+/NMP+/+/rds+/+) compared with single-transgenic (H-R172W+/+/rds+/+) mice. (B) Rod a-wave and cone b-wave averages show a significant enhancement in rod function ($P < 0.05$) and rescue in cone function ($P < 0.001$) in double-transgenic (H-R172W+/+/NMP+/+/rds+/+) compared with single-transgenic (H-R172W+/+/rds+/+) mice at 1 month of age. At 3 months of age, enhancement in rod function and significant ($P < 0.001$) improvement in cone function resulting from the supplementation of WT Rds in H-R172W transgenic mice (H-R172W+/+/NMP+/+/rds+/+) persisted. (C) Rod a-wave and cone b-wave averages at 11 months of age in H-R172W mice demonstrate a significant ($P < 0.001$) rescue in rod function in double-transgenic (H-R172W+/+/NMP+/+/rds+/+) compared with single-transgenic (H-R172W+/+/rds+/+) mice. However, at this age, double-transgenic mice lost their rescue of cone function provided by NMP supplementation. ERG wave amplitudes for (B) represent an average of 12 to 16 eyes for each genotype, and ERG amplitudes for (D) represent 8 eyes for each genotype.
the hypothesis that Rds functions differently in rods versus cones.

Here we observed that cone degeneration occurred in older animals in the WT background but was accelerated in the rds hemizygous background. This is probably a result of the dominant effect of the R172W protein combined with the moderate effect of the Rds haploinsufficiency phenotype. Although we observed no dominant effect of the L-R172W on rod function, L-R172W/rds+/- animals did exhibit the standard rds+/- haploinsufficiency phenotype. In this one aspect, the L-R172W/ rds+/- diverged from the human phenotype. R172W patients did not exhibit clinical signs of Rds haploinsufficiency; rather, they had pure cone degeneration. This disparity might have occurred because our L-R172W mouse model had slightly less R172W protein than what would be afforded by one WT allele (approximately 40% of total Rds in the mouse vs. 50% in the human). However, this disparity likely resulted from species-specific differences between mice and humans; mice exhibit a decidedly lower tolerance for Rds deficiency than do humans. For example, when examined on an rds+/- background, a typical Rds loss-of-function mutation, such as C214S, leads to an early-onset haploinsufficiency phenotype indistinguishable from that seen in rds+/- mice.30 Conversely, patients with the C214S mutation exhibit rod defects (e.g., night blindness), but not until later in life (fourth to fifth decade).51 Far from being a detriment to our models, the differential requirements of human rods versus mouse rods for Rds is further proof of the underlying differences in the role of Rds in rods and cones. In other words, human rods need less Rds than do mouse rods, whereas human cones and mouse cones need similar quantities of Rds (as evidenced by the similarity in cone-dominant disease patterns in humans and mouse models).

The second portion of our study dealt with characterizing the ability of wild-type Rds (NMP) to rescue the severe cone-rod degeneration seen in the H-R172W model and provided us with valuable information regarding the usefulness of gene transfer therapy to rescue dominant disease phenotypes. In the simplest case, one might hypothesize that genetic supplementation would have no effect on a dominant mutation. However, our data do not support this theory. At early time points with H-R172W, rod degeneration is not dominant. In other words, in terms of rod function, having one allele of H-R172W is better than not having it at all. This phenotype is best explained by hypothesizing that at early time points, R172W, if enough is present, it can support rod function, though not as well as native Rds. Thus, the observation that NMP can rescue rod function at early time points is not unexpected. In fact, the H-R172W+/NMP+/-/rds+/- rod function looks strikingly similar to the homozygous L-R172W+/+ phenotype: rod function at the lower end of the normal WT range. What is surprising is that the severe, early-onset, dominant-rod degeneration observed in the H-R172W+/rds+/- can be rescued almost completely. For example, at 1 month of age, the H-R172W cone degeneration is a completely dominant effect (not associated with haploinsufficiency), yet the presence of one allele of NMP provides considerable rescue of cone function (approximately 90% of WT). This observation is exciting from a mechanistic point of view and begs the question: how does NMP overcome the dominant structural and functional effect of H-R172W in cones?

From a gene therapy perspective, however, our gene transfer strategy is less promising. For a gene therapy to be clinically viable, rescue must be long term. Our data clearly show that over the long term, the presence of WT Rds does not provide rescue to cone function. That rescue persists to a certain level in rods is of little clinical usefulness because rod degeneration is not associated with the R172W phenotype in humans. Even with the persistent partial rescue in rods combined with observations at later time points, H-R172W/rds+/- has equal or better rod function than rds+/- (see Fig. 7 and our previous work55), which supports our contention that the R172W mutation does not have a dominant effect on rods in the traditional sense.

Interestingly, however, evidence suggests that the R172W allele may have a small, virtually undetectable dominant effect on the rod. First, the long-term rod ERG in L-R172W animals and R172W patients is at the far lower end of the normal range, though the divergence from normal is not statistically significant. Second, though NMP-mediated rescue of rods in the H-R172W line is certainly a more efficient rescue than of cones over time, this rescue also declines (Figs. 7C-D), in marked contrast to the rescuing ability of NMP in both the rds+/- and in the C214S/rds+/- loss-of-function mutation in which rescue persists at similar levels throughout the life of the animal.52 All these observations suggest that, though gene transfer therapy of the WT Rds may not be a suitable strategy for R172W patients, understanding the reasons for the differential requirement of rods and cones for Rds and the subtle differences in the effect of the R172W mutation in rods and cones is critical for the design of any form of effective treatment. Future rescue studies of the R172W model will likely make use of gene replacement with concurrent gene knockdown approaches. Studies on other dominant retinal degenerations (e.g., the P23H rhodopsin mutation) have used ribozyme53,54 or small interfering or small hairpin RNA (siRNA/shRNA) approaches as a way to decrease expression of a dominant mutant protein.55-57 Similar approaches could be used to overcome gain-of-function mutations in Rds. Because the dose of normal and mutant Rds is critical to disease phenotype and rescue, treatments will have to be fine-tuned to deliver the appropriate amount of therapeutics.

Here we have characterized a mouse model of cone dystrophy associated with the R172W mutation (L-R172W), which closely mimics the late-onset cone dystrophy seen in human patients harboring this mutation. Although gene replacement therapy alone is not likely to be an effective treatment strategy for R172W-associated macular degeneration, the temporary cone rescue we observed after supplementation with NMP raises exciting questions about the mechanism of the R172W-mediated degeneration. Future studies with this model in native and cone-dominant (Nrl-/-) retinas may enable us to gain greater understanding of the pathogenesis and intricacies of dominant cone dystrophies and may help us to elucidate some of the underlying reasons for differences in rod/cone biogenesis and degeneration.

Acknowledgments

The authors thank Chibo Li for generating the transgenic mice used in this study, Huijun Yang and Barbara Nagel for their exceptional technical assistance with histology, and Mark Ballard for his assistance with the manuscript.

References


