Iron-Chelating Drugs Enhance Cone Photoreceptor Survival in a Mouse Model of Retinitis Pigmentosa

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PURPOSE. Retinitis pigmentosa (RP) is a group of hereditary retinal degeneration in which mutations commonly result in the initial phase of rod cell death followed by gradual cone cell death. The mechanisms by which the mutations lead to photoreceptor cell death in RP have not been clearly elucidated. There is currently no effective treatment for RP. The purpose of this work was to explore iron chelation therapy for improving cone survival and function in the rd10 mouse model of RP.

METHODS. Two iron-chelating drugs, 5(4-(2-hydroxyethyl) piperazin-1-yl (methyl)-8-hydroxyquinoline (VK28) and its chimeric derivative 5(3(N-methyl-N-propargyaminomethyl)-quinoline-8-oldihydrochloride (VAR10303), were injected intraperitoneally to rd10 mice every other day starting from postnatal day 14. We investigate the effects of the two compounds on cone rescue at three time points, using a combination of immunocytochemistry, RT-PCR, Western blot analysis, and a series of visual function tests.

RESULTS. VK28 and VAR10303 treatments partially rescued cones, and significantly improved visual function in rd10 mice. Moreover, we showed that the neuroprotective effects of VK28 and VAR10303 were correlated to inhibition of neuroinflammation, oxidative stress, and apoptosis. Furthermore, we demonstrated that downregulation of NF-kB and p53 is likely to be the mechanisms by which proinflammatory mediators and apoptosis are reduced in the rd10 retina, respectively.

CONCLUSIONS. VK28 and VAR10303 provided partial histologic and functional rescue of cones in RD10 mice. Our study demonstrated that iron chelation therapy might represent an effective therapeutic strategy for RP patients.

Keywords: rd10 mice, iron-chelating drugs, photoreceptors, vision rescue, microglia

Retinitis pigmentosa (RP) is a genetically and phenotypically heterogeneous group of inherited retinal disorders. The most common forms of RP are caused by mutations in genes essential to rod photoreceptor function and metabolism, resulting in rod cell death and subsequent night blindness. Cones that are present at a low density throughout the retina, with a sharp peak in the center of the fovea and represent 5% of all photoreceptors in the human eye, do not express a mutated protein, and degenerate secondarily to rods. Although different approaches to treating retinal degeneration are being explored, few effective treatments are currently available for RP patients. Therefore, establishing efficacious treatment strategies would create new perspectives for patients with retinal degeneration.

Growing evidence suggests that multiple mechanisms are involved in inducing photoreceptor cell death in RP. Oxidative stress has previously been found to correlate with photoreceptor cell death in RP. Indeed, treatments with antioxidants showed a reduction in oxidative damage imposed by dying rods in the rd1 and rd10 mouse models of RP. In addition, retinal iron accumulation is reported in RP animal models. Administration of iron chelators was shown to reduce iron-associated oxidative stress and attenuates photoreceptor degeneration in RP mouse models. Moreover, neuroinflammation is reported to be associated with RP and activated microglia are observed in human RP patients and RP animal models. We have shown that suppression of microglia activation promoted photoreceptor cell survival in the rd10 mouse model of RP, confirming the involvement of neuroinflammation in RP progression. Furthermore, photoreceptor death is reported to be related to the downregulation of the mechanistic target of rapamycin pathway in RP mouse models. Collectively, it appears that multiple mechanisms contribute to photoreceptor cell degeneration in RP.

Due to the complex pathology involved in RP progression, RP treatment poses a perplexing challenge. In this study, we explore iron chelation therapy in the rd10 mouse model of RP. To this end, we tested two iron-chelating compounds, 5(4-(2-hydroxyethyl) piperazin-1-yl (methyl)-8-hydroxyquinoline (VK28) and its chimeric derivative 5(3(N-methyl-N-propargyaminomethyl)-quinoline-8-oldihydrochloride (VAR10303), in the rd10 mouse. We have previously shown that VK28 and
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VAR10303 exerted multiple effects including neuroprotection and neurorestoration in the animal models of Alzheimer and Parkinson’s diseases.21–28 VK28 has been shown to have neuroprotective activity against 6-hydroxydopamine and 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) neurotoxicity, iron accumulation, and microglial activation,27,29 while VAR10303 attenuates the apoptotic cascades (induction of B-cell lymphoma 2 [Bcl-2] and reduction of Bax), and increases the expression levels of Bcl-2, neurotrophic factors including brain-derived neurotrophic factor (BDNF) and glial cell line-derived neurotrophic factor (GDNF), and phosphorylation of Akt and Glycogen synthase kinase 3 (GSK-3 β).24–26 Here, we have demonstrated that the two compounds provided significant histologic and functional rescue of photoreceptors and improved visual behaviors in rd10 mice.

METHODS

Animals and Drug Administration

Wild-type (WT; C57BL/6) mice, rd10 mice, and Cx3cr1GFP/GFP mice were obtained from Jackson Laboratory (Bar Harbor, ME, USA). Cx3cr1GFP/GFP mice were used to cross with WT (C57BL/6J) mice to obtain heterozygous Cx3cr1+/GFP mice, and also to cross with rd10 mice to generate rd10/Cx3cr1+/GFP mice, in which retinal microglia express green fluorescent protein (GFP) in a model of retinal degeneration. The littersmates from Cx3cr1+/GFP mice and rd10/Cx3cr1+/GFP mice were used for the experiments. The genotypes of mouse litters were determined by PCR and confirmed by Southern blot analysis of genomic DNA from tail biopsies. Animals were bred and maintained at the Centralised Animal Facilities of The Hong Kong Polytechnic University on a 12-hour light-dark cycle with room illumination of approximately 50 lux and food ad libitum. All experimental procedures were approved by the Animal Subjects Ethics Sub-Committee (AESC) at The Hong Kong Polytechnic University and conducted in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

VAR10303 was synthesized by incorporating the neuroprotective N-propargyl moiety of the rasagiline into the antioxidant-iron chelator moiety of an 8-hydroxyquinoline derivative of VK28. VK28 or VAR10303 (5 mg/kg) was injected intraperitoneally to rd10 and rd10/Cx3cr1+/GFP mice every other day, 48-hours apart, starting from postnatal day 14 (P14). Rd10 mice in control groups received the same volume of PBS intraperitoneally to rd10 and rd10/Cx3cr1+/GFP mice, at the same time points.

Immunocytochemistry and Confocal Imaging

Mice were deeply anesthetized with a mixture of ketamine hydrochloride (30–40 mg/kg) and xylazine (3–6 mg/kg) at different time points. Retinas were dissected free of the vitreous and sclera in carb oxygenated Ames’ Medium (Sigma, St. Louis, MO, USA), and then fixed in 4% paraformaldehyde (PFA) in 0.1 M phosphate buffer, pH 7.4, for 0.5 to 1 hour. Some of retinas were sectioned serially at a thickness of 10 to 12 μm using a cryostat. Both whole-mounted retinas and vertical sections were blocked in a solution containing 4% normal goat serum (NGS), 1% bovine serum albumin (BSA), and 0.3% Triton X-100 in PBS for 1 hour. Retinas were then incubated in a primary antibody from overnight to 5 days at 4°C. The following primary antibodies were applied: rabbit antibody to GFP (1:500; Invitrogen, Carlsbad, CA, USA), rat antibody to C3a (1:500; AbD Serotec, Raleigh, NC, USA), rat anti-mouse CD68 (1:500; AbD Serotec), rabbit anti-red/green opsin (1:500; Chemicon, Temecula, CA, USA). After rinsing, a secondary antibody conjugated to either Alexa 488 or Alexa 594 (1:500; Invitrogen) was applied for 2 hours at room temperature. Whole-mounted retinas and vertical sections were rinsed and cover slipped in Vectashield mounting medium (Vector Laboratories, Burlingame, CA, USA).

Confocal micrographs of fluorescent specimens from retinal whole-mounts and vertical sections were captured using a Zeiss LSM 700 Meta Axioplan 2 laser scanning confocal microscope (Carl Zeiss, Oberkochen, Germany) equipped with argon and helium-neon lasers. Plan-Apochromat ×63/1.4 or ×100/1.4 oil immersion objective lenses were used. Images scale was calibrated, and if necessary, brightness and contrast were adjusted using Photoshop CS3 software (Adobe Systems, San Jose, CA, USA).

Data Analysis

For measurement of outer nuclear layer (ONL) thickness, some retinas were sectioned serially at a thickness of 10 to 12 μm using a cryostat. Vertical sections passing through the optic nerve head were analyzed by counting the number of photoreceptor nuclei stained with diamidinophenylindole (DAP). Two measurements were taken at 200 μm (central region) and 1 mm (midperipheral region) from the optic nerve on both sides. Quantification of surviving cone photoreceptors was conducted in retinal whole-mounts, using a ×40 objective (numerical aperture: 0.85). Sampling areas were two fields of 240 × 240-μm squares along the dorsal-ventral axis of retinal whole-mounts, 1 mm (midperipheral region) from the optic nerve on both sides. Numbers of surviving cones were counted per grid directly under microscope or from z-stacked confocal images. The raw counts were then converted into cells/millimeter squared.

Electroretinographic Analysis

Electroretinograms (ERGs) were recorded using an Espion ERG Diagnosys machine (Diagnosys, Littleton, MA, USA) as previously described by us.23 In brief, flash ERG was measured using a gold wire corneal electrode, a forehead reference electrode, and a ground electrode near the tail. Scotopic, rod-mediated responses were obtained from dark-adapted animals at the following increasing light intensities: 0.01 and 3 cd-s/m². Photopic, cone-mediated responses were performed following 10-minute light adaptation on the background light intensity of 30 cd/m². Recordings were obtained at the light intensity of 3 cd-s/m². From each animal, 15 waveforms were recorded and the values were averaged. The ERG b-wave amplitudes were measured from the trough of the a-wave to the peak of the first positive wave or, when the a-wave was not present, from baseline to the peak of the first positive wave.

Optokinetic Tracking

Optokinetic tracking was performed as previously described by us.23 Mice were placed on a platform positioned in the middle of an arena created by a quad-square of computer monitors. Vertical sine wave gratings (100% contrast) written in MATLAB (MathWorks, Natick, MA, USA) were projected on the computer monitors. The spatial frequencies tested were 0.05, 0.075, 0.1, 0.2, 0.3, 0.4, 0.5, 0.6 cycle per degree (cpd), at a constant speed of 12 degree per second. An infrared-sensitive small camera monitored the eye movement images.

ELISA Measurement

ELISA for TNF-α (cat no. 900-M54; PeproTech, Rocky Hill, NJ, USA), IL-6β (cat no. 900-K50; PeproTech), nuclear-factor kappa B (NF-κB) p65 (cat no. 7174C, Cell Signaling, Danvers, MA,
USA), caspase-3/7 (cat. no. 13503; ATT Bioquest, Sunnyvale, CA, USA), caspase-8 (cat. no. 22798; ATT Bioquest), hypoxia inducible factor (HIF)-1α (cat. no. DYC195-2; R&D Systems, Minneapolis, MN, USA), malondialdehyde (MDA; cat. no. 21012; Oxis Research, Portland, OR, USA), reduced glutathione (GSH), and oxidized glutathione (GSSG) expression in retinas were performed using ELISA kits, respectively, according to manufacturer’s instructions as previously described by us.34,35

Murine TNF-α ELISA development kit contains the key components required for the quantitative measurement of natural and/or recombinant murine TNF-α in a sandwich ELISA format. Mouse TNF-α is a glycoprotein composed of 156 amino acids produced mainly by monocytes and activated macrophages. The kit is recommended for assaying murine TNF-α (PeproTech).31,32

Total NF-kB p65 Sandwich ELISA Kit is a solid phase sandwich ELISA kit that detects endogenous levels of total NF-κB p65 protein. The kit detects both the phosphorylated and nonphospho-NF-κB p65 protein (Cell Signaling).31

Caspase-3 Assay Kit uses (Z-DEV/DI2R110) as fluorogenic indicator for assaying caspase-3 activity. Cleavage of R110 peptides by caspases generates strongly fluorescent R110 that can be monitored fluorimetrically at 510 to 530 nm with excitation of 488 nm, the most common excitation light source used in fluorescence instruments. This kit can be used to continuously measure the activities of caspase 3 and 7 in cell extracts and purified enzyme preparations using a fluorescence microplate reader or fluorometer (ATT Bioquest).31

Caspase 8 Activity Assay Kit is designed to monitor cell apoptosis by measuring caspase 8 activity. This kit uses (Ac-LEVD)2-R110 as a fluorogenic indicator for caspase 8 activity. Cleavage of rhodamine 110 (R110) peptides by caspase 8 generates strongly fluorescent R110, which is monitored at the emission between 520 and 530 nm with the excitation between 480 and 500 nm (ATT Bioquest).31

The Human/Mouse Total HIF-1α DuoSet IC ELISA kit contains the basic components required for the development of sandwich ELISAs to measure in cell lysates. The kit’s specificity for recognizing HIF-1α was demonstrated by Western blot analysis. An immobilized capture antibody specific for binds both phosphorlated and unphosphorlated (R&D Systems).31

Western Blotting Analysis

CD68 is a membrane glycoprotein present in lysosomes of macrophage-lineage cells32; the antibody used in this study is an affinity purified goat polyclonal antibody raised against a peptide mapping near the C-terminus of CD68 of mouse origin, and recognizes on Western blot a protein of 75 to 110 kDa specifically present in tissue macrophages that correspond to different degrees of protein glycocalyx related to cell activation and phagocytosis.34,36

Phospho-IkBα is a rabbit monoclonal antibody (clone 14D4) and produced by immunizing animals with a synthetic phosphopeptide corresponding to residues surrounding Ser32 of human IkBα. Phospho-IkBα detects endogenous levels of IκBα only when phosphorylated at Ser32. This Phospho-IkBα antibody recognizes a single band of 40-kDa by Western blot of extracts from HeLa and NIH3T3 cells and mouse tissues (Cell Signaling).34,37

The IkBα antibody is a mouse monoclonal antibody to 112B2 to NFKBIA, and nuclear factor of kappa light polyepitide gene enhancer in B cells inhibitor, alpha. The IkBα antibody is produced by immunizing animals with a synthetic peptide corresponding to residues near the carboxy-terminus of human IkBα. The monoclonal antibody detects endogenous levels of total IkBα protein and recognizes a single band of 39-kDa by Western blot of extracts from HeLa cells and mouse livers (Cell Signaling).30

Nuclear p53 antibody is produced by immunizing animals with a synthetic peptide corresponding to residues surrounding Ser20 of human p53. p53 immunoreactivity detected with this antibody is exclusively nuclear. This antibody is highly specific for detecting a major protein band at 53 kDa that has the same mobility as recombinant p53 (Cell Signaling).30

Phospho-p53 antibody is a polyclonal antibody and produced by immunizing animals with a synthetic phosphopeptide corresponding to residues surrounding Ser15 of human p53. This antibody detects endogenous levels of p53 only when phosphorylated at serine 15.38 This antibody does not cross-react with p53 phosphorylated at other sites and recognizes a single band of 53-kDa by Western blot of extracts from MvILu cells (Cell Signaling).30

Transferrin antibody (11D5) is a mouse monoclonal antibody, raised against placental transferrin receptor or transferrin of human origin, and is recommended for detection of transferrin of mouse, rat, and human origin by Western blotting. The antibody recognizes a band of approximately 72-kDa by Western blot analysis of extracts from Hep G2 whole cell and rat sympathetic neurons (Santa Cruz Biotechnology).39

Statistical Analysis

Data were represented as means ± SDs. ANOVAs with Bonferroni’s and Dunnett’s post hoc tests for multiple comparisons were performed with Origin (OriginLab, Wellesley Hills, MA, USA). Software programs written in MATLAB on full data sets were used to detect significant differences in the means. A P value less than 0.05 was considered statistically significant.

RESULTS

VK28 and VAR10303 Partially Rescued Photoreceptor Morphology in the rd10 Mouse Retina

Rd10 mice are a well-characterized mouse model of RP, in which a mutation of the beta subunit of rod-specific phosphodiesterase (PDE6β) gene causes the massive degeneration of rod photoreceptors followed by cone photoreceptors.40 Mutations of the same gene have been linked to some
forms of human RP. Rod degeneration begins around P18 and the peak of photoreceptor degeneration is reached at P25 in the rd10 mouse retina. Because of the late initiation and mild degeneration, the rd10 mouse is increasingly being used for research to develop new experimental therapies for RP. We treated rd10 mice with VK28 or VAR10303 starting from P14, which is a few days earlier before the initiation of rod degeneration, and performed morphologic analysis at P26 and P30.

To assess the effectiveness of VK28 and VAR10303 on rod survival, the numbers of photoreceptor nuclei (97% of photoreceptor nuclei in the ONL of the mouse retina are rods) were counted in retinal vertical sections. At P26, the ONL in rd10 mice contained 3 to 5 rows of photoreceptor nuclei, while 14 to 16 rows were observed in normal WT mice (\( P < 0.01 \); Figs. 1A, 1D, 1I). Compared with rd10 controls, VAR10303-treated eyes had a thicker ONL (\( P < 0.05 \); Figs. 1C, 1I), indicating the reduction of rod cell death in VAR10303-treated rd10 mice. A thicker ONL was also observed in VK28-treated rd10 mice compared with rd10 controls (\( P > 0.05 \); Figs. 1B, 1I).

To characterize the protective effect of VK28 and VAR10303 on cone photoreceptors, we labeled whole-mounted rd10 retinas with an antibody specific for red/green cone opsins, and studied cone morphology and quantified cones (Figs. 1E–H). Cones in WT mouse retinas show...
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**FIGURE 2.** VK28 and VAR10303 partially rescue photoreceptor function in the rd10 mouse retina. (A, B) Representative scotopic ERG responses to 0.01- and 3-cd-s/m² light intensities from P26 rd10 mice treated with PBS, VK28, or VAR10303 from P14 to P25. Scotopic ERG responses from C57BL/6 (blue curve) are shown as comparisons. Average scotopic b-wave amplitudes elicited at both 0.01- and 3-cd-s/m² light intensities are shown on the right panel. Data are expressed as the mean ± SD. (C) Representative photopic ERG responses to 3-cd-s/m² light intensity from P26 rd10 mice are on the left, and averaged photopic b-wave amplitudes elicited at 3-cd-s/m² light intensity are on the right. (D) Average scotopic b-wave amplitudes elicited at 3-cd-s/m² light intensity were from P30 rd10 mice treated with PBS, VK28, or VAR10303 from P14 to P29. (E) Average scotopic b-wave amplitudes elicited at 3-cd-s/m² light intensity were from P30 rd10 mice treated with PBS, VK28, or VAR10303 from P14 to P29. Values are means and SDs and were analyzed using the Student’s t-test. **P < 0.01, ns, not significant; n, number of mice in each group.

**FIGURE 3.** VK28 and VAR10303 improved visual acuity in rd10 mice. Photopic visual acuity was measured by the optokinetic response under light-adapted condition. (A) A single-frame video camera image of a mouse surrounded by 360° of sine wave gratings. (B) Average visual acuity was evaluated from P26 rd10 mice following treatments with PBS, VK28, or VAR10303. Results are presented as means ± SDs and were analyzed using the Student’s t-test. **P < 0.01. ns, not significant; n, number of mice in each group.

VK28 and VAR10303 Partially Rescued Photoreceptor Function in the rd10 Retina

Functional rescue of photoreceptors in drug-treated rd10 mice were monitored by ERG. Because a-waves were variable and small and couldn’t be measured reliably, we thus chose b-waves as an indirect measure of photoreceptor function for the comparison of visual function. Representative ERG recordings are shown in Figure 2. Following a 12-day treatment, we observed markedly increased amplitudes of scotopic b-wave in P26 rd10 mice receiving VAR10303 treatment, compared with PBS-treated rd10 controls (P < 0.01; Figs. 2A, 2B). Similarly, photopic ERG b-wave amplitudes were significantly larger in VAR10303-treated rd10 mice than in PBS-treated rd10 controls (P < 0.01; Fig. 2C), suggesting improved retinal function. On the other hand, VK28 didn’t show any obvious effects on the functional rescue of photoreceptors under both scotopic and photopic conditions at P26 (P > 0.05; Figs. 2A–C). However, the protective effect of VK28 became prominent following the long-time treatment with VK28. Scotopic b-wave amplitudes were significantly higher in VK28-treated P30 and P42 rd10 mice, compared with age-matched PBS-treated rd10 controls (P < 0.05; Figs. 2D–E), indicating a better retinal function rescue. Similarly, scotopic b-wave amplitudes were appropriately 3- and 2-fold higher in VAR10303-treated rd10 mice than in PBS-treated rd10 controls (P < 0.01) and P42 (P < 0.05) rd10 mice than their age-matched rd10 controls, respectively (Figs. 2D–E), suggesting a significant rescue of retinal function with VAR10303 treatment.

To evaluate spatial visual performance in rd10 animals treated with VK28 or VAR10303, we measured mouse optomotor responses to moving gratings of different spatial frequencies at P26 (Fig. 3A). We found that photopic visual acuity in VAR10303-treated rd10 mice was significantly better compared with age-matched rd10 controls. At P26, average cone density was comparable among PBS- and VK28- and VAR10303-treated rd10 retinas and WT retinas (data not shown). However, there was a substantial loss of cone photoreceptors in PBS-treated P30 rd10 retinas when compared with age-matched WT (P < 0.001; Fig. 1K). Treatments with VK28 or VAR10303 enhanced cone survival in rd10 retinas, with average density of cones markedly higher than PBS-treated P30 rd10 retinas (P < 0.01; Fig. 1K). Taken together, two iron-chelating drugs provided significant histologic rescue of cones in rd10 mice.
than that in PBS-treated rd10 controls ($P < 0.01$; Fig. 3B). However, the visual acuity in VAR10303-treated rd10 mice was still poorer than that in WT control mice ($P < 0.01$; Fig. 3B), indicating partial rescue of retinal function by VAR10303 treatment. We did not observe improved visual performance in VK28-treated rd10 mice at P26, which was consistent with ERG results in the same age of VK28-treated rd10 mice.

**Figure 4.** VK28 and VAR10303 reduced microglia activation and productions of proinflammatory mediators in the rd10 mouse retina. Microglia maintain a ramified morphology in Cx3cr1$^{+/GFP}$ control mouse retinas (A), and CD68 staining is undetectable ([E, I] arrowheads). Microglia show amoeboid morphology with few processes and rounded cell bodies in PBS-treated P26 Cx3cr1$^{+/GFP}$/rd10 retinas ([B] arrows), and CD68 staining is intense and widespread ([F, J] arrows). VK28 and VAR10303 treatments preserved the ramified appearance of microglia in Cx3cr1$^{+/GFP}$/rd10 retinas ([C, D] arrowheads), and reduced CD68 immunoreactivity ([G, H, K, L] arrows). (M) Western blotting analysis and densitometry of CD68 protein expressions from PBS-, VK28-, and VAR10303-treated P26rd10 mouse retinas and WT mouse retinas ($n = 8–12$ in each group). (N-O) ELISA analysis of TNF-$\alpha$ ([N]) and IL-6 ([O]) protein expression levels in the retina of WT and PBS-, VK28- and VAR10303-treated rd10 mice. Results are presented as means $\pm$ SDs and were analyzed using the Student’s $t$-test. *$P < 0.05$, **$P < 0.01$; ns, not significant. Scale bar: 50 $\mu$m.

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**VK28 and VAR10303 Suppressed Microglia Activation in the rd10 Retina**

Microglial activation has been shown to contribute to the pathogenesis in RP retinas. We investigated whether VK28 and VAR10303 treatments suppressed microglial activation in the rd10 retina. Microglia have a ramified appearance in Cx3cr1$^{+/GFP}$ control mouse retinas (Fig. 4A, arrowheads).
gradual morphologic change in microglia from a ramified appearance into an amoeboid shape with retracted processes and rounded cell bodies was observed in the PBS-treated Cx3cr1−/−/GFP/rd10 retina (Fig. 4G, arrows). However, microglia maintained ramified morphology in VK28- and VAR10303-treated Cx3cr1−/−/GFP/rd10 mouse retinas (Figs. 4J, 4D, arrowheads). To confirm the status of microglial activation, we performed CD68 staining, a marker for microglia activation. CD68 immunoreactivity was undetectable in Cx3cr1−/−/GFP control mouse retinas (Figs. 4E, 4I), while CD68 immunoreactivity was widespread and intense in the rd10 retina (Figs. 4F, 4J). VK28 and VAR10303 treatments led to reduction of CD68 immunoreactivity (Figs. 4G, 4H, 4K, 4L), which was further confirmed by Western blotting analysis (Fig. 4M).

Moreover, we measured the expression levels of the inflammatory mediator TNF-α and IL-6 by ELISA analysis. Compared with WT retinas, TNF-α and IL-6 protein expressions increased approximately by 2- and 3-fold in rd10 mouse retinas, respectively (P < 0.01; Figs. 4N, 4O). However, VK28 and VAR10303 treatments markedly reduced the protein expression levels of TNF-α and IL-6 (P < 0.01; Figs. 4N, 4O), indicating attenuation of neuroinflammation in drug-treated rd10 retinas.

Furthermore, we sought to gain more insight into mechanisms underlying the suppression of microglial activation by VK28 and VAR10303 treatments. Nuclear-factor kappa B is known to induce the transcription of proinflammatory genes. Nuclear-factor-kB is bound to IκB in the cytoplasm under normal physiological conditions. Under pathologic conditions, IκB is phosphorylated and degraded, leading to activation of NF-κB. Subsequently, NF-κB is translocated to the nucleus and activates the expression of proinflammatory cytokine genes. We thus studied the expression level of NF-κB in the nucleus, and IκB in the cytoplasm. Nuclear factor-kB exists in several dimeric forms, predominantly the p50/p65 heterodimer. ELISA analysis demonstrated a significant increase in phosphorylated-NF-κB p65 in PBS-treated rd10 retinas (P < 0.01; Fig. 5A), but the increases were significantly counteracted by VK28 and VAR10303 treatments, leading to a significant reduction in the expression levels of NF-κB p65 in rd10 retinas (Fig. 5A).

Meanwhile, Western blotting analysis showed that phosphorylated-IκBα (p-IκBα) expression increased by 4.2-fold in PBS-treated rd10 retinas, compared with WT retinas (Fig. 5B). After VK28 and VAR10303 treatments, p-IκBα expression levels were decreased by 1.4- and 2.6-fold in rd10 mouse retinas, respectively, when compared with PBS-treated rd10 retinas (Fig. 5B). Conversely, IκBα protein expression decreased by 10-fold in PBS-treated rd10 mice (Fig. 5C), compared with WT retinas. On the other hand, IκBα protein levels increased by 7.9- and 1.6-fold in VAR10303- and VK28-treated rd10 retinas, respectively (Fig. 5C). Taken together, our data showed that reduction of proinflammatory mediators by VK28 and VAR10303 treatments in the rd10 retina correlated to downregulation of NF-κB.

**The Mechanisms of Actions of VK28 and VAR10303 in the rd10 Retina**

To further explore the mechanisms underlying the rescue of cone photoreceptors, we evaluated the several mechanisms that are previously reported to contribute to photoreceptor cell death in RP. Altered iron metabolism is implicated in retinal degenerations including RP. We thus performed Western blotting analysis to assess the expression level of the iron metabolism-related protein transferrin. Transferrin expression have been shown to increase during the course of retinal degeneration in rd10 mice. We have confirmed the upregulation of transferrin expression in rd10 retinas (Fig. 6A).

Systemic administration of VK28 or VAR10303 significantly downregulated transferrin expression in rd10 retinas (Fig. 6A), indicating a possible protective therapeutic approach against iron overload-induced photoreceptor degeneration.

Moreover, oxidative stress causes photoreceptor dysfunction and cell death in RP mouse models. We measured several biomarkers of oxidative stress, including MDA, reduced GSH, GSSG, and GSH/GSSG. Malondialdehyde is a common lipid peroxidation product that accumulates in many pathophysiological processes. We found that MDA expression increased approximately by 2-fold in rd10 retinas compared with WT retinas (P < 0.01; Fig. 6B). VK28 and VAR10303 treatments significantly reduced the expression levels of MDA in rd10 retinas compared with PBS-treated rd10 retinas (P < 0.01; Fig. 6B).

The GSH/GSSG status is commonly used in measuring oxidative stress status, and a reduction in GSH/GSSG ratio reflects a reduced antioxidant capacity. A significant reduction in GSH/GSSG ratio was observed in rd10 retinas, compared with WT retinas (P < 0.01; Fig. 6C), and VAR10303 and VK28 treatments significantly increased the GSH/GSSG ratio (P < 0.01; Fig. 6C), indicating an increased antioxidant capacity.

Furthermore, we investigated whether VAR10303 and VK28 regulated the apoptotic pathway. Caspases are a family of genes important for the apoptotic response. We evaluated the expression levels of the initiator caspase, caspase-8, and the effector caspases, caspases-3/7. We found that the protein expression levels of caspase-3/7 and -8 were significantly higher in PBS-treated rd10 retinas than in WT controls (P < 0.01; Figs. 6D, 6E). Treatments with VK28 and VAR10303 significantly decreased the protein levels of caspase-5/7 and -8 in rd10 retinas (P < 0.01; Figs. 6D, 6E).

Hypoxia-inducible factor 1α, an oxygen-sensitive transcriptional activator, plays an important role in cellular response to systemic oxygen levels in mammals. We found that HIF-1α was...
upregulated in rd10 retinas (Fig. 6F), and the increases were counteracted by VAR10303 and VK28 treatments \((P < 0.01; \text{Fig. 6F})\). The key tumor suppressor protein p53, a downstream target of HIF-1α, has a diverse range of functions, including the ability to promote apoptosis. \(^48\) Western blotting analysis showed that total p53 protein level in the nucleus was comparable among WT retinas, PBS-, VK28- and VAR10303-treated rd10 mouse retinas (Fig. 6G). However, phosphorylated p53 (p-p53) expression level increased in rd10 retinas compared with WT retinas (Fig. 6G). Treatments by VK28 and VAR10303 dramatically decreased p-p53 protein levels in rd10 retinas (Fig. 6G). These findings indicated that downregulation of nuclear p53 was likely to be the mechanism underlying reduced apoptosis in photoreceptors in the rd10 retina after VK28 and VAR10303 treatments.

Taken together, our data suggested that VK28 and VAR10303 exerted protective effects likely through regulation of iron bioavailability, oxidative stress, and photoreceptor apoptosis, leading to partial rescue of photoreceptors in rd10 mice.

**DISCUSSION**

In this study, we investigate the effects of two iron-chelating drugs on the rescue of cone photoreceptors in the rd10 mouse model of RP. We demonstrated a significant rescue of the structure and function of cone photoreceptors by the two iron-chelating drugs, VK28 and VAR10303, for extended periods, during a time that they would ordinarily degenerate in rd10 mouse retina. Moreover, we showed that the two compounds...
exerted their effects by altering the profile of cone degeneration through the possible synergism of anti-inflammatory, antioxidant, and antiapoptotic mechanisms of actions in the rd10 retina.

Neuroinflammation is an underlying component of a diverse range of chronic neurodegenerative disorders. Downregulation of HIF-1α and Parkinson’s diseases. Moreover, HIF-1α proapoptotic proteins in the animal models of Alzheimer’s showed that VAR10303 and VK28 significantly downregulated caspase-3. Consistent with current findings, we have previously reported that downregulation of NF-kB was likely to be the mechanism by which the proinflammatory mediators were downregulated. Consistently, increased activity of NF-kB and its neurotoxic role in photoreceptor cell death have been previously reported in the rd1 mouse model of RP.

Iron-associated oxidative stress plays an important pathogenic role in neurodegenerative diseases including AMD and RP. Similarly, iron accumulation and increased expression of iron-regulating protein transferrin were shown in rd10 retinas. We have demonstrated that VAR10303 and VK28 downregulated transferrin expression in rd10 retinas, which might oppose iron overload-induced cone degeneration. Moreover, VAR10303 and VK28 counteracted oxidative stress by restoring antioxidant enzymes and limiting iron bioavailability in rd10 retinas, resulting in attenuation of oxidative stress injury in the rd10 retinas. The findings are consistent with our previous observation that two compounds are capable of chelating iron and suppressing iron-induced oxidative stress, and subsequently attenuating neuronal loss in Alzheimer’s and Parkinson’s diseases animal models. In accordance, a recent study demonstrated a reduction in retinal oxidative stress by the iron chelator desferrioxamin in rd10 mouse retina.

Furthermore, it has been shown that caspase-dependent upregulation is involved in photoreceptor cell death in RP. Indeed, the upregulation of caspase-3 has been reported in the rd1 mouse model of RP. Administration of VAR10303 and VK28 resulted in a significant reduction of caspase-3 and -8 protein levels, which might reflect the possible effect of the compounds on photoreceptor apoptosis. Consistent with current findings, we have previously shown that VAR10303 and VK28 significantly downregulated proapoptotic proteins in the animal models of Alzheimer’s and Parkinson’s diseases. Moreover, downregulation is observed during photoreceptor degeneration in several mouse models of RP. Downregulation of HIF-1α and p53 that we observed in the present study was probably the mechanisms underlying reduced apoptosis in photoreceptors in the rd10 retina.

Finally, VK28 and VAR10303 also protected rods from degeneration in the rd10 retina. The partial rescue of rod photoreceptor cells might subsequently improve the survival of cones. Indeed, previous studies suggested that rod-derived cone viability factor (RdCVF) maintained the viability and function of cones in the retina, and mice that lack RdCVF showed a gradual loss of photoreceptors. Therefore, the rescue of rods by VK28 and VAR10303 might partially account for the survival of cones in the present study.

In summary, we demonstrated that VK28 and VAR10303 provided significant rescue of cone photoreceptors in RP mice, and the neuroprotective effects of the two compounds on cone photoreceptors correlated to inhibition of microglial activation, oxidative stress, and apoptosis. Therefore, iron-chelation therapeutic strategy may hold great prospects for an effective treatment of RP.

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References

Iron-Chelating Drugs Protect Cones


